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Spectral Studies of Extra-Terrestrial Materials

Catarina Fernandes

Licenciatura em Geologia Aplicada e do Ambiente, Universidade de Lisboa

Submitted for the degree of Doctor of Philosophy

Faculty of Science, The Open University

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</tr>
<tr>
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</tr>
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Abstract

Experiments were made using a state-of-the-art UV-Vis microspectrophotometer (MSP) in order to assess if the instrument is suitable for use on spectroscopy of terrestrial and extra-terrestrial materials. This new instrument brings advantages that cannot be found in instruments currently in use: it requires only extremely small samples (min of \( \sim 2 \, \mu m \)) and it is very quick to use (little sample preparation and spectra taken in less than 2 min).

If suitable, the instrument could help to show the relationship between meteorites and their parent bodies and in the study of very small fragile samples, such as cometary samples and Interplanetary Dust Particles (IDPs).

Using samples of minerals commonly found in meteorites, it was concluded that the instrument is suitable for the study of these materials, however it has some limitations and certain conditions need to be met.

The method was then applied on two grains from comet 81P/Wild 2 returned by the space mission Stardust. Further limitations were found with these samples caused by the fact that they are covered in aerogel and embedded in gold foil. Results indicate however, that the samples seem to be composed of a mixture of different materials.

Results from the study of HED (howardites, eucrites and diogenites) type meteorites proved that if the conditions are met, the technique is suitable and comparable to other instruments and can be used to match the spectra of meteorites to that of their possible asteroidal parent bodies.

A complementary investigation studied the effects of impact by shock on the spectra of rocks using a Light Gas Gun and Near Infrared spectroscopy with the goal of investigating the effects of weathering on the spectra of asteroids. It was found that there is a change in the spectra of the samples and a relationship with a change in composition of the impacted area.
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1 Small Solar System Bodies

1.1 Introduction

Understanding the origin and evolution of the Solar System is one of the most important topics studied by the scientific community, as without knowledge of how the Sun and planets formed, it is difficult to understand how life originated, and whether it might be present on bodies other than Earth. The purpose of the work described by this thesis is to study and analyse extra-terrestrial materials, in particular materials which originated from small bodies, such as asteroids and comets, in order to comprehend some of the processes that the material experienced during its history. It is important to outline the mechanisms of formation of the Solar System in order to understand better the origins and compositions of extra-terrestrial material. Paradoxically, it is the study of these materials that leads back to theories of formation of the Solar System. The method selected to carry out this study was laboratory-based UV-Vis spectroscopy of grains derived from asteroids and comets. In this chapter, a brief introduction to the small Solar System bodies investigated is given, and a rationalisation of why the spectroscopic technique was selected, and what specific questions are hoped to be answered.

Since the Big Bang, circa 13.7 Ga (billion years) ago, stars have formed and died in a continuous cycle during which elements (and their isotopes) heavier than hydrogen are produced (Figure 1.1). The death of stars and the consequential spread of material feed the interstellar medium with heavier elements that are incorporated in molecular clouds. These clouds become unstable, collapse and contract, and start forming other stars. The Solar System is, therefore, composed of remains of previous stars.
The most accepted theory for the formation of planetary systems describes this process as beginning in a **dense cloud** (Figure 1.2) with higher concentrations of gas and dust than its surrounding medium. A region of slight over-density causes this gas and dust to contract when the cloud reaches a certain size, mass and temperature. As particles collide, other effects start to influence the collapse of the cloud, such as magnetic fields, rotation, and effects external to the cloud. Temperature and pressure rise and one, or a cluster, of **protostars** start to emerge (e.g. Larson 2003 and references therein; Cole and Woolfson 2002).

![Figure 1.1: Cartoon to illustrate the chain of events from the Big Bang through to the formation of the Solar System and the origin of life on Earth (credit: NASA/JPL-Caltech)](image-url)
When the temperature and pressure reach the stage where fusion reactions take place, the protostar becomes a star – the Sun in the case of the Solar System. As this occurs, the remaining dust, guided by angular momentum, flattens out, forming a disc of debris around the star – the *protoplanetary disc* (Figure 1.3) (e.g. Greaves 2005; Connolly 2005 and references therein). Within the disc, particles continue to grow by condensation, collisional accretion and coagulation and increase in size, leading to bodies of 0.1 to 10 km across – *planetesimals*. When it reaches approximately 10 km in diameter, a planetesimal has gained sufficient mass for its own gravitational field to attract more and more material, growing, and affecting other passing planetesimals. Collisions become more common, leading to fragmentation of planetesimals and then larger bodies from the accretion of these fragments – *planetary embryos*. In the Solar System, as planetary embryos continued to grow, the terrestrial planets formed in the hotter inner Solar System and cooler, gaseous giants are formed further from the Sun. The Sun’s T-Tauri phase, in which stellar winds removed a great quantity of material from the disk, removed most of the remaining material which had not been incorporated in the terrestrial and giant planets. This and further information can be found in several publications such as Cyr et al. 1998; Ehrenfreund and Charnley 2000; Alexander et

Figure 1.3: The protoplanetary disc HH-30 (credit: NASA images)

The gravitational perturbation of Jupiter and other giant planets prevented the accretion of some of the material into a planet, forming the asteroid belt, and flung icy bodies towards the edge of the Solar System, forming the Oort cloud (Figure 1.4).

Some of the knowledge about the processes of formation of the Solar System is derived from astronomical images of these processes occurring in other protoplanetary systems, assumed to be analogous to the early Solar System. Most of it, however, has been acquired from the study of meteorites which contain remnants of the various phases of the formation of the planetary system. From meteorites, we can also derive information of conditions and timescales of the events which took place as the Solar System formed. Furthermore, the development of new techniques and missions, which allow us access to other bodies (e.g. dust from pristine comets and interstellar space), takes the study of formation and evolution of ours, and other planetary systems one step further. A description of the objects from which the materials that were studied for this thesis were derived is presented in this chapter.
1.2 Interstellar Dust

Interstellar Dust Grains (ISD) constitute ~0.1% of the solid matter in the Galaxy, but they are a major constituent of diffuse and dense clouds in the interstellar medium (Freund and Freund, 2006). They are the building blocks for the formation of stars and planets, asteroids, comets and Kuiper Belt objects. The interstellar medium consists mainly of hydrogen gas, ~10% He atoms and ~0.1 % dust particles of C, N and O (Ehrenfreund and Charnley 2000 and references therein). These are present in the molecular cloud from where, in the case of the Solar System, the protostar and a protoplanetary disc evolved. Li and Greenberg (1997) developed models based on astronomical observations and laboratory measurements to develop theories of core and mantle components of ISD. They concluded that most particles have silicate cores with a mantle of complex organic molecules. There are also small carbonaceous particles and ices. These particles have different rates of accretion and different abundances depending on the surrounding environment.

Results from Li and Greenberg (1997) are accepted by other authors. Mann et al. (2005) confirmed that, at the early stages of the formation of the Solar System, the particles which are colliding with each other consist either of a silicate core and an organic refractory mantle, a few tenths of micrometres in size, or are very small carbonaceous particles with an outer mantle of volatile ices. Cold gas-phase chemistry can form simple species and short carbon chains, but the production of organic molecules can be enriched by thermal and energetic processes such as UV and cosmic ray irradiation. The organic refractory material started out as ices of simple chemical compounds (e.g. CO, N₂, O₂). After millions of years of UV photo-processing in interstellar space, the icy mixture changes into a carbon-rich and oxygen poor refractory material containing very different organic molecules such as carbon chains, aromatic compounds and Polycyclic Aromatic Hydrocarbons (PAH).
(e.g. Greenberg and Hage 1991; Ehrenfreund and Charnley 2000) which might have played an important role in the origin of life.

1.3 Comets

Comets are small bodies (a few km in diameter) that orbit the Sun and that, at some point in their orbit, as they come closer to the Sun (approximately between the orbits of Jupiter and Saturn), their ices and dust begin to sublimate, and produce a coma and a tail. Comets formed approximately 4.57 Ga ago from the presolar nebula in the same processes that led ultimately to the formation of the Sun and planets. The composition of comets is dominated by ice (water, carbon monoxide and dioxide, methane and others), making it implicit that they formed at a distance from the Sun where ice could condense. Comets and asteroids differ in their volatile content (high abundances of volatiles in comets) and orbits (comets have more eccentric orbits with higher inclinations) which are linked to the different locations where they were formed. Asteroids were formed at a distance from the Sun at which significant quantities of water-ice are unlikely to be present. However, 'old' comets, which have lost most of their volatile composition, may come to resemble asteroids.

The Kuiper belt and the Oort cloud are the major reservoirs of comets in the Solar System (Figure 1.4). Long period comets (with orbital period of more than 200 years) constitute the largest population. They inhabit the outermost fringes of the Solar System, where they compose the Oort cloud. The Oort cloud, approximately 10,000 to 50,000 AU from the Sun, has never been viewed directly, but its existence is inferred from observations of the orbits of long period comets (Oort 1950). These comets are believed to have formed in the outer Solar System at the same time as the other bodies and were then scattered outwards by the gas giants, forming the Oort cloud (e.g. Fernandez and Brunini 2000). Objects in the outer Oort cloud can be gravitationally perturbed by passing stars, causing some comets to be thrown...
inwards, close enough to the giant planets to be affected by their gravity, and become short period comets. Others may be thrown out of the Solar System and are lost. Some single-apparition comets have parabolic and hyperbolic orbits, which cause them to permanently exit the Solar System after one pass by the Sun (Oort, 1950).

Much closer to the Sun than the Oort Cloud is the Kuiper Belt, stretching beyond Neptune, from ~ 30 to 60 AU, a reservoir of comets with shorter orbital periods – less than 200 years (e.g. Duncan et al. 1988; Weissman 1995; Jewitt 2004 and references therein). The orbits of the comets from the Kuiper Belt and the Oort cloud are distorted by gravitational interaction with the gas giants, extra-Solar System influences, and collisions and close encounters with other bodies, causing their orbits to become shorter and closer to the Sun. Eventually, the comets are scattered out of the Solar System or collide with the Sun. Short period comets can be further divided into families according to the position and characteristics of their orbits (Jewitt 2004). Halley family comets are comets with short periods but various inclinations including retrograde orbits. They are thought to have come from the inner Oort cloud and captured by gravitational processes of the gas giants (Bailey and Emel'yanenko 1996). Jupiter-family comets have small orbits and small inclinations and eccentricities and are believed to originate from the Kuiper Belt, which has a similar inclination and distribution (Duncan et al. 1988). They are possibly a result of the fragmentation of Kuiper belt objects. Centaurs are dynamically intermediate between the Kuiper belt and the Jupiter-family comets. Most of them appear asteroidal as they have lost most of their volatile content or because their surfaces consist of non-volatiles, but some have been observed to have a coma. They are strongly affected by the gravitational fields of the gas giants and most are ejected from the Solar System. The remaining Centaurs become Jupiter-family comets (Jewitt 2004).
Because comets formed and remain most of their 'lives' on the outer fringes of the Solar System, they are thought to be relatively pristine in composition (Weissman 1985). This means that they have not been greatly altered by thermal or fluid processing. They spend most of their 'lives' far from the Sun where temperatures are low and therefore they are not affected by heating and solar radiation. Also, because of their small sizes (usually less than 50 km diameter), gravity does not play an important role and consequently internal heating is not significant. It is generally accepted (e.g. Weissman 1985; Feldman 1987; Greenberg and Hage 1991; Greenberg and Li 1998) that their composition must be different from the remaining bodies of the Solar System and they may have preserved the composition and isotopic ratios of the early Solar System.

As a comet draws closer to the Sun, solar radiation causes sublimation of the material near the surface of the nucleus of the comet, forming a coma. Although the
Comets are small (generally less that 50 km in diameter) the coma can be larger than the Sun. A gas tail is formed and ionised by UV radiation and solar wind, forming a straight plasma tail. This tail follows the direction of the Sun's magnetic field. The force of the Sun's radiation pressure also forms a dust tail. Because the dust particles are of different sizes and travel at different velocities, this is a curved tail, which follows the path of the orbit of the comet and can stretch many thousands of kilometres (e.g. Mendis and Ip 1977) (Figure 1.5).

Each time a comet approaches the Sun, and tail formation takes place, additional material is lost, gradually resulting in a decrease in size of the nucleus. This evolutionary sequence, however, is not believed to process material within the nucleus, as material is constantly removed rather than being altered in situ - the cometary nucleus remains almost pristine (some thermal processing of the near surface takes place). Unlike planets and many asteroids, the composition of comets has not changed to a great extent since their formation, and they are believed to be the most primitive bodies in the Solar System. Comets can thus provide information on the original material from which the Solar System formed (e.g. Feldman 1987; Rotundi et al. 2002; Brownlee et al. 2003; Green et al. 2004).

Figure 1.5: The Dust and Ion Tales of Comet West. Credit: Observatoire de Haute, Provence, France
1.3.1 Cometary Missions

There have been several missions to comets over the past twenty-five years. One of the first, following Vega 1, was ESA's Giotto mission, which flew by Halley's comet in 1986, took the first close-up image of a cometary nucleus (Figure 1.6), and, for the first time, directly measured the size of the nucleus (Reinhard 1986). Results show an elongate, non-spherical nucleus ~15 km by 7-10 km in size. Irregular and spherical structures, similar to impact craters, were seen. The comet has very low albedo and most of the cometary activity seems to come from a few locations on the illuminated surface (e.g. Keller et al. 1986; Reinhard 1986).

Figure 1.6: Comet Halley in a composite of images taken by the Giotto probe on 14 March 1986 from 1,500 km away. The nucleus of Halley's Comet, 16 km long, is visible. The image shows gaseous jets from sublimating ices spewing from the surface of the nucleus (Credit: ESA)

Results from the mass spectrometers PIA, PUMA1 and PUMA2 on Giotto, Vega 1 and Vega 2, respectively (Kissel et al. 1986; Reinhard and Battrick 1986), show that H2O is the dominant parent molecule in the coma followed by CO2, NH3 and CH4. Daughter species, present in different abundances, include C+, CO+, S+, H+, He+, O+, OH+, and H3O+. There is a striking richness in C+, which cannot be accounted for by photodissociation of CO, CH4, or CO2 (Greenberg and Li 1998). There is the possibility that carbon atoms are released from the comet's surface directly, or the source may be the dust grains. In the dust, H, C, N, O, Na, Mg, Si, K, Ca, and Fe were detected (Reinhard 1986). Results from the Vega missions show similar
compositions of the dust grains which match the composition of CI chondrites more closely (Kissel et al., 1986). However, mass spectra taken on some of the particles of the comet indicate that the CI chondrite model for cometary particles cannot be the sole model, as the particles are richer in carbon and nitrogen than expected (Kissel et al. 1986; Reinhard 1986). Other dust grains are rich in H, C, N and O (CHON) supporting models that describe comets containing radiation-processed ices (e.g. Schutte et al. 1992). Ratios of silicate, CHONs and mixed particles show a heterogeneity that is consistent with different origin location of the particles within comet Halley (Clark et al. 1987). Composition of other particles may support the idea that comets are aggregates of interstellar dust particles, consisting of a silicate core embedded into a non-volatile organic mantle produced from ices by ultraviolet radiation before solar nebula condensation (Kissel et al. 1986).

ESA followed up the success of the Giotto mission with the Rosetta mission (e.g. Schulz, 2009). This is the first mission planned to rendezvous with and land on a comet; the spacecraft, launched in March 2004, will deliver a lander (Philae) to the nucleus of comet 67P/Churyumov-Gerasimenko in 2014 and make in situ measurements of the nucleus.

NASA has also had two successful cometary missions. Deep Impact, launched in January 2005, fired a 360 kg copper-rich projectile into the path of comet 9P/Tempel 1 in July 2005. When the projectile hit the comet, a crater was formed that liberated vast amounts of ice, gas and dust in the process (A'Hearn et al. 2005). Results from cameras and the spectrometer on board the spacecraft show that water and carbon dioxide gases had asymmetric surface distributions, suggesting a non-uniform composition of the nucleus (Sunshine et al. 2006). It was also the first time that water ice had been observed on the surface of a comet (Sunshine et al. 2006; Sunshine et al. 2007). However the amount of ice detected was insufficient to produce the amount of water and by-products measured following the impact, so
there may be sources beneath the surface that supply the coma (Sunshine et al., 2006). The spectrometer on telescope Keck 2 in Hawaii detected eight gases in the ejecta coming from the impact of the projectile: water, ethane, hydrogen cyanide, carbon monoxide, methanol, formaldehyde, acetylene and methane (Mumma et al. 2005). The Deep Impact mission has been recently extended to the EPOXI mission which flew by comet 03P/Hartley in 2010 (A'Hearn et al. 2008).

NASA's other successful cometary mission was Stardust, launched on February 7th 1999 to comet 81P/Wild 2. This mission collected samples of the comet's coma and it has recently been extended to the NEXT mission which will study the effects of the copper projectile from Deep Impact on comet 9P/Tempel 1. The Stardust mission, and results from analysis of the samples returned from comet 81P/Wild 2 are presented in Chapter 5.

### 1.4 Meteorites and Interplanetary Dust Particles (IDP)

Thorough reviews of the classification and properties of meteorites can be found in Papike 1998; McSween Jr. 1999; Hutchison 2004, from which much of the information in this section is taken. Meteorites are mostly fragments from asteroids, although there are separate populations from the Moon and Mars. They fall at random over the Earth's surface. Meteorites can be sub-divided into two large divisions: differentiated and undifferentiated materials. The former come from parent bodies that have been subject to planetary processes of melting and segregation, leading to fractionation of elements and minerals at a variety of scales, from macroscopic to sub-micron. Differentiated meteorites include irons (composed of nickel-rich iron metal), stony-irons (sub-equal mixtures of nickel-rich iron and silicates) and achondrites (silicate-rich meteorites). Undifferentiated meteorites, mainly chondrites, are of specific relevance to this study. They are silicate-rich meteorites that emanate from parental sources that have not suffered episodes of
major melting and fractionation. A short definition is found in Hutchison (2004):
"Chondrites are agglomerate rocks of silicate, sulphide and, usually, metal, whose
crystal composition closely approach the composition of the Sun, less its
hydrogen, helium and other highly volatile elements". The grains from which
meteorites initially aggregated were interstellar silicates, processed through a
molecular cloud prior to accretion in the protoplanetary disk. Although chondrites
have experienced varying degrees of thermal and hydrothermal processing since
accretion, the most primitive of chondrites still retain a record of the original
primordial cloud. See Appendix 1 for classification of a meteorite.

Chondrites can be further sub-divided into carbonaceous, ordinary and enstatite
classes. Enstatite chondrites are rare meteorites with a very high abundance of the
Fe-poor silicate mineral enstatite, iron occurs mainly as metal or sulphide. This
indicates a very reducing environment of formation. Ordinary chondrites, as the
name suggests, are the most common chondrites in the meteorite samples
available. They are further subdivided according to their iron content into H (High-
iron), L (Low-iron) and LL (Low-iron, Low-metal). Carbonaceous chondrites are rich
in magnesium-rich minerals and organic compounds, including amino acids. They
are the most primitive of the chondrites and have the same composition as the solar
nebula when the Solar System was formed (without the volatiles). They are sub-
divided into seven groups, CI, CM, CO, CV, CK, CR, CH. The first six groups are
named after the first meteorite described of that type (Ivuna, Mighei, Ornans,
Vigarano, Karoonda and Renazzo, respectively); CH chondrites have very high iron
contents. CMs and CIs contain large quantities of water and volatiles and are
thought to be the most primitive of the carbonaceous chondrites. Most have been
altered by aqueous processes but have not undergone heating. Possible parent
bodies for these meteorites are discussed in the next section.
The main constituents of almost all chondrites are chondrules, which are near-spherical, sub-mm, silicate assemblages, composed of olivine and/or pyroxene set in a feldspathic mesostasis. All chondrites, (apart from the CI group) contain calcium-aluminium-rich inclusions (CAIs), which carry isotopic anomalies that date them back to the birth of the Solar System. CAIs are highly irregular objects, some of which are surrounded by layered rims composed of Al-, Ti-, Ca-rich, Fe-poor oxides and silicates that are stable at high temperatures (Hutchison 2004).

Chondrites also contain minor components that have distinct origins: over the past 30 years, interstellar and circumstellar grains have been separated from meteorites. These grains were originally identified by the unusual isotopic composition of the noble gases that they carried (Anders 1991). Subsequently, the isotopic composition of the grains themselves were found to be very different from Solar System material (Hoppe and Zinner 2000). Presolar grains identified so far, and that provided information about their stellar sources, are diamonds, silicon carbide (mainstream and type X), graphite, silicon nitride, corundum and spinel.

Dust grains from comets (and asteroids) have been captured from Earth’s atmosphere through collection on plates carried on U2 planes flying through the stratosphere (Brownlee 1985). These interplanetary dust particles (IDPs) are divided in three classes all of which have broadly chondritic composition. Two of the classes are anhydrous with a mineralogy dominated by pyroxene and olivine; the third class has hydrous silicates similar to terrestrial smectites. Pyroxene- and olivine-dominated IDPs appear to come from comets because of their fragile microstructures, high carbon abundance, high content of Mg-rich silicates and inferred high atmospheric entrance speeds (Sandford and Bradley 1989). Hydrous IDPs are more likely to be from asteroids (Sandford and Bradley 1989; Bradley et al. 1999). GEMS (glass with embedded metal and sulphides) are major components of the chondritic porous anhydrous IDPs. GEMS are glassy spheroids, 0.1 to 0.5 μm in
diameter, with nanometre-sized opaque (metal and sulphide) inclusions embedded in nonstoichiometric Mg-silicate glass, and chondritic bulk compositions (Bradley et al., 1999). Their physical and chemical composition and inferred exposure ages lead to the belief that they have an interstellar origin (Bradley 1994). The spectrum of GEMS matches the spectrum of interstellar molecular cloud dust, young stellar objects, and the M-type supergiant. The isotopic compositions of some IDPs rich in GEMS link them to a molecular cloud environment (Bradley et al. 1999). Visible reflectance spectroscopy has been applied to IDPs, in order to compare them with primitive meteorites and with astronomical data. The preliminary results show that chondritic IDPs are spectrally dark like carbonaceous chondrites; hydrated IDPs show reflectance characteristics similar to CM and CI meteorites and main belt C asteroids; some anhydrous chondritic porous IDPs have reflectance similar to outer P and D asteroids (Bradley et al. 1996).

Micrometeorites are an additional class of extra-terrestrial materials that provide information about the Solar System (Grady and Wright 2006; Genge et al. 2008) and are the most abundant sources of IDPs available for study (e.g. Genge, 1997 and references therein). Antarctic micrometeorites show a composition similar to that of CM (unmelted) and CR and CI (hydrous) carbonaceous chondrites and are relatively free of terrestrial contamination, although subject to atmospheric entry heating alteration (Genge and Grady 1998; Genge et al. 2008). Micrometeorites can be more representative of a wider range of parent bodies than meteorites because of the dynamic associated with their orbits (Genge, 1997) but, also because of their small size, might not be representative of the whole parent body (Genge et al. 2008).
1.5 Asteroids

Asteroids are thought to be the parent bodies of most of the meteorites that fall on Earth. Most asteroids occupy the main asteroid belt, the area between the orbits of Mars and Jupiter (2-4 AU from the Sun), and it is widely accepted in the scientific community that they are remnants of many small planetary bodies which, unlike the planets, were never able to accrete into a single body. The cause for this inability to form a planet was the strong gravitational influence of the newly formed gas giant, Jupiter, which perturbed these small bodies, causing too many collisions for them to be able to complete the accretion process to form a planet (e.g. O'Brien and Sykes 2011). Asteroids are still affected by this gravitational force: there are gaps in the asteroid belt - Kirkwood gaps - which coincide with orbits in resonance to Jupiter's orbit. If a body occupies an orbit in resonance with Jupiter it will be accelerated or decelerated and, eventually, its orbit is altered sufficiently, removing the body from that area. Mars can also have an effect on the asteroids of the inner asteroid belt. Together with Jupiter, it changes the orbit of some asteroids throwing them in the direction of the Sun. These asteroids may acquire an orbit which is close to or even crosses the orbit of the Earth, making them potentially hazardous. These asteroids are known as Near Earth Objects (NEO).

Several missions have had ‘close encounters’ with asteroids. The first was Galileo which flew by asteroid 951 Gaspra, obtaining the first high resolution image of an asteroid (Figure 1.7) and 243 Ida (Belton et al. 1992).
Other missions are shown in Table 1.1. The Hayabusa mission (JAXA) landed on comet 25143 Itokawa, collected samples and returned them to Earth (e.g. Tsuchiyama et al. 2011).

Table 1.1: Asteroid missions, asteroid target and types

<table>
<thead>
<tr>
<th>Mission Name</th>
<th>Encounter Date</th>
<th>Target Asteroid</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>Galileo</td>
<td>Oct 1991</td>
<td>951 Gaspra</td>
<td>Flyby</td>
</tr>
<tr>
<td></td>
<td>Aug 1993</td>
<td>243 Ida</td>
<td></td>
</tr>
<tr>
<td>NEAR</td>
<td>Jun 1997</td>
<td>253 Mathilde</td>
<td>Flyby</td>
</tr>
<tr>
<td></td>
<td>Feb 2000 – Feb 2001</td>
<td>433 Eros</td>
<td>Orbitr</td>
</tr>
<tr>
<td>Deep Space 1</td>
<td>Jul 2001</td>
<td>9969 Braille</td>
<td>Flyby</td>
</tr>
<tr>
<td>Hayabusa</td>
<td>Sept 2005</td>
<td>25143 Itokawa</td>
<td>Orbiter-lander</td>
</tr>
<tr>
<td>Rosetta</td>
<td>Sept 2008</td>
<td>2867 Steins</td>
<td>Flyby</td>
</tr>
<tr>
<td></td>
<td>Jul 2010</td>
<td>21 Lutetia</td>
<td></td>
</tr>
<tr>
<td>Dawn</td>
<td>Jun 2011</td>
<td>4 Vesta</td>
<td>Orbitr</td>
</tr>
<tr>
<td></td>
<td>Feb 2015 (planned)</td>
<td>1 Ceres</td>
<td></td>
</tr>
</tbody>
</table>

Asteroids were formed at the same time as the Sun, planets and comets. Being relatively small bodies (1 Ceres, the largest, is \( \sim 1000 \) km in diameter), they have not undergone much alteration since the formation of the Solar System (with the exceptions described in the following section), and many are considered to have a primitive composition. Asteroids could be considered the counterpart of comets in
the inner Solar System with regards to the importance they have for the study of the
processes of its formation. Asteroids are also studied for economic reasons - for
future resources, and, on another level, for protection in the event of a collision with
the Earth (Burbine et al. 2002).

1.5.1 Classification of Asteroids

Information about the composition of the asteroids derives from spectroscopic
measurements, both ground and space based, mainly in the ultraviolet, visible and
near-infrared wavelengths (UV/Vis/NIR/MIR). Their classification into different types
is based on albedo, depth of absorption bands and slope of the spectral continuum
(Clark et al. 2002). Asteroids were first classified according to their visible
wavelength reflectance spectra (Chapman et al. 1975) (see example in Figure 1.8),
and there are now several different taxonomic schemes followed (see examples in
Figure 1.10). Three characteristics are essential for this classification: the first one
discriminates the presence of a UV absorption feature caused by Fe$^{2+}$ charge
transfer (see next chapter); the second characteristic is the slope of the spectrum
above 550 nm – which depends on alteration products and the presence of Fe-Ni
metal; the third characteristic is the presence or absence of silicate absorption
features beyond 700 nm (Pieters and McFadden 1994). A short description of the
asteroidal classes of most relevance to this thesis work follows, detailing major
features in their spectra and possible meteorite associations. For a more exhaustive
description see Chapman et al. 1975; Barucci 1991; Pieters and McFadden 1994;
Burbine et al. 2002; Bus and Binzel 2002; Cellino et al. 2002; Gaffey et al. 2002 and
references therein. Figure 1.9 shows the distribution of the different types of
asteroids and their distances from the Sun.
Figure 1.8: Examples of the spectra and albedo of some asteroid types. 4 Vesta is a type V; 16 Psyche is a type M; 1 Ceres, 324 Bamberga and 2 Pallas are type C; 3 Juno and 230 Athamantis are type S. Image retrieved from http://www.observeasteroids.org/spectra.php. Data are from (Chapman et al. 1975).

Figure 6.7 The uneven distribution of various spectral classes of asteroids in the asteroid belt. E asteroids may be the source of enstatite chondrites (Table 6.1); S asteroids, ordinary chondrites and/or stony-iron meteorites; M asteroids, iron meteorites; and C asteroids, carbonaceous chondrites. D and P asteroids are dark, carbon-rich objects that may not be represented in meteorite collections. (Abstracted with permission from J. Gradie and E. Tedesco,“Compositional Structure of the Asteroid Belt,” Science 216, 1982, 1405–1407. Copyright 1982 American Association for the Advancement of Science)

Figure 1.9: Most common types of asteroids and their distribution according to distance from the Sun.
S-type related asteroids

S-type asteroids

Mainly present in the inner asteroid belt, S-type asteroids have a moderate to high albedo (0.10-0.22) and spectral features attributed to the presence of the silicate minerals olivine and pyroxene (weak absorption feature at 1 and 2 μm) (e.g. Burbine et al., 2002). Their spectra are similar to the spectra of ordinary chondrites (OC) but they are redder and the features are not as strong. This may be because of a slightly different mineralogy or it may be related to the effects of space weathering (Wetherill and Chapman 1988). These effects will be considered further in Section 1.5.2 and in Chapter 7. The asteroids of this class have two minor features at 600 and 670 nm associated with the spinel group and oxidised Fe-Ni metal (Hiroi et al., 1996). They have a steep slope below 700 nm. S-type asteroids have recently been divided into subsets with slightly different spectral characteristics (Gaffey et al. 1993 and references therein). Of these, only sub-type S (IV) seems to have similarities to the spectra of OC (Burbine et al., 2002 and references therein).

K-type asteroids

K-type asteroids are included in S-types on the Tholen classification. They have spectra with characteristics between S and C-types. They appear to be composed of mainly olivine and pyroxene, with carbonaceous matter and are compatible with CV and CO meteorites.

Q-type asteroids

Q-type asteroids are quite uncommon. Their spectra are somewhere between S and V-type asteroids. They have a strong absorption feature at 1 μm indicating the presence of olivine and pyroxene and features just below and above 1.7 μm. The slope of their spectra seems to indicate the presence of metal. There is a very strong similarity to the spectra of ordinary chondrites. These asteroids are common
among Near Earth Objects (NEOs); they are much smaller bodies and their surfaces are interpreted to be younger in terms of exposure to space weathering and, therefore, are not as altered as the asteroids in the main belt. This explains their stronger similarity with OC than with S and V-types.

**C-type related asteroids**

**C-type asteroids**

C-type asteroids populate most of the outer asteroid belt. They are thought to be composed of carbon/organic/opaque rich rocks with hydrated silicates. C-types have low visible albedo (0.03-0.10) and an essentially flat, albeit slightly reddish, spectrum. There are some weak features between 600 and 900 nm from the alteration of Fe oxides and a strong drop-off below 400 nm. Some asteroids of this type show a water feature at 3 μm, leading to the belief that they have been the subject of thermal aqueous metamorphism. A broad absorption at 700 nm, which is also present in carbonaceous chondrites, is related to the presence of Fe$^{3+}$ formed by alteration of phyllosilicates. These similarities indicate that C-Type asteroids are the parent bodies of carbonaceous chondrites. Some weakening of the features in the asteroids spectra may be caused by space weathering (see e.g. Burbine et al, 2002, section 1.5.2 in this chapter and Chapter 7).

**B-type asteroids**

The spectra of B-type asteroids are very similar to the spectra of C-types. The differences between these two types reside on a slightly higher albedo, bluish spectra and absence of a strong drop-off in the UV wavelengths on the B-type asteroids. They are also mainly found in the outer asteroid belt. B-types appear to be composed of hydrated silicates, carbon-rich/opaque/organic rich minerals with a high degree of metamorphism. Their spectra characteristics indicate that they could be the parent bodies of altered carbonaceous chondrites.
G-type asteroids

G-type asteroids have also quite similar spectra to C-types. They have a strong UV feature below 500 nm and some spectra show a feature at 700 nm, which can be caused by the presence of phyllosilicates such as clays and micas. Similarly to the previous 3 types, they might also be composed of hydrated silicates and carbon-rich/organic/opaque matter, which makes them suitable candidates for thermally metamorphosed, aqueous altered carbonaceous chondrites.

P-type asteroids

The letter P comes from Pseudo M-type because both these types have very similar spectra, albeit with quite different albedos. P-types have very low albedo (0.1) and featureless, reddish spectra. With the exception of these last two characteristics, their spectra are also similar to S-types and they are believed to be composed of silica/organic/carbon-rich and hydrated materials. They are found essentially in the outer asteroid belt seeming to form the tail of C-types.

D-type asteroids

D-type asteroids have dark, featureless and reddish spectra. Their spectra are similar to the spectra of S-types but they lack the strong drop-off in the UV wavelengths. The orbits of D-type asteroids are located further away from the Sun and most of them are in fact Trojans. Trojans are asteroids that share the same orbit as Jupiter. They appear to be composed of silicates and organic/carbon-rich material and possibly clays. There are some theories that they might have an icy interior. They have no apparent link to any type of known meteorite but they might be the parent body of some IDPs.

V-type asteroids

Vesta is the most referred to asteroid of the V-type. There are several smaller asteroids on the same 'orbit' as Vesta (vestoids) which seem to have a similar
composition. They have a moderate to high albedo and a very strong feature just above 700 nm. The spectra are similar to S-type indicating the presence of olivine, pyroxene and feldspars. Vesta appears to be the only relatively large body with an almost intact basaltic surface. Its spectra have been linked to howardite, eucrite and diogenite meteorites (HED). Eucrites seem to originate from the upper layers of the asteroid and diogenites from the interior. Howardites look like a mixture of both maybe a result of some impact/geological event. Vesta’s major feature is at 506 nm, consistent with the presence of augite.

1.5.2 Difficulties in correlation between meteorites and asteroids

There are several factors that make the characterisation of asteroids through their spectra a difficult task. The phase angle of the observation (angle between the light source-Sun and the detector-telescope), space weathering, particle size and even temperature change the spectra in different ways. Differences in mineralogy within the same body, such as a slightly more metallic region caused by a collision may also cause reddening and darkening of the spectra. The abundance of spectrally-dark minerals on a meteorite or asteroid may produce a featureless spectrum, even though there are minerals present with strong absorption bands (e.g. Pieters et al. 1993; Noble et al. 2001). This is true especially for carbonaceous chondrites in which, even though the mineral olivine is predominant, the presence of opaque minerals that are very fine grained, dispersed and very effective absorbers, transforms the spectra of these meteorites into almost featureless spectra (Clark 1983). The effects of opaque phases are a darkening and reddening of the spectra and also a change in position of the silicate absorption feature at 1 µm towards shorter wavelengths (Gaffey et al. 2002).
Space weathering is one of the major factors thought to be the cause of discrepancies between the reflectance spectra of meteorites and asteroids. Space weathering of asteroids is discussed in greater detail in Chapter 7.

Other than telescope observations and space missions, meteorites are the main source of information we have about asteroids. It is, however, essential to identify the meteorite’s parent body. It is thought that, in general, different groups of meteorites derive from different parent bodies, with the exception of some cases where the same parent body seemed to be the origin of different meteorite groups. But the meteorite collection available is greatly reduced, compared to the number of asteroids, and this collection may be biased. Examples of more fragile compositions may not be common because they are more easily destroyed, and more resistant compositions may be more common even though they may not be predominant in the asteroid belt (Burbine et al. 2002). Furthermore, the dynamics of their orbits might prevent some populations from reaching the Earth.

The spectra of asteroids can be compared to the spectra of meteorites taken in laboratories. Gaffey (1976) showed that different meteorite types appear to produce different spectra in Vis and NIR. Features and slopes are dependent on mineralogy, particle size and temperature. Examples of comparison of spectra of meteorites and asteroids can be seen in Figure 1.10.
Figure 1.10: Examples of asteroid spectra and their similarities to the spectra of some meteorites indicating a possible link. a) John L. Remo, "Classifying and Modeling NEO Material Properties and Interactions" in Hazards, p. 561. b) Comparison of reflectance spectra of the Tagish Lake meteorite with individual reflectance spectra of the T-, P-, and D-type asteroids. Each of the reflectance spectra of the asteroids (open squares) and that of the Tagish Lake meteorite (solid line) are scaled for the best fit with each other for spectral shape comparison.

1.6 Summary

In this Chapter, the basic features of small bodies in the Solar System were outlined. Much of this information has been acquired through remote observations by ground- and space-based telescopes and by space missions. The purpose of this thesis is to describe the work carried out on meteorites and cometary dust to
complement the remote observations. UV-Vis reflectance spectroscopy is used because it is widely used in remote sensing of distant objects (as seen above) and classification of asteroids. Furthermore, the instrument used in this work, the microspectrophotometer (MSP) is an instrument of high definition, non-destructive and quick (see Chapter 3 for more information on the instrument).

We know from results from the recent Dawn mission that the surface of Vesta is heterogeneous. By analysing a variety of HED meteorites, possibly including some with anomalous composition, I hope to identify which types of meteorites are represented on Vesta's surface. This, in turn, will yield an idea of the extent of overlay by impact bombardment, projecting fresh material from deeper down onto the surface of Vesta. The ultimate goal will be to help understand the formation and processes occurring in differentiated asteroids.

Despite the fact that ordinary chondrites are the most abundant in the collection of meteorites falling onto the Earth’s surface, no specific asteroid class has been identified as the source of OC, although S type asteroids come closest. It has been suggested that space weathering, which produces a regolith on the surface of the asteroids, is responsible for modifying the OC signal. This hypothesis has been tested by laser irradiation of OC and measuring of the resulting spectra (Yamada et al. 1999 and Sasaki et al. 2001). I will take a different approach and analyse the effect of impacts by shock in the spectra of basalts. If effective, this technique can be applied to OC and possibly establish a better link between OC and S-type asteroids.

In summary, the specific goals of the research project were to:

• Determine whether the technique used in this work, UV-Vis microspectrophotometry, could be applied to the study of terrestrial and extra-terrestrial samples
• Produce a database of spectra from minerals most commonly found in meteorites and cometary materials; compare these data with data collected using other instruments (specifically NASA's RELAB Facility at Brown University, which provides data for most Solar System missions)

• Perform the first UV-Vis analysis of cometary grains and compare with results by other authors and using other techniques

• Determine the UV-Vis spectral characteristics of meteorites from Vesta, and compare with results from other instrument systems (see below).

• Simulate space weathering of asteroids by impact processes, and record any changes in the spectrum of the samples. Assess whether results from the simulations can be applied to assist in interpretation of the effects of space weathering on asteroids, and how this might influence their links to meteorites.

In Chapter 2, some of the background theory behind spectroscopy is given. In Chapter 3 the MSP will be described together with a series of tests performed in order to better understand the capabilities and limitations of this technique.

In Chapter 4, the technique will be applied to the study of two grains from the comet Wild 2, collected by the Stardust spacecraft. This chapter will also include some analysis of the limitations of the technique on certain samples. Chapter 5 will describe and compare the spectra of HED meteorites with data from other techniques and results will be compared to the spectrum of the asteroid Vesta. Chapter 6 is a tentative approach to the effects of space weathering on the parent bodies of meteorites. Samples were impacted at hypervelocities and their UV-IR spectra were acquired. The final chapter brings together all the conclusions derived in each chapter and suggests further work.
2 Spectroscopy

2.1 Introduction

This work focuses on the study of solids using ultraviolet (UV), visible (Vis) and near infrared (NIR) techniques. In this chapter a summary of the theory behind each of the techniques will be given as well as a brief description of the instruments used. For more detailed explanations of spectroscopy theory, see, e.g. Hapke 1993; Clark 1995; Beran and Libowitzky 2004.

Spectroscopy studies the effects of the interaction between radiation and matter. Electromagnetic radiation is the propagation of energy as a wave. It consists of oscillating magnetic and electrical fields which are perpendicular to each other and perpendicular to the direction of propagation. Electromagnetic radiation (EMR) is classified according to its wavelength from gamma-rays to long radio waves. Different atomic or molecular processes are associated with each wavelength region (Figure 2.1). The wavelength ($\lambda$) of the radiation is coupled to its frequency ($f$ or $\nu$) by the relation:

$$c = f \lambda$$  
Equation 2.1

where $c$ is the speed of light in a vacuum ($3 \times 10^8$ ms$^{-1}$).

EMR also has particle-like characteristics, referred to as photons, which are transporters of energy. The energy per photon can be related to the frequency via the Planck–Einstein equation:

$$E = hf$$  
Equation 2.2

where $h$ is Planck’s constant and $f$ is frequency.
Combining equations 2.1 and 2.2:

\[ E = \frac{ch}{\lambda} \]

Equation 2.3 relates the energy of a photon to the wavelength of radiation. The incidence of electromagnetic radiation, or photons, into or onto matter causes changes in energy at the atomic and molecular level. Objects can emit, absorb, or reflect EMR, and the different energy transfer mechanisms result in different effects. For example:

**Rotational Spectroscopy** – Molecules with an electric dipole moment (i.e., molecules made from charged particles e.g., H\(^+\)Cl) change their orientation by rotating when exposed to electromagnetic radiation (Figure 2.1). The ability for the molecule to rotate in a solid depends on their shape and type of bonds. Rotation bands occur in the far infra-red and microwave regions of the spectrum, are beyond the scope of this study and, for that reason, will not be discussed further.

**Vibration Spectroscopy** – Molecules with an electric dipole can change their configuration by vibration around their equilibrium position in a crystal lattice (Figure 2.1). Absorption of electromagnetic radiation interferes with the frequency of lattice vibration, causing the lattice to jump from one vibrational state to one of higher energy. It can then return to a lower energy state, releasing photons and heat (Hapke 1993). Vibration bands occur in the infrared region of the spectrum. Specific structural or functional groups within a molecule give rise to features at characteristic wavelengths, and so vibrational (or infrared, IR) spectroscopy is a valuable method for the identification of materials. Vibrational mechanisms are the basis of IR spectroscopy and will be discussed further in Section 2.3.

**Electron transitions** – Absorption or emission of electromagnetic radiation can change a previously stable system. Photons that are introduced into the system
may alter the configuration of electron shells or orbitals\(^1\), causing electrons to experience transitions between energy states and change the energy level of the system (Figure 2.1). If an electron travels from a lower energy orbital to a higher energy one, then the system has absorbed energy and the electron is said to be in an excited state. If an electron moves from a higher energy shell to a lower energy one, the system emits energy. This is the mechanism that produces absorption in the NUV, Vis and NIR and will be described in Section 2.2.

\(^{1}\)Very simply, atoms can be regarded as a system in which electrons orbit a central nucleus comprised of protons and neutrons. The electrons have a higher probability of being found in specific regions around the nucleus; these are known as shells or orbitals, have different shapes and energies, and each may be occupied by up to two electrons. The innermost orbital is the single s orbital. Next are the three p orbitals, then five d orbitals and seven f orbitals. The letters s, p, d and f stand for sharp, principal, diffuse and fundamental, from the characteristics of spectral lines. Electrons in the outermost orbital are the bonding electrons.
2.2 UV-Vis Spectroscopy

Near UV-Vis spectroscopy concerns the excitation of valence electrons within an atom, where transitions between energy levels result in absorption (or emission) of energy at wavelengths below ~ 1 \( \mu \text{m} \). The electron transitions taking place are from movement of electrons between orbitals. In inorganic compounds, the subject of this work, electron transitions are dominated by one of two effects: crystal field effects and charge transfer effects. These transitions occur between (or within) atoms or ions.

Crystal field effects result from changes in the distribution of electrons in the outer shells of an individual ion (especially the transition metals, where electrons are
arranged in the $d$ orbitals). Co-ordinating anions (or ligands) create an electric field about a central transition metal ion - crystal field - with a specific symmetry and shape, depending on the number of anions, their distance from the central cation and their electronic charge. The interaction between anions and central cation force the outer $d$ and $f$ orbitals of the cation to split into different energy levels, allowing electron transitions between different orbitals to occur within the cation. These transitions are dependent on the cation, so each element will result in a different spectrum. Crystal field transitions are more common in minerals with partially filled $3d$ orbitals. Each absorption band corresponds to the transition of an electron from the ground state to one of the possible excited states. The transitions occur in the visible to NIR region of the spectrum, and are the cause of the characteristic colours of transition metal compounds and minerals. There are several rules for these transitions to occur. The spin-multiplicity selection rule states that, in a free ion, there cannot be a change in the number of unpaired electrons, i.e., A change in the spin of the electron. These Spin-forbidden transitions have very low probability of occurring and have very weak features. The Laporte selection rule states that transitions within the same quantum shell (between the same type of orbitals) for example between $d$ and another $d$ which involves a change in configuration of the orbital are strictly forbidden. If there is a lack of a centre of symmetry (such as in tetrahedral coordination), however, these transitions may occur or, in case of octahedral coordination, there are moments when molecular vibration causes the metal ion to leave its symmetrical central position e.g. (Burns 1970).

The main Charge Transfer effect gives rise to features in the visible part of the spectrum, and is known as the Inter-Valence Charge Transfer effect (IVCT). It occurs between neighbouring ions with different oxidation states in a lattice (Gaffey 1976). These transitions occur in minerals when valence electrons transfer back and forth between adjacent ions - the electrons are in shared molecular orbitals.
E.g., in \( \text{Fe}^{2+} \rightarrow \text{Fe}^{3+} \) an electron is transferred from \( \text{Fe}^{2+} \) to \( \text{Fe}^{3+} \) (they switch). The same also happens between \( \text{Fe}^{2+} \) and \( \text{Ti}^{4+} \). The IVCT effect is much stronger than crystal field effects and is particularly significant in mineral species, where the 'cages' of \( \text{SiO}_4^{4-} \) are held together by arrays of metal ions (\( \text{Mg}^{2+}; \text{Ca}^{2+} \), etc.). These ions are present in mafic minerals such as olivine, pyroxene and Fe phyllosilicates. Different mineral compositions will have a different 'signature' shown by the depth, width and wavelength position of the absorption bands created by the charge transfer effect (Gaffey et al. 2002). An example of a very characteristic feature is the strong \( \text{Fe}^{2+} \) band at 260 nm, which is shown as a significant drop-off in the spectrum (e.g. Loeffler and Burns 1974). If \( \text{Ti}^{4+} \) is present in a mineral (e.g., in the pyroxene augite) it interacts with \( \text{Fe}^{2+} \). An electron in \( \text{Fe}^{2+} \) becomes excited into a higher state, changing \( \text{Fe}^{2+} \) to \( \text{Fe}^{3+} \) and \( \text{Ti}^{4+} \) as \( \text{Ti}^{3+} \). This is seen in a charge transfer band at 340 and at 430 nm (Cohen 1972; Hapke 2001b). The 260 nm drop-off in anorthite and the charge transfer effects in augite can be seen in Figure 2.2.

![Figure 2.2: Bidirectional reflectance spectra of minerals found in the lunar surface. The spectrum of anorthite show the 260 nm drop-off caused by \( \text{Fe}^{2+} \). The spectrum of augite shows the charge transfer bands at 340 and 430 nm caused by the presence of \( \text{Ti}^{4+} \). From Hapke (2001) using data from (Wagner et al. 1987)](image_url)
2.2.1 Other factors that influence the spectra of minerals at UV/Vis wavelengths

Two important definitions in spectroscopy are those of specular and diffuse reflectance. Specular reflection occurs when the light is reflected from a smooth surface. In this case the reflection has one direction only depending on the angle of incidence of the incident radiation. In diffuse reflection, the radiation is reflected back in various different directions. This happens when the surface of the material is not smooth.

Scattering (when the path of radiation is changed in any way by a change in medium) can be divided into surface scattering, in which light is reflected from the surface of a mineral, and volume scattering, in which light is absorbed to some extent by the mineral and then reflected back to the detector (Figure 2.3). The difference between reflected and scattered light as a function of wavelength is dependent on the composition and internal structure of the material and is used to derive information about its chemistry and internal structure.

Figure 2.3: Schematic showing specular reflection surface scattering and volume scattering. Incident rays (full lines in black) can be specularly reflected - surface scattering (dotted lines in red) or scattered into the particle - volume scattering (dashed lines in blue)
In opaque minerals at UV-Vis wavelengths, the spectrum is dominated by surface scattering, showing a reflectance maximum within a low reflectance spectrum. Volume scattering dominates in the NUV-NIR range. The relative contribution of surface and volume scattering to the final spectrum is not, however, proportional to the quantity of a mineral present. If an opaque mineral (e.g. carbon, with a featureless spectrum) is present, it can significantly diminish the features of a silicate, turning the spectra flat even with just a small quantity (Wagner et al. 1987).

The scattering process, or transmission into the medium, is dependent on the absorption coefficient of the medium according to the Beer-Lambert law:

$$ T = e^{-ta} $$  

Equation 2.4

where $T$ is the transmission of electromagnetic radiation, $\alpha$ is the absorption coefficient which varies with wavelength of the incident radiation and $t$ is the thickness of the medium.

**Particle size** also influences spectra, even within minerals of the same composition. In a large (or thick) grain, there is a greater internal path for the light to travel before it is reflected back, resulting in a spectrum with a lower overall reflectance. As particle grain size decreases, incident light experiences a greater number of internal reflections as it travels through the material, which decreases the magnitude of surface reflectance and increases volume reflectance (Aronson et al. 1966). Materials with low absorption coefficients become brighter as particle size decreases: incident radiation hits more surfaces, or interfaces, after having passed a shorter distance through the absorbing medium, and so less light will be absorbed before it is reflected back out of the medium. Several studies have been made on the effects of particle size on a spectrum including (Cloutis et al. 1986; Hapke 1993; Pieters et al. 1993; Pieters et al. 2000; Hapke 2001a; Hapke 2001b; Noble et al. 2001; Noble et al. 2007).
2.3 IR spectroscopy

Infrared spectroscopy results from the excitation of vibrational motions of atoms within a molecule. There are several ways a molecule can vibrate (vibrational modes) and this is related to the linearity of the molecules of the compound and the Cartesian coordinates it can move towards. Linear molecules have 3N-5 vibrational modes (3 are the Cartesian coordinates x, y and z) and non-linear molecules have 3N-6, where N is the number of atoms. The frequencies at which the vibrations occur are the fundamental frequencies (assigned as \( \nu_1, \nu_2, \) etc - e.g. water molecule in Figure 2.4.

When the vibration produces a dipole in the molecule, there will be an absorption band in the spectrum and the vibration is said to be active in the IR. If no dipole is produced, the vibration is inactive in the IR, and there is no absorption band. The frequency at which a vibration occurs is dependent on the strength of the bond and the mass of the atoms and is, therefore, specific to the molecule.
Figure 2.4: Water vibrational modes. The three normal (or fundamental) modes are: (a) the symmetrical stretch ($v_1$), (b) the symmetrical bend ($v_2$), and (c) the anti-symmetrical bend ($v_3$). Image from http://www.webexhibits.org/causesofcolor/5B.html (12/09/2012)

Overtones occur at multiples of the fundamental frequency: at $2v_1$, $3v_1$, $2v_2$, etc.; each overtone is usually 30 to 100 times weaker than the previous. Combinations ($v_1+v_2$, $v_2+v_3$, etc.) occur when two (or more) fundamental vibrations are excited simultaneously. Vibrations of different modes can also occur at the same frequency, in which case they are known as degenerate.

Several other features and characteristics, which can be seen in Figure 2.5, can be identified in the IR in order to determine the composition of the sample:

The Christiansen Feature (CF) was first recognised as a mineral composition indicator by (Conel 1969). It is an emissivity maximum, or a reflectivity minimum (see Figure 2.5), resulting from similarity in refractive index between the sample and the surrounding material (Conel 1969; Cooper et al. 2002). The similarity between the refractive indices results in little backscatter and low absorption. This results in the IR radiation passing freely through the sample (Conel 1969; Cooper et al, 2002)
and reflectance values are at a minimum. Volume scattering dominates at wavelengths below the CF and surface scattering dominates above the CF (Salisbury et al. 1991; Hapke 1993). The wavelength at which the CF occurs (typically between 7.5 to 9 \( \mu \text{m} \) for silicates in air) may change with grain size, but it is fixed if the grain size is smaller than the wavelength. For rock samples, analysis is more complex than for a single mineral, because the various minerals in a rock have different CF, some of which may overlap (Conel 1969). If the CFs are close to each other they average, if they are separate they are shown as different features (Salisbury et al. 1991; Cooper et al. 2002 and references therein). The Reststrahlen Bands (see below) of minerals at a shorter wavelength may also mask the different CF (Cooper et al. 2002).

**Reststrahlen bands (RB)** are the fundamental molecular vibration bands located in the surface scattering dominated part of the spectrum, i.e., immediately above the CF. This results in high peaks in a reflectance spectrum. The RBs are much stronger than other absorption bands and can be easily seen in the spectra of silicates (Figure 2.5). Because silicates have such high absorption coefficients in the Reststrahlen bands they behave as if they were opaque, producing a mirror-like reflection (high reflectivity peaks). RB are affected by the particle size of the sample (Lyon 1964) and become quite weak in fine powder of some minerals. If the surface is roughened and the grain size diminished, multiple scattering starts occurring and the reflectance decreases – the Reststrahlen bands become weaker and weaker (Salisbury et al. 1991).

**Absorption bands** occur in the region between 4 and 7 \( \mu \text{m} \) where volume scattering prevails (see below). For this reason, the absorption bands are seen as troughs or minima. These are also characteristic of the composition of the sample and have the advantage of increasing in definition with decreasing grain size.
Transparency Features (TF) appear in the normally transparent part of the spectrum (between the Si-O fundamental vibration bands at around 4 and 8 \( \mu m \)). It is only at small grain sizes that particles become optically thin at this wavelength and where volume scattering begins to dominate. TF are generally present in particles with grain sizes < 75 \( \mu m \). The reason for this is that sizes of the particles that pass through this sieve size are predominantly much lower than 75 \( \mu m \) (<5 \( \mu m \)) and, which tend to be optically thin (Hunt and Logan, 1972; Salisbury et al., 1991).

In addition, (Salisbury and Wald 1992) have shown that larger particles are coated by finer particles, hence the finer particles influence the spectra to a greater extent. TFs are a broad peak in reflectance and. Like CF and RB, they change in wavelength with composition. For this reason they can be used as an identification tool, together with the CF, of finely powdered samples (Salisbury et al. 1991; Cooper et al. 2002).

![Figure 2.5: Example of a spectrum showing the different identification features. Spectrum of < 75 \( \mu m \) powder of olivine. Spectral features include: C, Christiansen feature; R residual Reststrahlen features; TF, transparency feature (from Salisbury, et al 1991)](image)

2.3.1 Other factors that influence the spectra of minerals in the IR wavelengths

There are several additional factors that influence the IR spectrum of a sample. The orientation of the crystals relative to the polarised beam changes the intensities of
the bands. If the orientation is parallel to the beam, the features are stronger. In order to obtain a representative spectrum, random crystal orientations and non-cleavage surfaces should be chosen. Different areas of solids have different absorbances because of the variation of shape, size and distribution of the particles – this is the mosaic effect. To minimize the mosaic effect, the particles should be smaller than the wavelength of the incident radiation. This will, however, cause the RB to decrease.

Particle size – Rocks and particles with a large grain size are dominated by surface scattering – higher spectral contrast in the RBs. As roughness increases and particle size decreases, there is multiple surface scattering (cavity effect) and volume scattering starts to dominate, resulting in shallower bands. At particle sizes < 5 μm, close to the wavelength of the incident light, the particles become optically thin even in the RB area, resulting in TF (see above). The CF of rocks also changes, becoming enhanced and shifting to longer wavelengths. The amount of shift depends on the type of rock (Cooper et al. 2002). Below the CF, where volume scattering dominates, bands behave in the opposite fashion. As particle size decreases, volume scattering increases, resulting in a greater spectral contrast.

Vincent and Hunt (1968) first described three different particle size effects, later also explained by Salisbury and Wald (1992):

Type 1: present in the RB of silicates, and is a reduction in spectral contrast with decreasing particle size. The strength of the RB in silicates however, are sufficiently high that they never completely disappear.

Type 2: present in calcite at 11.47 μm, where a switch between the influence of volume and surface scattering changes a shallow reflectance maximum into a minimum as particle size decreases.
Type 3: weaker reflectance features are also exhibited as troughs rather than maxima.

Surface scattering gives reflectance peaks from the very strong molecular vibration bands, and is dominant in particles down to 5 μm in size, where type 1 behaviour is shown. If the particles become optically thin, at intermediate particle sizes, then volume scattering will become more significant and type 2 behaviour dominates. If type 3 behaviour is shown, it implies that all particles are optically thin and that that volume scattering is dominant (Salisbury and Wald, 1992). When the volume scattering starts to dominate, the shape of the RB is changed and the wavelength shifts (Vincent and Hunt 1968).

Porosity is a complex function of grain shapes and sizes and it may be higher in samples made of finer grains, because electrostatic forces cause adhesion and the particles form bridges and branches with large cavities between them (Salisbury and Wald, 1992). These cavities will act as a ‘photon trap’, altering the spectrum of a sample by reducing the spectral contrast of the RB (Salisbury and Wald 1992). Smaller particles cling to larger ones influencing the spectrum as described above (TF description) strongly changing the shape and wavelength positions of the RB (Salisbury and Wald 1992). Even in a sample where larger particles are dominant, smaller grains stick to the surface of the larger grains, covering them and the resulting spectrum will strongly reflect the influence of the smaller particles as described above. In packed samples, the porosity of fine particulate samples is reduced, and they become optically thicker, hence reducing volume scattering. However, packing may orientate the grains which leads to other alterations in the spectrum explained above.

Temperature and pressure also interfere with the spectra: Low temperatures cause higher reflectances and low pressure changes the CF, shifting it ~ 0.5 μm to shorter wavelengths, and deletes TF.
2.4 The FTIR Spectrometer

For the measurement of NIR to MIR spectra of our samples, we used a Fourier Transform Spectrometer (FTS). The main components of the instrument are:

- A polychromatic (multiple wavelengths) globar source - a silicon carbide rod which emits radiation from ~4 to ~15 μm, approximate to a black body radiator.

- A potassium bromide (KBr) beam splitter

- One movable mirror and one fixed mirror

- The sample chamber

- DTGS detector – deuterated triglycine sulphate detector.

Figure 2.6: Scheme showing the light path through the Fourier Transform Spectrometer
Figure 2.6 shows the light path through the spectrometer. A parallel beam of monochromatic radiation is emitted by the source hits the beam splitter, half is reflected to the fixed mirror and half to the movable mirror. They are both reflected back from the mirrors to the beam splitter where they are again divided and ideally 50% will reach the sample and 50% the detector. If the distances between the beam splitter and the mirrors (retardation) are the same or differ by an integral number of wavelengths, the beams leaving the beam splitter are in phase and interfere constructively, creating a very strong signal (peaks in Figure 2.7). As the difference in distance between the mirrors increases, different wavelengths will produce peaks at different positions and the signal diminishes (Figure 2.7), cancelling it if the difference is half integer wavelength number. As the movable mirror is moved away or closer to the fixed mirror, the detector sees radiation alternating in intensity. Fig 2.6 shows the interferogram that is produced by the detector with alternating intensities dependent on the distances between mirrors. If the source is polychromatic, however, as it is in this case, the interference pattern of the different frequencies of emitted colours will overlay and the beam will see a more complicated intensity fluctuation. A Fourier Transformation is applied by the computer software and it becomes possible to calculate the original frequencies of the light emitted by the source and remove it from the interferogram. Different 'sets' of wavelengths interfere differently (destructively or constructively) at each position of the mirror and at each position a data point is taken. The result is a sum of data points, each with information on how light at all wavelengths behaved at that mirror configuration. The spectrum of a sample is dependent on how that sample absorbs or emits light which is, in turn, dependent on the composition of the sample. The sample is positioned in the sample chamber, just before the detector. After the light passes through the sample, when the Fourier transformation is applied, the result is a normal absorption spectrum: by moving the fixed mirror over a period of time and
applying the transformation to detector signal at small intervals resulting in a spectrum of the sample at all frequencies.

![Graph of interferogram](image)

Figure 2.7: Interferogram showing the differences in signal with the distances between mirrors

The application of IR spectroscopy to minerals and meteorites will be discussed further in Chapter 6.
3 Microspectrophotometer operation

The instrument used for this work was an optical microspectrophotometer system supplied by Craic Technologies (San Dimas, Ca). The system is based on a Leica DMR microscope, with the regular objective lenses removed and fitted with Cassegrain reflecting mirrors (Figure 3.1). The light source is a 75W xenon lamp. The square aperture through which light was focussed varied in size with selected magnification, from 10 x 10 μm to 2 x 2 μm. The microspectrophotometer combines a microscope and a UV/Vis range (200-800 nm wavelength) spectrometer allowing viewing and magnification of a sample at very small scales (the samples can be as small as 2 μm) and measuring of the intensity of light in the UV/Vis region. This allows the distribution of organics or mineral types within small clumps of materials to be determined simply by selecting the specific area from which to gather the spectral data. This technique is non-destructive and quick as it needs almost no sample preparation. The microspectrophotometer also has a spectral resolution of > 2 nm (the Full Width at Half Maximum of the smallest resolvable spectral feature. This is the ideal spectrophotometer resolution but other factors in the microscope, sample and even the software can affect this number). The instrument also allows measurement of spectra by transmittance and reflectance in various types of samples. For this work we used only reflectance spectroscopy.

In reflectance mode, the light is focused onto the sample by the objective. The light reflected by the sample is then collected by the objective and the magnified image is projected into the spectrometer aperture (Figure 3.1). There is also a camera fitted which allows us to take photographs of the samples.
Inside the spectrometer (Figure 3.2) the light is reflected by a concave mirror (1) which then focuses the light onto the diffraction grating (2). The diffraction grating or monochromator separates the light into different wavelengths. The different wavelengths are then reflected by another concave mirror (3) onto the CCD (Charged Couple Device) (4) so that they can be read individually. This CCD array is positioned at the focal plane of the monochromator so that the entire spectrum can be detected.
Because the light has been separated into different wavelengths the intensity of the separated wavelengths will be lower. Sometimes it is too low to be detected. CCDs are integrating detectors. This means that if the signal we are receiving is too low to be detected, we can increase the exposure time of the detector to the incoming light to improve the signal-to-noise. The CCD is a device which responds to incoming photons of light using light sensitive elements (pixels) and produces a voltage. When a photon hits the pixel it generates an electron-hole pair. This electron is called the photoelectron. This information is accumulated, transferred and converted into a digital signal that can be stored and displayed.

There are several corrections that must be applied to the measured spectrum in order to obtain a true spectrum. The spectrum must be corrected for **dark current** and the varying **sensitivity** of the detector at different wavelengths.

A **dark current** occurs when some signal is still generated on the detector even if there is no light. This dark current results from the production of electrons that originate not from the light, but from thermal electrons and this increases with temperature. The way to correct this effect is to take a **dark scan**, i.e. a background exposure with no light. This dark spectrum is then subtracted from the sample and
reference spectrum (explained below) so that the effects of temperature are not shown in the final spectrum.

The Sensitivity varies with wavelength because neither the detector nor the light source have a flat spectral response. This problem is solved with a reference scan. The reference scan reflects the source effects and the reference imperfections. The sample scan is divided by this reference scan which eliminates the variation in detector sensitivity. Therefore the final spectrum will be a relative spectrum: the spectrum of the sample relative to the reference material. The ratio will be different if the standard used is different, but the features of the sample spectrum will be there and will be related to the instrument sensitivity. The instrument sensitivity can still be a problem if the signal is too low. This problem will be further analysed and discussed in the following section.

In summary the final spectrum (corrected) C will be:

$$ C = \frac{S - D}{R - D} \quad \text{Equation 3.1} $$

where $S$ is the sample spectrum, $D$ is the dark scan and $R$ is the reference scan.

$C$ will eliminate the system spectral response to give a corrected spectrum of the sample relative to the reference, which is a standard material with a known spectrum.

The spectrum is wavelength calibrated by exposing the spectrograph to a standard emission light source whose spectral lines are at well determined wavelengths. The pixel values of the signal are compared to the lines of the known source and wavelengths are attributed.

The software (see Figure 3.3) supplied with the instrument is equipped with several tools which include a calibration check. This calibration is done every day at the beginning of the session, after the lamp has stabilized. It ensures that the
wavelength conversion is calibrated and that all parts are aligned correctly. For this calibration check Holmium Oxide and Fluorilon™ diffuse white reflectance standards are used.

3.1 Performance analysis of the instrument

The microspectrophotometer is a state-of-the-art instrument which makes taking spectra of samples very easy and quick. The final output is a spectrum that is corrected with reference and dark scans.

The software (Figure 3.3) works in the following way: the scan duration and number of counts are first defined. A dark scan and a reference scan (the reference we used is a NIST traceable Fluorilon™ white reflectance standard) are then taken, followed by the sample. The final result has the correction for dark scan subtraction and the ratio with the reference standard; however, dark scans and reference scans are saved separately in the specified folder, allowing data to be retrieved for any off-line processing.
In order to understand fully the performance of the microscope, and to characterise the accuracy of the dark and reference scans, some tests were made.

3.1.1 Characterisation of dark scan

The spectrometer measures the way light is reflected off a sample. This is caused by the energy transitions that occur between the atoms when they are hit by the light, i.e., transition of electrons between energy levels. If there is no light in the system, there should be no transition and therefore a featureless spectrum. However, even without a light source, some signal can be detected: the dark current (discussed previously).
Figure 3.4 shows a typical dark scan from this instrument. The spectrum has a noticeable gradient, as well as fine scaled structure. It is necessary to know if these features and this gradient are reproducible and can be quantified.

For this purpose, a series of dark scans were taken throughout the day. The lamp was switched on and left for 1.5 hours (time assumed it would take to stabilize the lamp; the supplier advised only half an hour; this issue is discussed later). One dark scan was taken every half hour so that not only the reproducibility of the features and the gradient were analysed but also changes in time (Table 3.1). The results were plotted in different ways to illustrate the results.
### Table 3.1: Times at which the series of dark scans were taken and description of the figures

<table>
<thead>
<tr>
<th>Dark Scan Number</th>
<th>Time of scan</th>
<th>Spectrum Number</th>
<th>Description</th>
<th>Spectrum Number</th>
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<td></td>
<td>1</td>
<td></td>
<td>1</td>
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<td>T₀+30m</td>
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<tr>
<td>3</td>
<td>T₀+1h00m</td>
<td>3</td>
<td></td>
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<td></td>
</tr>
<tr>
<td>5</td>
<td>T₀+2h00m</td>
<td>5</td>
<td>Respective dark scans normalised to the median of dark scan 1</td>
<td>5 Ratio of respective dark scans to dark scan 1</td>
<td>5 All the spectra are normalised to their own mean and then divided at each point by the mean of all the other normalised spectra at that same point.</td>
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<tr>
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<tr>
<td>7</td>
<td>T₀+3h00m</td>
<td>7</td>
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<td></td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>T₀+3h30m</td>
<td>8</td>
<td></td>
<td>8</td>
<td></td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>T₀+4h00m</td>
<td>9</td>
<td></td>
<td>9</td>
<td></td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>T₀+4h30m</td>
<td>10</td>
<td></td>
<td>10</td>
<td></td>
<td>10</td>
<td></td>
</tr>
</tbody>
</table>

Figure 3.5: Dark scans normalised to the median of the first scan taken

After the spectra were taken, they were normalised to the median of the first spectrum so that variations became easier to analyse (Table 3.1, Figure 3.5). It shows that the features are reproducible and so is the gradient; however there is a
shift in the scan over time. This shift reaches values of 1% which is quite a significant value. All the features, gradients and shifts are present when there is no light source so they represent the instability of the instrument itself.

The fine scale features result from the pixel to pixel sensitivity variation; the cause of the gradient is unknown. Since features and gradient are reproducible over the entire wavelength range, they are not considered to be problems; the shift in the value of the dark current must be caused by temperature stabilization. If the shift decreases with time, then the spectra must be taken after the instrument stabilizes. If not, a dark scan must be taken before every spectrum taken from the sample – whilst this is not difficult, it is time consuming. In order to verify if the shift decreases with time, the scans were plotted as a ratio of each scan to the last scan taken (Table 3.1, Figure 3.6).

Figure 3.6: Ratio of each dark scan to dark scan 10

Figure 3.6 shows that after 4 hours, the shift decreases to ~0.1%/half hour which is acceptable. This means that if we allow 4 hours for the lamp to stabilise we only need to take one dark scan and the error will be sufficiently low.
An alternative solution for this problem is to calculate the uncertainty associated with the shift in dark current during the time of the experiment. This would mean that any fine-scaled structures in a spectrum smaller than that error should be discarded as artefacts (Figure 3.7). To calculate that error several steps were made. The first step was to normalise each scan by dividing each one by its own mean. Then find the mean of the ten scans at each wavelength. The last step was to divide each of the previously normalised scan at each wavelength by the mean found in step two (Figure 3.7).

The purpose of these steps is to find how much each feature varies during the time of the experiment, and find the standard deviation (Figure 3.8) which will give us the uncertainty of the scans.

![Figure 3.7: Scans normalised to their own mean and divided by the mean of all scans at each wavelength](image)
The highest value for the standard deviation is $\sim 0.12\%$ at one place, but the typical values are $0.05\%$, which means that any feature smaller than this error should be ignored. However, this value is taken from data from the full time of the experiment including the spectra at the time at which the lamp is not stabilised. This will give us a highest error than waiting 4 hours for the lamp to stabilise. This value is negligible compared to other effects which will be discussed in the next sections.

For samples which require results with high precision, the best method is to wait for the lamp to stabilise. If this is not possible then the best method is to take a dark scan immediately before each of the samples. This way the drift will not be too significant.
3.1.2 Characterisation of reference scan and reference standard

As explained above, part of the calibration procedure is to take a spectrum of a reference material. This reference scan removes the variation in detector and source sensitivity.

The reference scan that is saved as a file by the automated software is not corrected for dark current (i.e., it does not have the dark scan subtracted). This calculation is not made until the time the sample spectrum is taken. In order to analyse any spectrum taken by using the reference scan on the software, the dark scan must be subtracted manually. In the plots presented in the rest of this chapter, all the spectra have the dark scan subtracted. The reference scan minus the dark scan also gives us an idea of the sensitivity of the CCD at each wavelength (Figure 3.9)

![Figure 3.9: Reference scan, dark scan and the difference between them giving us an idea of the sensitivity of the CCD](image)

The results show that for a typical integration time of ~2000 milliseconds, between 200-300nm the signal is very poor. Results over this wavelength range then, are
likely to be more subject to error. This problem may be overcome if a spectrum is acquired over a longer integration time. It will increase the number of counts and show more reliable results. However, this may result in saturation of the features at longer wavelengths, but the problem is overcome by acquiring a separate spectrum across a shorter wavelength range.

Another factor to take into consideration is the calibration of the white standard itself. The reference standard used in this work is the NIST traceable Fluorilon FW-99-01c Diffuse White Reflectance Standard which is suitable for wavelengths between 250 and 850 nm. The supplier provided the data for the calibration of this standard. The data provided refer to wavelength intervals of 25 nm so data were interpolated to 1nm intervals. The results show that, although the reflectance should be flat, there are some variations in response throughout the spectrum and a severe drop off in reflectance at wavelengths below ~350nm (Figure 3.10). This means that all the data that we obtain have to be corrected for this factor. All data presented from now onwards will be corrected for the white reference calibration factor. Together with the low signal of the instrument at these wavelengths, the uncertainty in calibration of any data under 250nm is too large for any confidence to be attached to the results.
Because the white reference standard has a relatively rough surface and the precision of the microscope is so high, we needed to ensure that a reference scan taken in different places on the reference would not be too different from each other and therefore interfere with the results. If that was not possible then a precision error should be estimated.

Another test that was needed, after the above experiments with the dark scan, was to analyse if the reference spectrum shifted with time, how much it did shift when a light source is included, and if after some time this shift decreases.

The first test to be done was to take a series of scans at different locations on the reference standard. In total 5 scans were made one after the other. One single dark scan was taken before this series.

Figure 3.11 shows the ratio of each scan with the first one taken (Table 3.2), so that the differences are made clearer.
Table 3.2: Information on figure 3.11 scans. Spectra was taken on different place on the reference and at consecutive times.

<table>
<thead>
<tr>
<th>Reference Scan Number</th>
<th>Time of scan</th>
<th>Fig. 3.11 Spectrum number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$T_0=1h30m$</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>$T_1=T_0+3m$</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>$T_2=T_0+7m$</td>
<td>3</td>
<td>ratio of respective reference scans to reference scan 1</td>
</tr>
<tr>
<td>4</td>
<td>$T_3=T_0+10m$</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>$T_4=T_0+13m$</td>
<td>5</td>
<td></td>
</tr>
</tbody>
</table>

Figure 3.11: Ratio of all the reference scans to reference scan 1. Spectra was taken on different place on the reference and at consecutive times.

We can see that there are some differences related to the spot from where the spectrum is taken; the surface of the standard is not uniform at the scale of the aperture of the instrument (10 $\mu m$ for a magnification of 15x). This means that, because the final spectrum is relative to the reference, the results will be different.
each time we measure them as going back to the exact same spot on the reference at different occasions will be almost impossible. These differences however are lower than 0.5% at wavelengths longer than 300 nm. Further tests may give us more information on the significance of this scatter and how to best correct these effects.

The second test to be made is related to the variations over time. A number of scans were made keeping the focus on the exact same location of the reference and taking spectra at different intervals of time (Figure 3.12, Table 3.3). One dark scan was taken before this set of experiments.

Table 3.3: Information on figure 3.12 scans. Scans were taken at the same location at the times referred

<table>
<thead>
<tr>
<th>Reference Scan Number</th>
<th>Time of scan</th>
<th>Spectrum number</th>
<th>Fig. 3.12</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>T₀=01h30m</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>T₁=T₀+5m</td>
<td>2</td>
<td></td>
<td>Ratio of respective reference scans to reference scan 5</td>
</tr>
<tr>
<td>3</td>
<td>T₂=T₀+15m</td>
<td>3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>T₃=T₀+35m</td>
<td>4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>T₄=T₀+75m</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Figure 3.12: Ratio of all the reference scans, taken at different times and the same location, to reference scan 5.

Figure 3.12 shows that there are systematic drifts of up to about 3%. These drifts, however, seem to decrease after about one hour after the start of the experiment. It is important to check if the drift stops or diminishes after a certain time, define that time and check if greater precision (<1%) can be achieved. At this point, the drift observed in the first hour is too significant for results to be useful. More tests need to be done at longer intervals of time to see if it is repeatable. The drift may be dependent on the stabilization time of the lamp or to external factors.

At this point there are two problems of accuracy presented to us: A drift of the spectra in time and the roughness of the surface of the reference standard. The following sections are dedicated to the tests made in order to overcome these problems.
3.1.3 Investigation of time shift

It is clear from previous experiments that there is a drift of the spectra in time. This drift, however, seemed to diminish after a certain amount of time which leads to believe that it is related to stabilization of the instrument.

It is necessary to find out how long it takes the system to stabilize enough so that the error is at least as low as 1% instead of the 3% shown until the 3rd hour.

A series of reference scans were taken every 30 minutes for a total of 4.5 hours, after the lamp has been on for 1.5 hours (Table 3.4). One dark scan was taken before each scan so that the drift also seen in the dark scan (Figure 3.5-3.8) is not a problem. The light was focused on the surface of the reference standard and the spectra were taken always in the same spot so that the roughness of the standard does not interfere with the results.

Table 3.4: Information on figure 3.13. Scans were taken always at the same location, at the times referred

<table>
<thead>
<tr>
<th>Reference Scan Number</th>
<th>Time of scan</th>
<th>Spectrum Number</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$T_0+1h30m$</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>$T_1=T_0+30m$</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>$T_2=T_0+1h00m$</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>$T_3=T_0+1h30m$</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>$T_4=T_0+2h00m$</td>
<td>5</td>
<td>Ratio of the respective reference scan to reference scan 10</td>
</tr>
<tr>
<td>6</td>
<td>$T_5=T_0+2h30m$</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>7</td>
<td>$T_6=T_0+3h00m$</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>$T_7=T_0+3h30m$</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>9</td>
<td>$T_8=T_0+4h00m$</td>
<td>9</td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>$T_9=T_0+4h30m$</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
In order to better analyse after how long and how much this drift start to be reduced the data was plotted as a ratio of all the spectra with the last one taken (after 6 hours from the moment the lamp was on) on Figure 3.13, Table 3.4.

The plot shows that the drift strongly decreases after 3.5 hours. Above 300 nm there is a drift of ~1% after 2 hours and that drift is reduced to ~0.2% after 3.5 hours similar to what happened with the dark scan (Figure 3.5-3.8).

3.1.4 Investigation of out-of-focus reference standard

A possible solution for the roughness of the surface of the Fluorilon White Reflectance Standard could be defocusing the beam of light so that a larger area of the standard was scanned. If proved, this could also be a system used to overcome the fact that the aperture of the spectrometer is so small. One of the major advantages of this Microspectrophotometer is that the spectra are taken from very small areas, from 2 to 10 nm. Although being excellent if the sample is very small and for single composition samples, this also means that when samples of rock are
analysed, each spectra taken will most likely be of a simple grain and not representative of the whole sample.

There may be two solutions for the problem of the roughness of the white reference standard: Put the instrument out of focus so that a large region of the surface is observed giving an average of the area, or take a series of in focus scan at various places, average them and calculate the standard deviation to find the error.

A small test was made in order to see if defocusing the light spot on the reference standard was viable and accurate enough. For this experiment one single dark scan was taken and then a series of scans at different times. The time lapses were the same as the previous experiments. The reference scans were made on one single place on the standard surface but defocused so that the area analysed was larger.

The results were plotted as a ratio of all the scans to the first one taken (Table 3.5).

Table 3.5: Information on figure 3.14 scans. Scans were taken with the reference out of focus

<table>
<thead>
<tr>
<th>Reference Scan Number</th>
<th>Time of scan</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>$T_0=01h30m$</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>$T_1=T_0+5m$</td>
<td>2</td>
</tr>
<tr>
<td>3</td>
<td>$T_2=T_0+15m$</td>
<td>3</td>
</tr>
<tr>
<td>4</td>
<td>$T_3=T_0+35m$</td>
<td>4</td>
</tr>
<tr>
<td>5</td>
<td>$T_4=T_0+75m$</td>
<td>5</td>
</tr>
</tbody>
</table>

Fig. 3.14: Description of scans.
It is obvious that a drift is still present. Even at wavelengths where the signal is quite high (Figure 3.9), the spectrum drifts about 0.1%. Any feature smaller than ~0.2% should not be trusted. They don’t seem to be larger than the previous experiments with focussed standard so it looks like an out of focus scan may be a solution for the roughness problem. However the results are not conclusive as the time drifting may disguise the results. Further tests must be made in order to prove the accuracy of using an out of focus beam.

To analyse the effects of defocusing the reference measurement, data were obtained using a rock sample calibrated with reference spectra, taken with a range of defocusing (Table 3.6, Figure 3.16, Figure 3.17).

The process of this experiment was to take a spectrum with the beam focussed on the sample and on the white standard and increasingly defocusing it by decreasing the distance of the objective to the sample in steps of known distances.

All these spectra were taken after the lamp has been on for 4 hours.
Table 3.6: Distance of objective from focused position and information on figure 3.15 scans taken on the white standard and figure 3.16 scans taken on a sample

<table>
<thead>
<tr>
<th>Reference Scan Number</th>
<th>Distance from focused position (µm)</th>
<th>Fig. 3.15 Description</th>
<th>Spectrum Number</th>
<th>Ratio of respective scan to the focused scan</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>4514</td>
<td>1</td>
<td>1</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>9028</td>
<td>2</td>
<td>2</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>13514</td>
<td>3</td>
<td>3</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>18056</td>
<td>4</td>
<td>4</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>22570</td>
<td>5</td>
<td>5</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>31598</td>
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<td>10</td>
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<td>11</td>
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<td>12</td>
<td>284382</td>
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<tr>
<td>13</td>
<td>464942</td>
<td>13</td>
<td>13</td>
<td></td>
</tr>
</tbody>
</table>

Figure 3.15: Ratio of all the out of focus scans to focussed scan, taken on white reflectance standard.
These results indicate that there seems to be an illumination issue; the less focused the sample is, the less light it has as a result of the same amount of light being spread over a larger area.

There is also a larger difference in the wavelength dependence as a function of focus. This might be explained by scattering. The surface of the sample is rougher and the scattering from the different grains may compensate the lack of light. The shape of the spectra can also be influenced by the size of the grains in comparison with the wavelength. All these effects mean that defocusing the light beam is not a solution to compensate for the roughness of the reference materials.

### 3.1.5 Conclusions

When using an instrument for the first time, it is important to evaluate all the characteristics, potential and limitations of that instrument. For that reason, a series of tests were conducted on the microspectrophotometer. These tests included the stability of the CCD, by testing the dark scan; the stability of the source and the sensitivity of the system at all wavelengths, by testing the reference scan; and the homogeneity of the diffuse white reflectance standard used. The results show that
there are a few limitations and that a certain method is necessary in order to obtain an acceptable degree of uncertainty. There are several steps that can be made and these depend on the level of uncertainty necessary for the work which will be discussed at the end of this section.

One of the problems encountered was the fact that the reference standard was not homogenous at the micron scale (aperture of the spectrometer). The solution found was to take a series of reference scans, in order to have a representation of the whole standard, average them, and find the uncertainty associated with it.

The same steps were made as in Figure 3.7 and Figure 3.8: each scan was normalised by dividing them by their own mean. Then the mean of the ten scans at each wavelength was found and each of the previously normalised scans were divided by this mean at each wavelength. This procedure was made on the three different magnifications (15x-aperture of 10 μm; 36x-aperture of 4 μm; 52x-aperture of 2 μm) of the microspectrophotometer (Figures 3.17-3.22).

![Figure 3.17: Scans normalised to their own mean and divided by the mean of all scans at each wavelength. The scans were taken on a 15x magnification - 10 μm](image-url)
Figure 3.18: Standard deviation of the normalised scans on figure 3.17

Figure 3.19: Scans normalised to their own mean and divided by the mean of all scans at each wavelength. The scans were taken on a 36x magnification - 4 μm
Figure 3.20: Standard deviation of the normalised scans on figure 3.19

Figure 3.21: Scans normalised to their own mean and divided by the mean of all scans at each wavelength. The scans were taken on a 52x magnification - 2 µm
Figure 3.22: Standard deviation of the normalised scans on figure 3.21

The plots show that the uncertainty increases with the magnification; however, this uncertainty is a fraction of the wavelength and always lower than 7%. This is the uncertainty if we only take one reference scan of the reference standard. The line in red on the plot is the uncertainty divided by $\sqrt{20}$. This is the uncertainty we take 20 reference scans and average them every day. This is less than 0.5% at a 15x magnification.

As a result, different steps can be taken according to the level of resolution and uncertainty necessary for the work in progress:

For a quick measurement with no need for a high resolution results, then the method is to just wait for $\sim1$ hour after the lamp is on, take one dark scan, one reference scan before each sample scan and let the system do the automatic corrections. This will give us an uncertainty of $\sim3-4\%$. 
If the accuracy of the results needs to be higher, there are several steps that must be taken:

- Wait 4 hours for the system to stabilise;
- Take one dark scan and one reference scan after the instrument is stabilised;
- Take the spectrum of the sample as if it were a reference so that all correction may be done manually;
- Do all the correction (subtract the dark scan and divide the reference scan) and include the white standard correction given by the supplier (Figure 3.10)

This will give us an uncertainty of ~2-3%

For samples which require very high accuracy, then at least 20 reference scans must be taken from the reference standard as shown above. This will lower the uncertainty to <1%.

3.2 Application to minerals

UV-VIS reflectance microspectrophotometry is a non-destructive and rapid technique which requires very small amounts of material. This makes it an invaluable tool for the study of meteorites and other fragile and rare samples. The UV-Vis microspectrophotometer (MSP) is, however, an instrument which has only recently been used in planetary sciences (Bradley et al. 1996) although it has extensive applications in ore mineralogy (Criddle and Stanley 1993). Because there is a lack of high (spatial) resolution reflectance data for silicate minerals, the technique was tested on various samples, to build up a spectral library. We first analysed single grains of different minerals, followed by analyses of meteorites and their components, to aid in the process of matching meteorites with potential parent bodies. As described in Chapter 1, asteroids are the parent bodies of most of the meteorites we have sampled and they are classified according to their visible
reflectance spectra. The spectra taken in laboratories of meteorites are compared with the spectra of asteroids in an attempt to find the parent body of each type of meteorite (e.g., Pieters and McFadden 1994; Binzel et al. 1996; Hiroi et al. 2001; Burbine et al. 2002; Pieters et al. 2005b; Hiroi et al. 2007; Hiroi et al. 2008 and references therein.) By extending the wavelength range down towards the UV, I hoped to find additional spectral features that could be matched between meteorites and asteroids. Further to this aim, there is also the case of the presence of organic matter within carbonaceous chondrites which may hold clues to the origin of life. To have a non-destructive technique which may, at a preliminary stage, identify the presence of specific minerals and organics in extra-terrestrial samples, as well as aid in the task of identifying meteorite parent bodies, using very small samples, was the reason for the study described in this section.

The minerals used in this study were chosen because of their abundance and/or significance in meteorites. In this chapter, only the results of two minerals are shown as example of the quality of the spectra obtained, analysis of the artefacts and errors that can be generated by the instruments and provide an overall assessment of its performance. Results from other minerals can be seen in Appendix 1.

As outlined in Chapter 2, UV-Vis spectra arise from the movement of electrons between energy levels in an atom, or between two separate atoms. Previous work (e.g., Burns, 1970, Gaffey, 1976, Hapke 1993) has shown that features in the UV-Vis spectra of minerals are caused by two main processes:

1) **Charge transfer**, which may be either the transfer of charge (electrons) between metal and oxygen in a crystal lattice (e.g., O²⁻ - Fe²⁺), or the transfer of charge between two metal ions in the lattice (e.g., Fe²⁺ - Ti⁴⁺). The wavelength at which the feature occurs is related to the site symmetry of the ions involved in the transition, and not necessarily to mineral species (olivine, pyroxene, etc). Table 3.7 lists the
wavelengths at which some of the most prominent features occur, and the metal-oxide charge transfer with which they are associated.

Table 3.7: Expected metal-O charge transfer absorption bands, after (Cloutis et al. 2008)

<table>
<thead>
<tr>
<th>Metal Site Symmetry</th>
<th>Wavelength (nm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe$^{3+}$ Octahedral</td>
<td>234, 280, 340, 395</td>
</tr>
<tr>
<td>Fe$^{3+}$ Non-octahedral</td>
<td>217</td>
</tr>
<tr>
<td>Fe$^{2+}$ Tetrahedral</td>
<td>343</td>
</tr>
<tr>
<td>Fe$^{2+}$ Octahedral</td>
<td>250, 267, 380</td>
</tr>
<tr>
<td>Ti$^{4+}$ Octahedral</td>
<td>259, 296</td>
</tr>
<tr>
<td>Ti$^{3+}$ Octahedral</td>
<td>216, 242</td>
</tr>
<tr>
<td>Cr$^{3+}$ Octahedral</td>
<td>169</td>
</tr>
<tr>
<td>V$^{5+}$ Octahedral</td>
<td>253, 303, 322</td>
</tr>
<tr>
<td>Co$^{3+}$ Octahedral</td>
<td>~ 330</td>
</tr>
<tr>
<td>Mn$^{3+}$ Octahedral</td>
<td>&lt;200</td>
</tr>
<tr>
<td>Ni$^{2+}$ Octahedral</td>
<td>&lt;200</td>
</tr>
</tbody>
</table>

(2) Crystal field transitions are the second process by which features in UV-Vis spectra are produced. These are transitions in which an electron moves between energy levels in an individual ion. The wavelengths at which features associated with such transitions occur are related to site symmetry and to mineral species. Table 3.8 lists some of the main absorbance features for both low (LCP) and high (HCP) calcium pyroxene.

Table 3.8: Expected absorption bands in pyroxenes (Cloutis 2002)

<table>
<thead>
<tr>
<th>Cation and Site Occupancy</th>
<th>Mechanism</th>
<th>Location of Associated Absorption Bands, µm</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ca$^{2+}$ in M2 octahedral site in LCP</td>
<td>Spin-forbidden crystal field transitions</td>
<td>0.425, 0.445, 0.480, 0.505, 0.545</td>
</tr>
<tr>
<td>Fe$^{2+}$ in M2 octahedral sites in LCP</td>
<td>Fe$^{2+}$ - Fe$^{3+}$ charge transfer</td>
<td>0.69</td>
</tr>
<tr>
<td>Fe$^{3+}$ in M2 octahedral site in LCP</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zr</td>
<td>Crystal field transitions</td>
<td>0.345, 0.360, 0.410, 0.540, 1.5</td>
</tr>
<tr>
<td>Li</td>
<td>Crystal field transitions</td>
<td>0.420, 0.440</td>
</tr>
<tr>
<td>Na</td>
<td>Crystal field transitions</td>
<td>0.445, 0.640</td>
</tr>
<tr>
<td>Ca</td>
<td>Crystal field transitions</td>
<td>0.46</td>
</tr>
<tr>
<td>Fe$^{2+}$ in M1 octahedral site in HCP</td>
<td>Crystal field transitions</td>
<td>0.445, 0.640</td>
</tr>
<tr>
<td>Ti$^{4+}$ in M1 octahedral site in HCP</td>
<td>Crystal field transitions</td>
<td>0.46</td>
</tr>
<tr>
<td>V$^{5+}$ in M1 octahedral site in HCP</td>
<td>Crystal field transitions</td>
<td>0.445, 0.640</td>
</tr>
<tr>
<td>Mn$^{2+}$ in M1 octahedral site in HCP</td>
<td>Crystal field transitions</td>
<td>0.46</td>
</tr>
<tr>
<td>Cr$^{3+}$ in M1 octahedral site in HCP</td>
<td>Crystal field transitions</td>
<td>0.345, 0.360, 0.410, 0.540, 1.5</td>
</tr>
</tbody>
</table>

*The latter three bands may only appear as an unresolved broad absorption feature.
Spin-forbidden transitions produce weaker features in spectra than either metal-oxide transfer or crystal field transition. Spin-forbidden transitions occur when an unpaired electron in one orbital pairs with an unpaired electron in a second orbital, changing its spin direction in the process. Such transitions are associated with ions in which all the outer electrons are unpaired, and so have the same spin, e.g., Fe$^{3+}$ and Mn$^{2+}$.

If not already provided in powder form, minerals were ground in an agate pestle and mortar. With the exception of the minerals used for grain size analysis, samples were not sieved and a variety of grain sizes were used. It is possible to observe the size of the grains relative to the aperture of the spectrometer in the images provided.

Further sample preparation was minimal: grains were simply placed on to a quartz slide, which was laid on the MSP stage. Analyses were taken with the 15x magnification. The reason why a quartz slide was used was to minimise effects of the medium surrounding the grain as those observed on the scans taken on Stardust samples which were pressed into gold foil (see Chapter 4 for more information). Unlike glass, quartz is transparent to UV wavelengths and is used in UV studies in transmission for this reason. The purpose was to use as little sample and sample preparation as possible. For all samples, the same number of counts (50) and Integration Time (1000 ms) was used in order to compare results between samples. Previous sections of this chapter contain a description of the MSP instrument and the data reduction procedures.

In some cases (where data were available) the results from the MSP were compared to those from the Reflectance Experiment LABoratory (RELAB) instrument at Brown University, and also with those from the spectroscopy laboratory at the University of Winnipeg (UNIWIN; Cloutis et al., 2008). The purpose is to infer if the MSP can be used in this field where data are often shared and
compared for validity. Further to this, the results from RELAB and UNIWIN are the ones used for comparison to asteroid spectra and their link to meteorites. If the results from this study are not comparable with RELAB or UNIWIN data, then the MSP might not be an appropriate instrument for the analysis of minerals and extraterrestrial samples.

It must be pointed out that the geometry and sample sizes used by both the RELAB and UNIWIN instruments are completely different from each other and from the OU MSP and, therefore, some differences in results are expected. The RELAB instrument is a bi-directional reflectance spectrometer for visible to near-infrared wavelengths (range of 0.3 to 2.6 $\mu$m) with a spot size > 1 mm (Pieters and Hiroi 2004), http://www.planetary.brown.edu/relabdocs/RelabManual2006a.pdf. At least 20 mg, with an optimum sample size of ~ 500 mg, of powdered sample is required, and each sample takes ~2 hours to run. The phase angle (angle of incidence of the source + angle of detector to the vertical) must be at least 10 degrees. The UNIWIN instrument Ocean optics S-2000 covers from 200 to 859 nm and the phase angle is 0 degrees http://psf.uwinnipeg.ca/Sample_Database.html. The sample size is not described by Cloutis et al (2008) but a reference to a sample holder 20 mm wide and 10 mm deep was used for the BaSO$_4$ standard to avoid light interactions with the sample holder. In contrast, the OU MSP can acquire a spectrum from 0.25 to 0.8 $\mu$m from a single grain < 20 $\mu$m across, in less than 2 minutes; the incident and reflected beams are normal to the sample. However, since the RELAB, UNIWIN and OU instruments are reflectance spectrometers operating across similar wavelength ranges, any features characteristic of the minerals should be present in spectra from both instruments at the same wavelengths, even if not with the same signal/noise ratio.
Rhodonite and apatite are used in this section as examples that show the capabilities and limitations of the instrument. A test of the influence of the grain size in the spectra of minerals is also shown.

### 3.2.1 Rhodonite

Rhodonite is a pyroxenoid, with Mn$^{2+}$ replacing Ca$^{2+}$ as the major cation. Fe$^{2+}$ and Zn$^{2+}$ may also substitute for Ca$^{2+}$ (Klein and Hurlbut Jr. 1993). Figure 3.23 shows images of five rhodonite grains for which spectra were acquired; the associated spectra, and the average of the five spectra, are in Figure 3.24. The composition of this sample can be found in Table 3.9.

Table 3.9: Composition of the rhodonite analysed. The composition analysis was performed using a Scanning Electron Microscope

<table>
<thead>
<tr>
<th></th>
<th>F  (wt%)</th>
<th>Mg (wt%)</th>
<th>Si (wt%)</th>
<th>Ca (wt%)</th>
<th>Mn (wt%)</th>
<th>Fe (wt%)</th>
<th>O  (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rhodonite</td>
<td>1.2</td>
<td>0.3</td>
<td>21.6</td>
<td>5.9</td>
<td>23.6</td>
<td>10.3</td>
<td>37</td>
</tr>
</tbody>
</table>
A characteristic of the spectra taken from our samples is a series of very weak features occurring at wavelengths above 450 nm, which we infer to be some type of analytical artefact. Cloutis et al. (2008) noted a similar effect in many of their spectra, where “spectral artifacts in the form of narrow (~ 3 nm wide) peaks and valleys appeared in the 200-212 nm and 390-400 nm regions” (Cloutis et al. 2008). They concluded: “Our interpretation of this is that the light path is effectively different for BaSO₄ versus a mirror or directly through air because some of the light will scatter within the BaSO₄ before reaching the spectrometer, thereby effectively
increasing its path length versus the standard mirror surface measurement." [Note: Cloutis et al. employed BaSO₄ as the standard against which their samples were compared, whereas a Spectralon standard was used in this study]. The specific wavelengths noted by Cloutis et al. were regions where the deuterium light source was known to have strong emission features. It is unlikely that completely the same explanation holds for the data acquired herein: we are using a xenon light source which has strong emission features around 475 nm and above 800 nm (see Figure 3.9), whilst the multiple narrow peaks and valleys occur in most spectra above ~450 nm.

Rhodonite was selected as a mineral to study because, whilst it is not a mineral known to be common in meteorites, it does possess a combination of transition metals (Mn, Fe and Zn) present in the M1 and M2 sites which should give a spectrum with additional features to those seen in pyroxenes. This seems to be the case. There is a very strong absorbance at around 300 nm, along with weaker, but still distinct, absorbances at ~410 and 550 nm, and a broad minimum around 700 nm (seen most clearly in rhodonite 2).
All five of the spectra are very similar in shape, but there are differences in brightness, redness and definition of the spectra which seems to be related to the shape and size of a sample. The spectrum of Rhodonite 1 is redder and with more features than the other spectra taken by the MSP. It is also the largest of the grains, and one that completely fills the field of view of the MSP. It was assumed that as long as the aperture of the MSP covered the sample, then we would be acquiring a representative spectrum of the material. Results from different grains indicate that this assumption may not be true, and that the entire field of view had to be filled with grains. Generally, the samples were mounted on a quartz slide, transparent to UV-Vis light. However, it is possible that the 'signal' from a small sample surrounded by what is, in effect, 'no signal', may be 'diluted', preventing the instrument from detecting a distinctive spectrum, or that the slide is causing specular reflection. This problem is considered again in Chapter 4, where it is clear that the area surrounding the sample can influence the final spectrum.
The data can be compared with results from a mineral with similar composition (Px044) analysed by both the UNIWIN and RELAB systems (Figure 3.25). The spectrum of Rhodonite 1 and Px044 show the same deep feature at 300 nm, possibly from charge transfer between Mn$^{3+}$ and oxygen (Langer and Abu-Eid 1977). The feature at ~ 410 nm is from Mn$^{3+}$ in octahedral coordination in the M1 site, whilst that at ~ 550 nm, seen in the RELAB spectrum but only very indistinctly in the UNIWIN spectrum, is possibly a spin-forbidden transition associated with Fe$^{2+}$ in the M1 octahedral site (Cloutis 2002).

The spectrum with the strongest similarity to the results from UNIWIN and RELAB is the one taken on a grain that fills the field of view which supports the statement that there might be some influence from stray light scattered into the instrument.

Figure 3.25: Spectrum of rhodonite 1 taken on the MSP, compared with results of the same mineral (Px044) taken by the RELAB and UNIWIN systems. Sample Px044 is 45-90 μm powder. UNIWIN and RELAB data from http://psf.uwinnipeg.ca/Sample_Database.html and http://www.planetary.brown.edu/relab/ respectively.
3.2.2 Apatite

Apatite is a group of phosphate minerals. Five separate grains of apatite were analysed; their images are shown in Figure 3.26 and their composition can be found in Table 3.10. The spectra of all five grains and of one of similar composition take by RELAB can be seen in Figure 3.27.

Table 3.10: Composition of the apatite analysed. The composition analysis was performed using a Scanning Electron Microscope.

<table>
<thead>
<tr>
<th></th>
<th>F (wt%)</th>
<th>P (wt%)</th>
<th>Al (wt%)</th>
<th>Ca (wt%)</th>
<th>O (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Apatite</td>
<td>8.6</td>
<td>16.6</td>
<td>3.0</td>
<td>38.2</td>
<td>36.7</td>
</tr>
</tbody>
</table>

Figure 3.26: Images taken by the MSP of five separate grains of apatite; associated spectra are given in Figure 3.27. (a) apatite 1, (b) apatite 2, (c) apatite 3, (d) apatite 4, (e) apatite 5. The black square in the middle is the aperture of the spectrometer and is 10 x 10 μm in size.
Figure 3.27: Spectra taken on the MSP of five different apatite grains compared with results from a similar mineral taken by RELAB. The RELAB sample is a 125-180 μm powder.

Three of the five grains (1, 3 and 5) have very similar spectra: a steep drop in reflectance up to ~400 nm, then becoming flat and almost featureless. In contrast, grains 2 and 4 show several pronounced features: there is a deep absorbance at 350 nm and a broader and shallower feature at 520 nm, centred at 490 nm. The overall spectrum reddens towards the highest wavelengths measured. The apatite spectrum measured by the RELAB system also has a reddish slope with a deep absorbance at 340 nm. There are also minor features at 580 nm and 700 - 750 nm; if these features occur in the OU spectra, they are hidden by 'noise'. It is possible that the 340 – 350 nm absorbance is from either Mn$^{5+}$ or Cr$^{5+}$ substituted into the PO$_4^{3-}$ anion (Grisafe and Hummel 1970). The longer wavelength absorptions might result from Mn$^{7+}$ in the PO$_4^{3-}$ anion (Grisafe and Hummel 1970).

The clearer and flatter grains (grains 2 and 4) yield much better spectra than the other three grains which have rougher, and therefore, more poorly-focussed...
surfaces (Figure 3.26). Grains 2 and 4 also have spectra that are the most comparable with data from RELAB. This result indicates that the flatter the grain, the better focussed is the microscope and the more detailed and complete is the final spectrum.

3.2.3 Grain size

As referred to above, the grain size of the sample influences the spectrum of the sample. Because the MSP has such a small aperture, a series of spectra were taken of the mineral orthoclase at a range of grain sizes, in order to assess the extent of grain size effect. Results are shown in Figure 3.28. Orthoclase mix is a mixture of all the grain sizes (before the sample was sieved).

![Figure 3.28: Spectra of the mineral orthoclase with a range of grain sizes taken by the MSP. Spectra taken on the MSP were averages of spectra taken in at least 5 separate locations on the sample.](image)

The results show that grain size influences the spectra and are consistent with previous work by other authors (e.g. Cloutis et al. 1986; Hapke 1993; Pieters et al. 1993; Hapke 2001b; Noble et al. 2001). Reflectivity and redness of the spectra decreases with increasing grain size as expected, with the smallest size fraction (<
30 μm) having the brightest and redder spectrum and that with grain size > 125 μm the lowest reflectance and less red spectrum. Pieters et al. (1993) also showed that the finest fraction of a suite of minerals had the highest reflectance when the samples were ground. The mixed sample had a very high and red spectrum, almost as much as the finest sample, probably because the finest grain size makes up a large percentage by volume of the mix and tend to 'cling' to the largest particles, covering them (Pieters et al., 1993; Noble et al., 2001)

It is reasonable to question why there should be a grain size effect with the MSP, given that the size of the aperture (10 x 10 μm) is below that of the smallest grain size fraction. However, there is an effect, and it is interpreted to be caused, in the same way as Pieters et al (1993), by light not being able to be transmitted through the grain and scattered back to the detector in larger particles. We also need to consider the fact that all grains within the field of view are going to have an influence on the final spectrum.

3.3 Conclusions

Previous work (summarised in Table 3.7 and Table 3.8) has shown that specific absorption bands are associated with the presence of certain cations (see e.g., Gaffey 1976; Wagner et al. 1987; Cloutis 2002; Cloutis et al. 2008). Despite the fact that an MSP has rarely been used for silicate mineralogy, results should have some similarity with results from other instruments in order to be of use to the field. The capacity of the instrument may affect the definition of absorption or reflectance bands, however features related to the composition and crystal structure of minerals should show at the same wavelengths independently of the instrument used.

There is a variety of results from the different minerals, with some having spectra that show few identifiable characteristics, whilst others produce spectra that display
features very similar to those of the RELAB or UNIWIN instruments, which are widely used by the scientific community for the study of minerals and meteorites.

From the results described in this chapter, several conclusions can be drawn from the discrepancies between the spectra from the MSP and the RELAB and UNIWIN instruments:

**Instrument Geometry:**

There are differences between the three instruments. The light path in the MSP has an angle of incidence on the sample of 0° and at the detector of 0° (i.e., phase angle of 0°). The RELAB instrument is bidirectional, with an angle of incidence of 10 - 30° degrees (usually set at 30°) and an angle of detection of 0° (a phase angle of 10 - 30°). A phase angle of 0° may cause some interference from specular reflection from both the sample (if we have a cleavage or straight surface) and the sample holder (discussed further down). UNIWIN has a phase angle of 0°, but the aperture of the instrument and the size of the sample are not known. It is assumed that the spectrometer is not attached to a microscope for small samples. Nevertheless, some results from the MSP are very similar to those from RELAB and UNIWIN, which might mean that the phase angle has a smaller influence on the results than other effects.

**Sample size:**

In the case of olivine (see appendix A), the grains are smaller than the aperture of the spectrometer and they may, in some cases, be smaller than the wavelength of the incident light. This may lead to light not being reflected from the sample. It can also be easily seen with the naked eye that they are very dark, possibly because of their size, and the results are dark, featureless spectra.
Surface Roughness:

The area surrounding the sample being analysed can influence the results. The quartz slide where the scans were taken for this chapter is transparent to these wavelengths and should emit no signal. If the beam does not hit the sample only or if the spot size is larger than expected, then the results might be diluted or, given that the slide is flat, there might be some specular reflection from the slide. Any features might then be too small to be discernible. Grains that have high relief and surface roughness are very difficult to focus on. This shouldn't influence the results, however there are consistencies on the results that show that badly focused samples produce 'bad' spectra and well-focussed samples produce spectra comparable to other instruments. The reason for this is unknown but it might be related to the configuration of the instrument or scattering effect. The same problem with focus was analysed in section 3.1.4. The beam might be focussing on something else or the light path might be altered in some way.

At this small scale with an aperture in the order of micron scale, cleavage at the surface of the grains may cause unusual visual effects caused by surface scattering, which may not have enough information on the composition of the grain. On the RELAB instrument the spot size is much larger (>1mm) so back scattering and irregularities of the grains are averaged out. It can be seen on the MSP results that different grains produce different spectra, some comparable to RELAB and UNIWIN results (when the grains are well focussed, occupying the majority of the field of view, bright enough but not with a flat surface) and others have featureless or darker spectra. Furthermore, results from the study of the reference standard in previous sections also show that the very small aperture and high magnification of the MSP lead to significant differences in spectra even when the sample is considered smooth and even to the naked eye (and other instruments).
All instruments may produce artefacts in the spectra that might cause its interpretation very difficult or even misleading. One of the artefacts visible in every spectrum, and first identified in the spectra of olivines, is that of low amplitude variations. These are reproducible so they are not caused by noise, which would be different in every spectrum. Even Fixed Pattern Noise, which has a constant pattern, should have been removed by the corrections applied. Calibration and corrections should reduce any influences outside the actual spectrum of the sample but it might be that there is some error with the corrections or the calibration or another factor related to the geometry of the instrument that causes these artefacts. One of the reasons might be because the calibration and reference material is not uniform at this scale. Another possible explanation is the much higher spatial resolution of the MSP (<2 nm) which is not achieved by most instruments, resulting in smoother spectra. The artefact seen close to UV wavelengths might also be caused by the reference material which is not suitable for NUV and also by the fact that signal from the instrument at these wavelengths is very poor.

3.4 Further work

This instrument seems to have many limitations when used in the field of mineralogy and planetary sciences. A number of conditions must be met in order to obtain similar results to those of other instruments used in the field but there is no exact formula as to how to obtain reliable results. More tests need to be made to assure the utility of this instrument. It is recommended that the following work is carried out.

- Scan a higher diversity of minerals some with little variation in composition
- The same mineral must be analysed in a variety of grain sizes
- Different sample holder must be used in order to access its influence on the final spectra
• Different amounts of samples must be scanned in order to find how this will affect the results.

Other suggestions will be presented in the following chapters. Chapter 4 describes an attempt to apply this technique to the study of micron-sized samples returned from the Stardust spacecraft from comet 81P/Wild 2. In Chapter 5, spectra taken from meteorites that have been ground to powders are presented. It is expected that mixtures of powders of different compositions might show more promising results. Meteorites are rare and their study would have much to gain in a technique that is non-destructive and requires only small amount of samples.
4 Stardust

4.1 Introduction

Comets are thought to have remained mostly unaltered since their formation in the early Solar System (see Chapter 1). Analysing material from comets can provide valuable insight into, and further our understanding of, the processes that formed the Solar System and which occurred prior to the formation of the Sun. For that reason, several missions have been sent to comets, mostly for imagery and analysis of the coma e.g. ESA’s Giotto mission (Reinhard 1986; Reinhard and Battrick 1986) or investigation of surface and subsurface composition, e.g., NASA’s Deep Impact mission (Mumma et al. 2005; Sunshine et al. 2006; Sunshine et al. 2007). However, one key mission had a fundamentally different approach: it would retrieve samples from the comet and return them to Earth, where state-of-the-art laboratory techniques could be used to analyse them. This mission was known as Stardust and provided the opportunity to study cometary samples using techniques that cannot be utilised in-situ.

The Stardust spacecraft (Brownlee et al. 2003) aimed to collect two sample sets; material from the coma of comet 81P/Wild 2 and the stream of interstellar dust entering the Solar System. Comet Wild 2, with a 5 km diameter nucleus, was chosen because up until 1974, it had an elongated orbit, with aphelion at 25 AU, perihelion ~ 5 AU and a period of 57 years (Sekanina 2003), so had not come close in to the Sun. In 1974, comet Wild 2 experienced a close encounter with Jupiter, giving it a new period of 6.4 years (with an aphelion of ~5.3 AU and perihelion of ~1.5 AU). The change in orbit meant that comet Wild 2 had not completed many orbits around the Sun since its encounter with Jupiter, and so would be relatively unaltered. At the same time, the Ulysses spacecraft detected a stream of ‘fresh’
interstellar dust entering the Solar System (Grü et al. 1993) making the Stardust mission the perfect opportunity to collect both cometary and interstellar samples.

The Stardust spacecraft was equipped with an aerogel collector grid which collected samples from the comet on one side and interstellar dust on the other side (Figure 4.1). Aerogel, a very low density material (it is made of 99.8% air and has a density of 3 mg/cm$^3$) derived from silica gel by supercritical drying, is extremely thermally resistant and can also support 4000 times its weight. As a result, it is ideally suited to the retardation and capture of particles travelling at high velocity from the comet and within the interstellar dust stream.

![Aerogel cells and collector grid of the Stardust sample collector (images courtesy NASA/JPL). The collector assembly is about 50 cm across and the cells are 1 to 3 cm thick](image)

The Stardust spacecraft was launched in February 1999, and made three orbits around the Sun before encountering Comet Wild 2 on January 2$^{nd}$ 2004. During
transit the spacecraft passed through the interstellar dust stream, collecting samples on two occasions (Figure 4.2) during the journey to the comet and while the spacecraft was moving away from the dust stream direction (reducing the impact speeds encountered). On encounter, the spacecraft passed in front of comet Wild 2, collecting samples from the coma. The collected grains were then returned to Earth, a capsule containing the collector assembly landing in the Utah desert on January 15th 2006. Figure 4.2a shows the path taken by the Stardust mission. An image taken by the Stardust spacecraft (Figure 4.2b) shows the cometary nucleus to have an irregular surface that is cratered in appearance.
On impact with the aerogel, captured particles form tracks (Burchell et al. 2008). Three different track morphologies (A, B and C) have been identified (Hörz et al. 2006). A-type tracks are typically single, long, narrow, carrot-shaped tracks with the impacting particle found at the track termination point. The particles that form these tracks are thought to be strong consolidated grains, whilst the particles that form the remaining tracks are thought to be less coherent or more volatile (Hörz et al. 2006).

B-type tracks (Figure 4.3) are bulbous and are formed when the particle heats and explodes, losing its gases resulting in the widening of the track at one end (Hörz et al. 2006). The remaining sample can be collected at the end, and occasionally on the walls of, the track. C-type tracks are similar to type B but exhibit the wider feature only (Hörz et al. 2006).
4.2 Results on Stardust grains from other investigations

Preliminary results published in a Science special issue (Keller et al. 2006; McKeegan et al. 2006; Sandford et al. 2006a; Sandford et al. 2006b; Zolensky et al. 2006a) show a range of compositions and possible sources for the material recovered from both the aerogel and the aluminium foil craters. The material collected is rich in amorphous silicates and with some crystalline silicates such as olivine and pyroxene. These silicates are formed at high temperatures, indicating that they were formed closer to the sun and then transported to the Kuiper belt before the aggregation of the comet. (McKeegan et al. 2006). The composition of this material is consistent with theories indicating a mixing of solar system and interstellar matter within the protoplanetary disc. Most material, however, seems to originate from the Solar System and not presolar materials (McKeegan et al. 2006).
Samples collected from comet Wild 2 show similarities in chemical, mineralogical and isotopic composition with IDPs, micrometeorites and carbonaceous chondrites suggesting a continuum between asteroidal matter and comets. Together with analysis of refractory material (Chi et al. 2009) they seem to confirm the idea that they are composed of material formed in different locations, through different processes and were then subject to radial mixing in the early solar nebula (Dobrica et al. 2009).

The absence of hydrous phases is indicative of no or very mild aqueous alteration (Zolensky et al. 2006 and Keller et al. 2006). However results on a TEM and EMPA analysis of sulphides from comet Wild 2 (Berger et al. 2011) reveal that they have been formed via low temperature aqueous processes. The authors suggest that sulphides such as troilite formed in the inner Solar System and have been transported to the Kuiper belt region by radial mixing and were then subject to aqueous alteration. The liquid that led to the formation of these sulphides formed by aqueous alteration would have been formed by micro-impacts or short-lived radionuclides.

4.3 Samples

Initial studies of the returned samples were undertaken by Preliminary Examination Teams (PETs) from various institutions across the globe. The PETs, along with the techniques to be used, were established before sample return. As many non-destructive techniques as possible were chosen for preliminary examination, to minimize sample loss. Our team was selected to measure the spectrum of the grains between 250 nm (in the near ultra violet; UV) to 800 nm (near infrared; IR) with a non-destructive, state-of-the-art instrument. UV/Vis spectroscopy was to be used to characterise the grains, and is complementary to other spectroscopic techniques. UV/Vis spectroscopy should be able to produce results that could be
matched directly with spectra acquired remotely (telescope or spacecraft instrumentation). Together with other analysis techniques, UV/Vis spectroscopy can give a deeper insight to the materials, environments and processes that occurred from the central to the outer region of the Solar System and around other stars by comparing the returned Stardust samples from comet Wild 2 to primitive meteorites, IDPs, and spectra acquired remotely.

The team received two grains, extracted from two different tracks. The sample reference numbers (C2054,0,35,1,0 and C2054,0,35,28,0) refer to the aerogel cell from which they were removed and their position relative to the track on the aerogel grid. Here, we will refer to them as grain A and grain B respectively.

The grains were extracted from aerogel and pressed into one gold foil by Lindsay Keller from the NASA Johnson Space Center (JSC). Images of grains A and B (Figure 4.4) on the gold foil were taken at the OU using a high spatial resolution FEG-SEM (Zeiss Supra 55 field emission gun scanning electron microscope).
Figure 4.4: SEM images of Stardust grain A (top and middle) and B (bottom). In the middle image, the aerogel coating is clearly visible.
Figure 4.5: SEM analysis results on grain A (top) and grain B (bottom). Only the Si and O of the aerogel was detected, the Au peak is from the gold foil substrate on which the particle is supported.

The samples are approximately 10 µm (grain A) and 8 µm (grain B) across. A preliminary analysis of the samples noted that they seemed to be grain aggregates, possibly melted together. Both grains were surrounded by aerogel, which probably melted on impact and fused with the sample (Figure 4.4). Figure 4.5 shows the SEM
analysis of the grains where only the Si and O of the aerogel can be detected. Because of the non-destructive nature of this investigation (as required by the Stardust Preliminary Examination Team), the samples could not be removed from the melted aerogel. As a result, spectra had to be obtained through the aerogel. Being primarily composed of silica, aerogel should be transparent at UV/Vis wavelengths. However, impurities or other aspects of the aerogel may affect the ability to obtain UV/Vis spectra of the samples. For this reason, calibration tests were carried out in order to evaluate the effects of the aerogel on the final spectra. Samples of flight-spare aerogel from the Stardust mission were made available for analysis.

4.4 Aerogel and gold foil tests

As a preliminary test, and because the samples were pressed into gold foil (see previous chapter for influences of the background on spectra), a spectrum was taken of a gold foil and then of gold foil through several pieces of aerogel of different thicknesses. Unfortunately, because of their transparency and difficulty in handling, it was not easy to measure the exact thickness of the aerogel pieces, but they were all slightly more than 2 mm thick. The results of the tests on aerogel and gold foil are showed in Figure 4.6.
Figure 4.6: Reflectance spectra of gold foil and of gold foil covered with several pieces of aerogel with different thicknesses. Because of the brittle and transparent nature of the aerogel, it is not possible to determine the relative thicknesses of the samples of aerogel.

The spectrum of gold foil itself is shown as the black curve in Figure 4.6. It is characterized by a single broad minimum between 300 and 500 nm, followed by a rapid rise to ~575 nm (the yellow region of the spectrum), then a more gentle increase up to 800 nm. It is clear from Figure 4.6 that the aerogel produces a difference in the gold spectrum, particularly by reducing the intensity of the reflected light. The figure shows that all the spectra taken through aerogel are darker than the spectrum of gold foil. This difference was not unexpected, because to look at, aerogel is not completely transparent, so some scattering of light must occur. Most of the features in the gold spectrum are still present, but reduced in intensity - the features are wider and not as deep. This effect will be referred to as smoothed from this point on. There are very few differences between the individual aerogel spectra; where there is a difference, it occurs above 700 nm. The 700 nm occurrence might be because of impurities and faults in the gold foil and/or the aerogel. Despite the
fact that the gold foil is clean and of extreme purity, at these magnifications several flaws and faults are visible.

Because gold has such a high reflectance, particularly above 500 nm, and is, apart from that, almost featureless, it was considered that this test would be more successful if made on a coloured surface. The experiment was repeated on a piece of green paper. Again the aerogel thickness was similar but could not be determined. The results are shown in Figure 4.7.

![Reflectance spectrum of a sheet of green paper (black curve) and from the same green paper covered with several pieces of aerogel with different thicknesses. Because of the brittle and transparent nature of the aerogel, it is not possible to determine the relative thicknesses of the samples of aerogel.](image)

As was observed for aerogel on gold foil, the results show a darkening and smoothing of the spectrum. There is only a small change in the aerogel on green spectrum at 300 nm, but this may also be because of some impurity on the paper and/or the aerogel. As described before, at these magnifications, gold, paper and aerogel are very heterogeneous.
On the basis of these results, it seems that aerogel will not introduce a significant difference to the spectra of the Stardust samples, except to reduce their reflectance, which would make it very difficult, if not impossible, to discern any weak features. It must also be considered that the flight aerogel may contain impurities, even though it was made in an extremely clean environment.

4.5 Sample data collection and analysis

The next step was to take spectra of the two cometary samples we received. Because of the small size of the particles, the only magnification available for use was 52x, giving an aperture of 2 x 2 μm. Several spectra were taken from different areas of the samples, in order to determine if they are homogeneous. Pictures of the samples and where the aperture was placed are shown in Figure 4.8 and Figure 4.9 shows all the spectra taken on the samples.
Figure 4.8: Images taken by the microspectrophotometer of both the samples. Within the circles is where the aperture was placed on the scans shown in Fig. 4.9 (black square).
By analysing the spectra of the two particles a few facts can be deduced. The spectra seem to be flat, featureless and very similar to each other below 500 nm. There is a common, sharp rise in reflectivity at 500 nm and then a gradual increase in reflectance up to 800 nm. The reddening and the features are different for each spectrum which leads us to believe that there are differences in composition between grains and even within the same grain. In order to analyse the results more thoroughly, and compare the samples with each other, the results of the two grains were plotted separately (Figure 4.10a and b)).
It seems clear that there are differences between the samples and within the same sample. Some spectra present the same features but they seem smoothed. The most striking characteristic of the spectra that we can observe is how similar they are to the spectrum of gold. They are flat, featureless and very similar at wavelengths below 500 nm, above which there is a pronounced reddening of the
spectra. This is a concern, because if the spectra are so similar to the gold spectrum, it presumably means that the gold foil is influencing the results. This should not happen, as the aperture of the spectrometer is smaller than the grains (as can be seen in Figure 4.8), and the spectra are not taken on the gold. It is however possible that light is reflected and scattered in different directions by the gold foil and that some of this light reaches the aperture of the spectrometer. If this is true, then spectra taken closer to the border of the grains should have a higher reflectance than spectra taken in the middle of the grains. Examination of Figure 4.8 and Figure 4.9 shows that this is, indeed the case, and there is a relationship between the position of the aperture and the brightness of the spectrum. When the aperture is closer to the border of the sample, the spectrum becomes much brighter. So, for sample A, spectra A and A2 are brighter than A1 and A3, and were taken closer to the margins of the sample. The same is true for sample B, where spectrum B is brighter than B2 or B3, and was taken from the edge of the grain. It seems that reflectance from the gold foil does influence the measured spectra. These results confirm the observations in chapter 4 where it was noted that the area surrounding the sample had an influence on spectra, possibly caused by the scattering of light from that area into the aperture of the instrument. Unfortunately the sample cannot be removed from the gold foil, but we can attempt to correct for the presence of the gold. In an attempt to analyse better the characteristics of the spectra of the grains without the influence of gold, the spectra of the former were divided by the spectrum of the latter. The results are shown in Figure 4.11.
Once the spectrum of the gold foil is removed, more pronounced features can be observed in the spectra of the grains. It is obvious that there are differences between grains and within the same grain as stated previously. There is one further step that has to be taken before we can attempt to interpret the Stardust spectra, and that is to remove any effects resulting from the presence of aerogel. The results
shown in Figure 4.6 allow a spectrum of aerogel to be calculated, by taking the spectra in Figure 4.6, and dividing them by the spectrum of gold foil. The results of this procedure are shown in Figure 4.12.

Figure 4.12: Reflectance spectra of three samples of aerogel with slightly different thicknesses on gold foil (black), and an average of these spectra (purple). All spectra have been corrected for the presence of gold foil by dividing the (aerogel + gold) spectrum

The final spectra for Stardust Grain A and Grain B are shown in Figure 4.13 and Figure 4.14. The measured spectra have been corrected for contributions from gold foil and from aerogel. It must be noted that it is impossible to determine how much the spectrum of gold and aerogel will influence the sample, because these depend on the proximity to the border of the grain (for the gold) and the thickness of the aerogel. It is however expected that the main features should still be observable and accurate, considering the transparency of aerogel and the flatness of the spectrum of the gold foil below 500 nm.

The corrected spectra for the two grains are similar, but there are differences between and within them.
Sample A: The four scans taken from sample A are quite similar to each other; their differences reside mainly in the brightness and relative intensity of features, which may result from small differences in composition. All the scans show a small minimum at ~ 330 nm. This feature is present in minerals like diopside (Gaffey 1976), where the cause is Fe³⁺, and in augite and ilmenite, where the cause is Fe²⁺ – Ti⁴⁺ charge transfer (Gaffey 1976 and Wagner 1987). Additional minima also occur at 380 - 400 nm, 440 nm, 550 nm, 640 – 660 nm, 730 nm and 760 – 770 nm. Table 5.1 summarises possible assignations for these features, based on comparison of data from other studies. The first of these may be caused by Fe²⁺-O or Fe³⁺-O charge transfer in minerals like enstatite, as may the feature at 550 nm. A small shoulder at 730 nm occurs in magnetite (Gaffey 1976). A weak minimum at 760 nm is present in phyllosilicates and is caused by Fe charge transfer. In hedenbergite, the feature is caused by Fe²⁺ - Fe³⁺ charge transfer (Cloutis 2002). It is clear that Sample A is not a single grain, but is an assemblage of different minerals. A very tentative conclusion, given the uncertainty in the corrections that have been applied to the data, is that Sample A seems to be of at least two different types of pyroxene, plus magnetite and a phyllosilicate.
Figure 4.13: Corrected spectra from Stardust grain A, taken at the locations shown in Figure 4.8

Table 4.1: Summary of features seen in Stardust Sample A

<table>
<thead>
<tr>
<th>Wavelength (nm)</th>
<th>Species</th>
<th>Transition</th>
<th>Mineral Phase</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>330</td>
<td>Fe^{3+}; Fe^{2+}; Fe^{2+}-Ti^{4+}</td>
<td>Charge transfer</td>
<td>Diopside, augite</td>
<td>1, 2</td>
</tr>
<tr>
<td>380 - 400</td>
<td>Fe^{3+}; Mn^{3+}</td>
<td>Charge transfer</td>
<td>Enstatite</td>
<td>3, 4</td>
</tr>
<tr>
<td>440</td>
<td>Mn^{3+}</td>
<td>Crystal field transition</td>
<td>Diopside, augite</td>
<td>3</td>
</tr>
<tr>
<td>540 - 550</td>
<td>Fe^{3+}; Mn^{2+}</td>
<td>Spin-forbidden CFT</td>
<td>Enstatite</td>
<td>3</td>
</tr>
<tr>
<td>640 - 660</td>
<td>Cr^{3+}</td>
<td>Crystal field transition</td>
<td>Diopside, augite</td>
<td>3</td>
</tr>
<tr>
<td>730</td>
<td>Fe^{3+}</td>
<td>Charge transfer</td>
<td>Diopside, magnetite</td>
<td>1</td>
</tr>
<tr>
<td>760 - 770</td>
<td>Fe^{3+}</td>
<td>Charge transfer</td>
<td>Olivine</td>
<td>5</td>
</tr>
</tbody>
</table>

References: (1) Gaffey, 1976; (2) Wagner, 1987; (3) Cloutis, 2002; (4) Cloutis et al., 2008; (5) Hunt 1977

Sample B: The scans taken from sample B are also quite similar to each other and to the scans taken from sample A. They seem to be slightly brighter, but this might be caused by a stronger influence of the gold foil spectrum, because the sample is smaller and the percentage of gold foil in the field of view is higher. Other differences might be caused by small changes in composition. Sample B exhibits a
strong absorbance at 360 – 380 nm, and broad features at 400 nm and 520 - 560 nm, as well as weak absorbance features at 320, 330, 450, 620 and 710 - 720 nm. Again, these are all features that can be assigned to silicate minerals and magnetite. Sample B, like Sample A, is an assemblage of different mineral grains and not a single grain.

Figure 4.14: Corrected spectra from Stardust grain B, taken at the locations shown in Figure 4.8

In summary, after correction for gold and aerogel, the spectra analysed here are consistent with the presence of silicate minerals such as both high- and low-calcium pyroxenes and olivines, and opaque minerals such as magnetite. These minerals have been found in Stardust grains by other authors. These analyses were performed on the craters that some grains left on the aluminium foil (Zolensky et al. 2006b), or, when conducted on the grains collected on the aerogel, using more invasive techniques that required the slicing and embedding of the grains in other material (Zolensky et al. 2006b; Leroux et al. 2008).
4.6 Comparison with previous data

4.6.1 Comparison with IDP

There has only been one other attempt to determine the reflectance spectra of interplanetary dust particles using a microspectrophotometer (Bradley et al. 1996). The authors used a microspectrophotometer to measure the visible spectrum of 5 chondritic smooth IDP and 4 chondritic porous IDP collected from the stratosphere (Figure 4.15). They also measured the spectra, at the same wavelengths, of the carbonaceous chondrite Allende (Figure 4.16).

A strong fall off below 400 nm is visible in all the spectra collected by Bradley et al. (1996) (Figure 4.15). The reason for this is not certain but the authors attribute it to two possibilities. One is the change from diffuse to specular scattering conditions and another is the positioning of diaphragm apertures close to the edge of minerals that may cause diffraction effects in the UV. This fall-off does not happen with the spectra taken by the MSP at 400 nm but in some spectra there is a strong artefact below 350 nm. There is not, however, a direct connection to the positioning of the aperture.

Sample results in Bradley et al. (1996) show that anhydrous IDPs, even with different compositions, show similar, slightly flat to reddened slopes. Their composition varies from single grain to chondritic element assemblies. The spectra of single grains are, in general, brighter and the C-rich compositions are usually the darkest. Samples rich in C and aromatic compounds show a distinctive red slope, especially above 500 nm, e.g., particle D in the top plot in Figure 4.15. The spectra of the Stardust particles analysed in this work (Figure 4.13 and Figure 4.14) also have a reddened slope at wavelengths above 500 nm, but it is not certain that the cause is the presence of aromatic compounds. Bradley et al. (1996) conclude that the presence of magnetite formed by heating during atmospheric entry of the IDPs.
can be identified by a darkening of the spectrum at UV-Vis wavelengths, but no other conclusions are drawn regarding mineralogy. The spectra of Stardust particles taken by the MSP show more detail, with various features visible (described above). This might be caused by the fact that the MSP has greater sensitivity and resolution, but the influence of the gold foil and the aerogel cannot be discarded. Because of the strong drop-off below 400 nm, no detail can be seen in the spectra taken by Bradley et al (1996). The lack of C in the Stardust samples may also account for the more detailed, less flat spectra compared with the ones in their work. Both works seem to indicate that some IDPs and Stardust samples are aggregates of different minerals with Fe silicates such as olivine and pyroxene and other Fe bearing minerals.

Figure 4.15: Reflectance spectra from four 20-40 mm diameter fragments of matrix from Allende CV3 chondrite. Fragments B, C and D are shifted upwards from A by 1, 2 and 3 reflectance units, respectively. The figure and caption are Figure 3 from Bradley et al. 1996
4.6.2 Comparison with CV and CM chondrites

IDPs are thought to be of cometary or asteroidal origin with chondritic composition (see Chapter 1). Hydrated IDPs have similar spectra to CM carbonaceous chondrites. For this reason, spectra were acquired from the CM2 meteorite Murchison and the CV3 chondrite Allende.
Murchison is a carbonaceous chondrite rich in organic materials and has been subjected to aqueous alteration (e.g. Hutchison 2004). Murchison has been extensively studied particularly because of its amino acid content (e.g. Kvenvolden et al. 1970). Amino acids such as purines and pyrimidines are building blocks of DNA and the connection between their presence in meteorites and the existence of life on Earth is being studied (e.g. Martins et al. 2008). Allende is a CV3 meteorite, rich in refractory materials (Calcium- and Aluminium-rich Inclusions (CAIs)) (see Chapter 1 for more details). Figure 4.17 shows images taken by the MSP of Murchison (left) and Allende (right) showing the aperture of the spectrometer (black square). Figure 4.18 shows the spectra of both meteorites for comparison.

Samples were prepared in the same way as the mineral samples in Chapter 3 – powdered samples placed on a quartz slide, but in this case there was sufficient material to fill the field of view of the MSP.
The Allende spectrum shares the low reflectivity seen in Bradley et al (1996) as does Murchison. Both spectra are flat, dark and featureless, possibly as a result of being rich in carbon. Both spectra are featureless with a slight reddening of the spectra above ~ 620 nm, with Murchison darker and redder than Allende.

4.7 Conclusions

Analysis of two particles from Comet Wild 2 demonstrated the utility of the microspectrophotometer for non-destructive characterisation of small samples. There were analytical problems because the particles were embedded in aerogel and pressed into gold foil, but careful acquisition of spectra from gold foil and from aerogel supported on gold foil allowed a qualitative correction for these materials, and deduction of the UV-Vis spectra of the cometary particles.

Results from the spectral measurements showed that the two particles (Grain A and Grain B) had very similar spectra, implying similar compositions. Comparison of the spectra with mineral data suggested that the particles were a mixture of silicates.
(e.g. pyroxene and olivine), with opaque minerals (such as magnetite). These results are in keeping with the findings from analyses of Wild 2 particles by other techniques, which show that comet Wild 2 is composed of a large variety of material (e.g. Keller 2006). These include silicates and refractory minerals, which were formed at very high temperatures, suggesting a mixing of material from the inner to the outer protoplanetary disc, and organics and other material which have not been heavily processed since their accretion in the comet (e.g. Weissman 1985; Feldman 1987; Greenberg and Hage 1991; Greenberg and Li 1998). Silicates present in the interstellar medium are amorphous whereas a major fraction of the grains found in Stardust samples are crystalline, indicating that they were produced in the Solar System (Brownlee et al. 2006). The suggestion that they might have formed by annealing of amorphous pre-solar grains is countered by their isotopic compositions which indicate Solar System origin (McKeegan et al. 2006). The varied range of olivine and low-Ca pyroxene indicates a varied range of formation conditions and the presence of Ca-rich pyroxenes further supports this hypothesis since they are indicative of processed Solar System materials (Zolensky et al. 2006b).

These observations contradict the previous belief that comets were dominated by pristine pre-solar grains or interstellar material. Instead, they are a collection of a variety of material - pristine and processed - of pre-solar and solar origin. The main aspect of these results is that the study of comets provides not only valuable insights into the processes and conditions of formation of the Solar System but also clues to the question: "Has the origin of life on Earth been triggered by the impacts of cometary material?"

Analysis of carbonaceous chondrites by UV-Vis spectroscopy has revealed little about their composition except for the fact that they have high abundance of carbon in their composition which turns the spectra flat and dark (e.g. Wagner et al. 1987; Clark 1983).
5 HED meteorites and their parent bodies

5.1 Introduction

As described in Chapter 1, one of the purposes of this work is to identify if the microspectrophotometer (MSP) can be an aid in the study of meteorites and their parent bodies. This means, not only tentatively identifying aspects of the composition of meteorites and IDPs, but also helping to establish links between these and their parent bodies whether they are asteroids, comets or Mars.

Chapter 5 contained a description of the application of the instrument to the study of cometary grains and carbonaceous chondrites and their possible parent bodies. In this chapter, result from HED meteorites (definitions to follow) and links to possible parent bodies are presented. If successful, this technique can also be applied to the study of recently discovered basaltic micrometeorites (Gounelle et al, 2009) and contribute to the investigation of their parent body.

5.2 HEDs and Vesta

Howardites, eucrites and diogenites (HED) are differentiated meteorites, i.e., their parent body has experienced some melting and processing. Diogenites are generally spinel-bearing orthopyroxenites (Hutchison 2004). Eucrites are composed mainly of calcic plagioclase and Ca-poor pyroxene and have textures similar to those of terrestrial basalts and gabbros and may include breccias (Duke and Silver 1967). Howardites are all breccias, and are composed of eucrite and diogenite debris plus some chondrite fragments (Hutchison 2004). They contain solar wind gases, implying that they come from the surface of the HED body. The chemistry, and particularly the oxygen isotopic composition of the three groups of meteorites suggests that they originate from the same parent body (Mittlefehldt 1994; Mittlefehldt and Lindstrom 2003; Greenwood et al. 2005). Pb-Pb age dating of HED
show that they have a crystallization age of at least 4.51 Ga, making them amongst the oldest basalts in the Solar System (Tera et al. 1997) and references therein.

![Figure 5.1: Laboratory measurements of the spectral reflectivity of the Nuevo Laredo meteorite (eucrite) and telescope data from Vesta taken from McCord (1970)](image)

The first link between HEDs and Vesta was made by McCord et al. (1970) using Vis-NIR spectroscopy data collected by two telescopes (Figure 5.1). They identified a 900 nm feature associated with Fe$^{2+}$ in the M2 site of magnesium-rich pyroxene, and a shallow band between 500 and 600 nm, possibly from Ti$^{3+}$ in pigeonite, and suggested that basaltic achondrites (a eucrite) were the closest match to the spectrum of Vesta. Since then, it has been generally accepted that Vesta is the parent body of the HED meteorites (see for example McSween Jr. 1999; Burbine et al. 2002; Keil 2002; Pieters et al. 2005a; McSween et al. 2011 and references therein) with different sides of Vesta with more similar spectra to either howardites, diogenites or eucrites. In his paper, Takeda (1997) further classifies HED meteorites into more specific categories, such as eucrites originating at different depths in the crust (Figure 5.2 and Figure 5.3), and links the different types to areas and geological features of the asteroid observed by telescopes, such as large craters.
TABLE I. Explanation for subclasses of the howardite, eucrite, diogenite achondrites.

<table>
<thead>
<tr>
<th>Class</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diogenites</td>
<td>Monomict or unbrecciated achondrites with major orthopyroxene</td>
</tr>
<tr>
<td>Eucrites</td>
<td>Pyroxene-plagioclase achondrites</td>
</tr>
<tr>
<td></td>
<td>A. Non-Cumulate Eucrites</td>
</tr>
<tr>
<td></td>
<td>a. Ordinary (O-) eucrites (monomict or unbrecciated)</td>
</tr>
<tr>
<td></td>
<td>- Basaltic rocks with homogeneous host pigeonite and fine exsolution lamellae</td>
</tr>
<tr>
<td></td>
<td>b. Surface (S-) eucrites (passamonte-type or lava-like eucrite)</td>
</tr>
<tr>
<td></td>
<td>B. Cumulate (C-) eucrites</td>
</tr>
<tr>
<td></td>
<td>(i) Binda-type</td>
</tr>
<tr>
<td></td>
<td>(ii) Moore Co.-type (Serra de Magé)</td>
</tr>
<tr>
<td></td>
<td>C. Polymict eucrites</td>
</tr>
<tr>
<td></td>
<td>Polymict breccias of various types of eucrites</td>
</tr>
<tr>
<td></td>
<td>- With &lt;10% (modal) orthopyroxene</td>
</tr>
<tr>
<td>Howardites</td>
<td>Polymict breccias of diogenites and eucrites</td>
</tr>
<tr>
<td></td>
<td>&gt;10% (modal) orthopyroxene</td>
</tr>
</tbody>
</table>

Figure 5.2: Subclasses of eucrites, howardites and diogenites by Takeda, 1997

Figure 5.3: A model of a layered crust of the HED parent body with four hypothetical impacts, producing various polymict breccias, polymict eucrite (top two), howardites (bottom) and polymict eucrites with <10% orthopyroxene

In summary, eucrites are basaltic rocks that seem to originate from the crust of 4 Vesta, Diogenites are similar to plutonic rocks composed mainly of Mg-rich pyroxene with some plagioclase and olivine, possibly from deeper within the asteroid and howardites are surface breccias mainly composed of pieces of diogenites and eucrites with clasts of carbonaceous chondrite.
Dawn Mission

Vesta and Ceres are the largest bodies in the asteroid belt and their study is essential in order to understand better the processes of melting and planetary evolution of the Solar System. For this reason, the Dawn mission was designed with the purpose of orbiting and studying first the asteroid 4 Vesta (Figure 5.4) and then the asteroid 1 Ceres. Ceres does not have any recognised associated meteorites; the study of HEDs does not tell us all the information about Vesta that we could get from a purposely built mission. A good review of the mission, its scientific justification and payload of the Dawn mission can be found in Russell and Raymond (2011).

Figure 5.4: NASA’s Dawn spacecraft obtained this image of the giant asteroid Vesta with its framing camera on July 24, 2011. It was taken from a distance of about 3,200 miles (5,200 kilometers). Credit: NASA/JPL-Caltech/UCLA/MPS/DLR/IDA. Image and caption from http://dawn.jpl.nasa.gov/multimedia/dawn_vesta_image_full_072411.asp

Preliminary results show that dark material found on the walls of craters on 4 Vesta could be shocked eucrite/impact melt, possibly with clasts of carbonaceous chondrites (Reddy et al. 2012a). The link between the HEDs and the asteroid are also strengthened by VIR maps from the spacecraft, which show mostly howardites-
like spectra at the surface, with diogenites-like material inside craters and eucrite-enriched howardites at shallow depths (McSween et al. 2012).

5.2.1 UV-Vis Microspectroscopy of two Eucrites and one Diogenite

Spectra of the meteorites EETA 79006 (eucrite), Johnstown (diogenite) and Juvinas (eucrite) were taken by the MSP. The sampling and data collection is the same as described in Chapter 3: a few grains of each meteorite were placed on a quartz slide and analysed on the MSP. In this case the samples of the eucrites are powdered and the diogenites are small grains of ~ 50 μm diameter. Following from the results described in Chapter 3, care was taken to ensure that there was sufficient material to fill the field of view of the MSP. This meant that all light entering the MSP came from the sample. Figure 5.5 shows an image of one of the site where the spectra were taken in all three meteorites. Figure 5.6 shows the average spectra of the same meteorites. Spectra presented is an average of at least 5 spectra taken at different sites on the sample.
Figure 5.5: Images taken by the MSP of grains of HED meteorites. a) Juvinas, b) EETA 79006, c) Johnstown. The black square in the middle is the aperture of the spectrometer and is 10 x 10 µm in size.

Figure 5.6: Visible spectra of the HED meteorites taken by the MSP.
The spectra seen in Figure 5.6 seem to be consistent with the spectra of the same meteorites taken by other authors (such as McCord et al. 1970; Gaffey 1976). The same shallow band between 450 and 600 nm is visible. Gaffey (1976) attributes the similarity of the spectra of the three meteorites with the fact that their composition is dominated by the same mineral: the Ca-poor pyroxene, hypersthene, mixed with varying amounts of plagioclase. Because of the lower resolution and longer wavelength range of the spectra given by Gaffey (1976), the shallow band centred at \(~\) 500 nm is visible as a narrow, weak absorption feature. Another shallow band just before 700 nm and a peak above this wavelength is also visible both in the work of Gaffey (1976) and the OU data (Figure 5.7).

The most prominent feature seen in Figure 5.6 is that of the deep absorbance centred between 300 and 400 nm. This is not recorded by Gaffey (1976), as his wavelength range did not reach down to that value. The feature reaches a minimum at 340 nm in the eucrites, and 320 nm in the Johnstown diogenite. By analogy with the results from single minerals documented in Chapter 3, and results by Reddy et al. (2012b) it is likely that this absorbance results from Charge transfer effects. From Figure 5.6, it can be seen that above 400 nm, the two eucrites, EETA 79006 and Juvinas, have a similar reflectivity, albeit Juvinas has a redder spectrum and the features seem to be sharper. Johnstown has a higher reflectivity and sharper features than the eucrites. The reason for the difference between the eucrites and diogenites is probably because the diogenite is almost all orthopyroxene, whilst the eucrites are mixed with plagioclase.

As with the data taken from minerals in chapter 3, the spectra from meteorites were also compared, when possible, to the ones taken at the RELAB facility. Figure 5.7 a) and b) show the spectra of Juvinas and Johnstown (respectively).
Figure 5.7: Spectra of a) the eucrite Juvinas taken by the MSP and in the RELAB facilities; b) the diogenite Johnstown taken by the MSP and the RELAB facilities. RELAB results have been offset by +0.4 to aid the comparison.
The spectrum of Juvinas taken by the MSP is quite similar to the one taken at the RELAB facilities but slightly redder. It also seem to present more detail in some areas such as more visible peak at ~580 nm and the broad absorption at ~600-700 nm is also more clear.

The opposite occurs with the spectra of Johnstown. The RELAB spectrum shows a much clearer and sharp peak in reflectivity at ~500 nm than the MSP and the peak at ~700 nm is also more evident. However, the band at ~420 nm is clearer on the MSP spectrum and the shallow band between 500 and 700 nm is also deeper. Overall the spectrum from the MSP is less red than the one taken by RELAB.

In order to understand the contribution the MSP technique might make to the link of meteorites to their parent bodies, the spectra taken by the MSP was compared to that of asteroid 4 Vesta taken by the SMASSII survey (Xu 1994; Xu et al. 1995) (Figure 5.8).

![Figure 5.8: Spectra of Juvinas, EETA 79006 and Johnstown taken by the MSP and the asteroid 4 Vesta taken by the SMASSII survey (Xu, 1994 and Xu et al. 1995). The spectrum from Vesta has been offset by +0.4 for better comparison](image-url)
Even though all spectra share some similarities, it is the spectra of Juvinas that is more similar to the spectra of 4 Vesta especially in terms of redness. The spectrum of the asteroid is very noisy and this prevents the smaller and shallower feature from being observable, if they are, at all, present. Just like the initial results from McCord (1970), the HED meteorites seem to be a very good match to the asteroid 4 Vesta. This similarity together with the comparison with the RELAB spectra shows that the MSP is a suitable technique to use in the field of meteorites and their links to parent bodies if the conditions noted in chapter 4 are followed.

**Applications to the study of micrometeorites**

A new type of micrometeorite has been reported by Gounelle et al. (2009). It is the first basaltic micrometeorite found and analysis shows that its parent body is not the same as for other extra-terrestrial basaltic samples available: Mars, the Moon and Vesta. It seems to originate from a non-Vestoid asteroid and it provides, therefore, a new opportunity to study planetary crusts in the Solar System. Given its small size (140x100 \(\mu\)m) it is an ideal candidate for study with the microspectrophotometer in order to attempt to identify its parent body. It is the perfect example of how this technique can aid in the study of parent bodies, when the sample available is too small for other spectrometer and too unique for destructive techniques.

**5.3 Conclusions**

By analysing the spectra of the HED meteorites EETA79006 (eucrite), Juvinas (eucrite) and Johnstown (diogenite), it is clear that the MSP is a suitable technique for the study of the spectra of meteorites and a tentative approach to relate and link their spectra to the spectra of possible parent bodies.

It can be seen in Figure 5.5 that the parameters identified in Chapter 3 and 4 for the good collection of suitable spectra – clear and focused image, samples covering the entirety of the field of view. When compared with spectra taken by other authors
such as McCord (1970) and Gaffey (1976) and spectra taken from the RELAB instrument, the results are very similar and the same features are present. Furthermore, when compared with the spectrum of the asteroid 4 Vesta, as it would be expected, considering the above is true, the spectra of the HED meteorites taken by the MSP show a visible resemblance to that of the asteroid.

The conclusions of this chapter are consistent with the conclusions from Chapter 3 and 4. For future work, some care must be applied in finding suitable samples and a database of spectra of different types of meteorites must be taken for comparison with the spectra taken from asteroids either by telescope or by space missions such as Dawn. There is no doubt that Dawn will bring many answers, and possibly even more questions, relating to the formation of the asteroids 1 Ceres and 4 Vesta and, consequently, the formation of the solar system. The study of meteorites will also aid in the understanding of results from the spacecraft.
6 The effects of high velocity impacts on the IR spectrum of basalts

6.1 Introduction

Space weathering is a modification process that changes the optical properties, physical structure or chemical or mineralogical properties of the remotely sensed surface of an airless body (Clark et al. 2002). The main agents that cause space weathering on asteroids are bombardment of the surface by solar wind and by micrometeorites (e.g. Pieters et al. 2000; Hapke 2001b; Chapman 2004 and references therein). The most significant by-product of space weathering is the production of tiny grains of iron metal (nanophase iron) in the outer rims of grains exposed to the bombardment (e.g. Hapke 2001b; Sasaki et al. 2001 and references therein). Space weathering is believed to be one of the causes of the differences between the spectra of meteorites and their presumed parent bodies. These differences mainly consist of a reddening and darkening of spectra and reduction in strength of the absorption features. Asteroids seem to be affected by space weathering to different degrees, even across different regions of the same body; some do not show any signs of weathering. The Galileo spacecraft showed that the asteroid Gaspra (S-type) had colour differences related to topography which indicated more altered and fresher surfaces. Belton et al. (1992) and Belton et al. (1994) described even greater colour differences on the asteroid Ida, associated with fresh impacts and their ejecta. There seems to be some process that transforms the spectra of Ida from close to an ordinary chondrite (OC) to an S-type asteroid. Other studies of space weathering on asteroids found that there is a range of maturity with a variety of spectra from Q-type to S-types within the NEA population (Binzel et al. 1996).
Many experiments have been made in order to determine the cause of space weathering. Because no samples of asteroidal soil were available (until the return of small amounts of material from the Hayabusa mission in 2011), lunar samples have been used in laboratory experiments as an analogue of asteroid regolith. There are, however several factors that must be taken into consideration when applying lunar soil studies to the study of asteroid weathering. Firstly is the fact that asteroids are much further away from the Sun than the Moon. For this reason they will not be as affected by solar wind as the Moon. Secondly, the speed of the impactors in the asteroid belt is lower (Clark et al., 2002). Mineralogical differences are also an important factor: on asteroids, the most common mineral is olivine, whilst on the Moon, the predominant mineral is feldspar (Clark et al. 2002). This fact affects the quantity of Fe available for the formation of nanophase metallic iron particles (or submicroscopical reduced iron; SMFe). The main outcome of experiments on lunar soil is the detection of SMFe particles on the upper surface of grains (Hapke 2001b). This metallic iron, produced by vapour deposition, causes the reddening and the darkening of the spectra and also the decrease in strength of the spectral features.

### 6.1.1 Laboratory simulations

In 1993, Pieters and Fincher studied the influence of the finest fraction of particles on spectra. They concluded that weathering is surface-related and not volume-related, adding to the idea that the alteration of spectra is caused by coating of particles with SMFe. The optical effects of submicroscopical reduced iron are non-linear and related to particle size. Because opaque and packed grains inhibit multiple scattering, carbonaceous materials and bare rock surfaces are less readily space weathered.

Yamada et al. (1999) and Sasaki et al. (2001) used pulsed laser irradiation of OC in order to simulate micrometeorite impacts. The results show that both olivine and
pyroxene (olivine more than pyroxene) spectra are altered (darkened and reddened) which leads to an increased similarity between the spectra of the altered OC and of S-type asteroids. They also observed that the grains in the sample were coated with SMFe which may have been the cause of the alteration of the spectra. Brunetto and Strazzulla (2005) and Brunetto et al. (2007) simulated micrometeorite bombardment and solar wind by laser ablation of silicates and irradiation with different ions at different energies. The irradiation produced reddening and darkening of the spectra of the samples. The authors state that solar irradiation must be a very strong effect closer to the Sun, whilst at further distances micrometeorite bombardment seems to be the dominant agent of weathering.

Hapke (2001b) described several previous experiments to investigate space weathering of lunar samples and analysed the causes of formation of SMFe. Meteorite impact vitrification (meteorite impact effect), in which impact melts and vitrifies material, is one of the processes thought to be responsible production of SMFe. However, the results obtained by other authors do not reproduce lunar conditions (Pieters et al. 2000; Noble et al. 2001). Hapke was also unable to mimic space weathering, even using experimental conditions closer to those of the lunar environment (Hapke 2001a).

The source of ion irradiation (solar wind effects) is plasma of solar composition coming from the Sun at speeds of 500 km s$^{-1}$ with a flux of ions of mainly hydrogen and helium and some heavier ions (Brandt 1970). Experiments with irradiation of solid surfaces irradiated with H ions do not show any darkening or reddening of the spectra (Hapke 2001 and references therein). On the other hand, after irradiation of hydrogen on loose powders there was a strong darkening and reddening with temperatures too low to allow melting of the minerals (Hapke 2001 and references therein). The only samples that did not darken were those without Fe in their composition, which shows that Fe must be present in the first place (Hapke 2001
and references therein). Irradiation with helium also produces darkening in powders containing Fe, so inert gas sputtering also causes darkening (Hapke 2001 and references therein). The samples have to have a rough surface and porous texture in order for them to be darkened by sputtering. On experiments with alumina, the surfaces darkened were the sides and under the samples, not on the irradiated surface - samples must be of loose powders. Compacted powders will not show these results (Hapke 2001 and references therein).

The purpose of the work presented here was to simulate the alteration effects of micrometeorite impacts on asteroids, not by irradiation or by laser pulsed heating, but by shock. The first part of the study was to undertake impact experiments on terrestrial basalt to see whether any measurable effects were produced. If so, the procedure could be applied to meteorite samples, in the first instance, to those with identified parent bodies, in order to observe whether the impacts would change the spectra of the meteorites to spectra more similar to the assumed parent body.

Because of the few features present in the measured spectra of individual minerals by the MSP in Chapter 4, it was decided that a search for any effects of impact would be undertaken in the Mid Infrared (MIR), a region of spectrum known to contain features from the main minerals present in basalt. As described in Chapter 3, the instrument used was a FTIR spectrometer in the department of Atmospheric Physics at the University of Oxford.

6.2 The Light Gas Gun

One option for producing impact shock would be to use a Van de Graaff (VdG) to shoot particle into a rock. The particles must, however, be conductive and sub-micron sized in order to obtain hypervelocities, and the only instrument available at the time was one that shot a fine iron metal powder. Because it would be difficult to distinguish SMFe produced by shock reduction of silicates from fine-grained iron
from the projectile embedded in the sample, use of the VdG was ruled out. The instrument used for the impact experiment was a **Light Gas Gun (LGG)**, available at the Hypervelocity Impact laboratory (HVI) at the Open University. The LGG was the preferred instrument because it produces hypervelocity impacts into targets by firing any type of particle with a wide range of sizes – for micron-sized powders to millimetre size particles.

The LGG used in these experiments is shown in Figure 6.1 a) and b). This gun is one of only two in the world that can shoot into targets from different inclinations, allowing impact into liquids. The figures show one of the possible sample chambers at the end of the gun and the gun in a vertical position (Figure 6.1b). The sample chamber shown in the picture also allows a high speed camera to be attached at various angles.

![Figure 6.1: a) The Open University Light Gas gun with a medium sized chamber attached. The empty ports are connections for cameras or other detectors b) The LGG in a vertical position (images courtesy of Chris Hall) ![Figure 6.1: a) The Open University Light Gas gun with a medium sized chamber attached. The empty ports are connections for cameras or other detectors b) The LGG in a vertical position (images courtesy of Chris Hall)](image)

A schematic of the different sections of the LGG is shown in Figure 6.2 and Figure 6.3.
Figure 6.2: A schematic of the LGG showing the different components and chambers (courtesy of Chris Hall)

The firing mechanism is a 12 G (gauge) shotgun cartridge, specially loaded with up to 10 g of rifle powder. A variety of powders can be used which will influence the speed of the projectile. After the firing mechanism there is a piston with 2 'O' - rings
that seal the back of the pump tube; a 0.5 mm thick aluminium shear disk seals the other end. The pump tube is evacuated and filled with gas, the type and pressure of which will help control the speed of the projectile. Soft projectiles with 4.5 mm diameter may be fired from the rifle barrel directly or hard projectiles may be fired in a sabot (Figure 6.4). If a sabot is used, the maximum diameter of the projectile is 3.5 mm.

![Figure 6.4: A four piece sabot (a) filled with basalt powder and introduced in the LGG rifle barrel (b). The sabot is ~4mm diameter](image)

After the rifle barrel is the blast chamber. It contains adjustable vanes to control shock waves and any leaking gases. This chamber also contains the sabot stop plate (Figure 6.5) which catches all four pieces of the sabot whilst the projectile continues in a straight line through the 8 mm hole in the sabot stop plate.
Following the blast chamber in the range assembly is the time-of-flight chamber. It consists of two vertical ribbon lasers with detectors on opposite sides of the chamber. As the projectile passes through the first laser beam, causing an interruption of the signal on the detector, a timer starts and when the projectile passes through the second laser beam, the timer stops. Knowing the distance between the lasers and the time it took for the projectile to travel between them the speed of the projectile can be calculated. With the speed and mass of the impactor, the impact force can be determined.

The last chamber is the target chamber (Figure 6.6). Three different sizes of target chambers can be fitted. In this project, the smaller chamber was used as the samples were only 4x4 cm. Every component of the LGG beyond the rifle barrel is evacuated.
As mentioned above, different powder types and different gases at different pressures result in different velocities. In these experiments rifle powder of the type R19 was used with hydrogen gas. The pressure is 45 bar giving a speed of \(\sim 5 \text{ km s}^{-1}\). Because of the high pressure and velocities involved in this procedure, the LGG is locked within a steel door, the lab is evacuated and a remote trigger is used from outside the lab to fire the LGG (Figure 6.7).

Figure 6.6: The LGG target chamber. The target is attached to a plate at the end of the chamber

Figure 6.7: The target chamber locked on the steel door (the remaining components of the LGG are behind the steel door) (a) and the remote firing trigger (b)
6.3 Shots and samples

Two different types of shots were attempted on the LGG: one with micron-sized basalt powder to simulate micrometeorite impact and the second with mm-sized grains to simulate the effects of larger impacts. Several shots were made, some with no success either because the gun did not fire or because the projectiles did not hit the target. Two slabs of impacted basalt were eventually available for spectroscopic analysis. The sample names and the LGG shot settings are given in Table 6.1 and their images in Figure 6.8.

Table 6.1: LGG shot settings for the two targets used (1 Torr = 1 mmHg = 133.322 Pa)

<table>
<thead>
<tr>
<th>Sample</th>
<th>Projectile Type and Lead</th>
<th>Propellant Type and Pressure</th>
<th>Stop Disk</th>
<th>Vacuum in Range</th>
<th>Pump Tube Gas Pressure</th>
<th>Shot Velocity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Slab 1</td>
<td>1mm diameter 75-125 μm Basalt Powder</td>
<td>R19 10g Hydrogen 45 bar</td>
<td>0.5 cm Hard Aluminium</td>
<td>8.5 x 10^-1 ton</td>
<td>45 bar</td>
<td>3.68 km/s</td>
</tr>
<tr>
<td>Slab 2</td>
<td>1mm diameter basalt grain</td>
<td>R19 10g Hydrogen 45 bar</td>
<td>0.5 cm Hard Aluminium</td>
<td>8.0 x 10^2 ton</td>
<td>45 bar</td>
<td>1.39 km/s</td>
</tr>
</tbody>
</table>

Figure 6.8: Results of LGG impacts onto basalt slabs: Slab 1 (a) hit by 75-125 μm powder and Slab 2 (b) hit by a 1 mm basalt grain. The crater formed by the basalt grain impact can be seen at the top left. Each slab is ~ 4 x 4 cm

Because there was the possibility of the single grain projectiles not hitting the target, six 1 mm grains were fired into 4 slabs (sections of slab 2). As can be seen in the picture, two projectiles hit the target but one of them hit between the slabs. For the analysis, the crater on the top left (see Figure 6.8b) was chosen. This crater had an approximate diameter of 1 cm. The impact with the basalt powder formed a series of
mm-sized craters across an area of approximately 3 cm in diameter. An additional slab of basalt was prepared by scratching the surface with a second piece of basalt. The objective of this experiment was to roughen the surface of the slab, in order to determine whether changes in the spectra of the impacted basalts were caused by an increase in roughness, rather than other physical changes to the sample. This final sample will be referred to as 'scratched basalt'.

6.4 Spectroscopy results for the samples in the Mid Infrared

In this section, several results acquired with the FTIR spectrometer with the instrument described in Chapter 2, are presented. The positions of the identified features were found by hovering the cursor over the spectrum and noting where the first minimum was for the absorbance and Christiansen Features (CF), and the maxima for the Reststrahlen Bands (RB) and Transparency Features (TF) (see Chapter 2 for a description of CF, RB and TF).

Spectra of the meteorites Juvinas and Allende and the mineral orthoclase were taken to visualise some of the characteristics of their spectra and to help with the interpretation of data from the analysis of the basalt slabs. Allende is a carbonaceous chondrite, Juvinas is a eucrite, which has a basaltic origin, and orthoclase is a potassium feldspar, not often present in these meteorites. Results are plotted in Figure 6.9.
Figure 6.9: MIR spectra of the meteorites Juvinas and Allende and the mineral orthoclase taken on the Oxford FTIR. CF is the Christiansen feature in all the samples and TF the transparency feature in orthoclase

The MIR spectra of these samples have clear features that differ between the materials. Orthoclase, a terrestrial mineral sample, has a clear absorbance from water at 2.8 μm and overtone and combination bands (see Chapter 2) at around 6 μm. Above 6 μm, there is a sharp drop in reflectivity in the spectrum, forming the Christiansen Feature (CF) with a minimum at 7.5 μm. This sample of orthoclase has been ground to a grain size of < 125 μm. For this reason, and as explained in Chapter 2, the Reststrahlen Bands (RB) are weak and a broad transparency feature appears. It is difficult to determine the exact position of the TF because it is a broad feature but its maximum is at 11.4 μm.

The spectrum of the meteorite Juvinas (a eucrite) seems to have a small H₂O feature at 3 μm, but this might be caused by terrestrial contamination. There is a combination of other features between at ~5.5 and 7 μm which may be a mixture of water contamination and features from minerals such as pyroxene. The CF of the meteorite Juvinas is somewhat broad and difficult to measure, but its minimum
appears at 8.2 μm. This might be caused by the CF of different minerals that are present in the meteorites being far apart and broadening the CF of the meteorite. Another reason might be the fact that this sample is also a powder, causing the RB to become much weaker reducing the sharpness of the CF. The TF of Juvinas is at 12 μm.

Allende is a CV3 carbonaceous chondrite, and is known to have experienced slight amounts of aqueous alteration (e.g. Hutchison 2004); however, water bands cannot be distinguished in the spectrum. The fact that the sample is a fine powder should have increased the depth of any water bands relative to the continuum. The lack of water bands may be explained by the presence of organic (opaque) matter in Allende, which suppresses the water bands (Clark, 1983). C=O and C=C bands normally appear near 6.13 μm. A series of small bands can be seen in this area of the spectrum of Allende, but because there is interference from water bands and other minerals such as olivine and pyroxene in this region of the spectrum, the bands cannot be attributed to the presence of carbon (Salisbury et al. 1991). The CF of Allende is quite difficult to discern because the RB are almost undistinguishable (sample is a very fine powder < 50 μm). Its minimum is at 9 μm. The TF of Allende peaks at 12.4 μm.

The most obvious difference between the spectra of the meteorites and the mineral is the lack of water bands (which are present at ~ 3 and 6 μm in the orthoclase) in the meteorite samples. Even when they are present, they are small and are likely to be the result of terrestrial contamination. Orthoclase is a mineral with a degree of transparency which is reflected in the higher features in both the volume and surface part of the spectrum (refer to Chapter 2 for definitions). Allende is also the finer powder. Because the samples are powdered, RB are strongly reduced and the TF can be seen at wavelengths around 12 μm.
For comparison purposes, the different sizes of basalt powder (from the same sample as the impacted slabs) and a non-impacted area of a basalt slab (representing basalt as a solid rock) were also analysed on the FTIR at MIR wavelengths. The reasoning behind this was that differences in spectra following impact might have been caused by breaking the rock to powder, rather than changing the chemistry or structure of the minerals. The basalt was taken from the same piece as the target slabs, ground and sieved. Five different samples size-fractions were produced: 30 - 63 µm; 63 - 75 µm; 75 - 125 µm; 125 µm – 0.4 mm and a powder mix of grain sizes < 125 µm. Results can be seen in Figure 6.10.

![Figure 6.10: MIR spectra of basalt powder with different grain sizes and non-impacted basalt, taken on the Oxford FTIR](image)

The results are as expected, according to the effect of grain size (see Chapter 3 for explanation and references). The only samples with RB present are the coarser 125 µm to 0.4 mm and the solid rock sample. In the spectra from the finer grain sizes, the RB disappear and a TF appears at 12.1 µm, which is not present in the rock
sample or the coarser 125 μm to 0.4 mm fraction. The CF on the powdered samples remains at the same wavelength (7.9 μm) with different grain size which is also to be expected with grain sizes smaller than the wavelength. The exception is the sample with basalt grain from 125 μm to 0.4 mm. In this case, the CF is found at shorter wavelength (7.8 μm), an intermediate position between the CF of the powdered samples and the rock sample with a CF at 7.6 μm. In the volume dominated part of the spectrum (wavelengths < 8 μm), as particle size decreases, the reflectance increases resulting in higher peaks – the rock sample has the lowest reflection under the CF and the smaller powders the highest. All sample show water bands at 2.7 and 6 μm, although the latter might have the influence of some minerals such as olivine and pyroxene, present in basaltic rocks. At 4 - 7 μm, volume scattering dominates and the absorption bands are troughs instead of peaks. These increase with decreased particle size becoming diagnostic features (Salisbury et al. 1997). This can be seen in Figure 6.10 where the absorption bands of smaller sized fractions are more sharp and deeper relative to the continuum than the coarse samples or the rock sample. In the mixed powder, the results substantiate the fact that small particle sizes cling to larger ones influencing the spectrum. This can be seen by the fact that the mixed powder, even though it has particles of all sizes, behaves like a smaller particle sized sample.

The results presented here were compared with the results from Cooper et al. (2002) for validation. One of the samples of basalt was chosen (Basalt 1) and the positions of the CF and TF in his rock and powdered samples are shown in Table 6.2. Because of the slightly different results for the fraction 125 μm – 0.4 mm, those are also shown in the table. Different positions of the CF and TF between the samples of Cooper et al. (2002) are caused by the fact that the sample may have different compositions. The table shows that the difference between the CF of rock and powder in Cooper et al (2002) and the results of this work are approximate 0.24
µm for Cooper et al. (2002) and 0.3 µm for this work. The slight discrepancy might be, again, cause by difference in composition (in his work, Cooper et al (2002) finds that the changes in the positions of CF in powders and rocks vary according to the composition of the sample), but also because the method for finding the position of features of this work is different and less accurate.

Table 6.2: Comparison of the position of the CF and the TF between rock and powdered samples of basalt and the same results from the sample Basalt 1 from Cooper et al (2002)

<table>
<thead>
<tr>
<th>Cooper (2002) CF Powder Basalt 1</th>
<th>Cooper (2002) CF Rock Basalt 1</th>
<th>Difference between solid and powdered basalt</th>
<th>Our CF powder</th>
<th>Our CF sample 125 µm 0.4 mm</th>
<th>Difference between solid and powdered basalt</th>
<th>Cooper (2002) TF Powder Basalt 1</th>
<th>Our TF powder</th>
</tr>
</thead>
<tbody>
<tr>
<td>8.18 µm</td>
<td>7.89 µm</td>
<td>0.24 µm</td>
<td>7.9 µm</td>
<td>7.8 µm</td>
<td>0.3 µm</td>
<td>12 µm</td>
<td>12.1 µm</td>
</tr>
</tbody>
</table>

The purpose of this work is to analyse the effects of impact shock on the IR spectra of basalt samples. Spectra of Slab1 (impacted and non-impacted area) and the crater on Slab 2 are given in Figure 6.11.

![Figure 6.11: MIR spectra of Slab 1 (powder impact), Slab 2 (cratered) and a non-impacted area of the basalt taken on the Oxford FTIR](image_url)
There is a noticeable difference between the impacted area on Slab 1 and the non-impacted basalt. The CF has changed from 7.4 to 7.9 μm and a much broader feature from the non-impacted to the impacted spectrum. The water feature at ~2.8 μm was enhanced with the impact, the RB were suppressed and the peak was slightly shifted to longer wavelengths. This may be caused by 3 reasons:

• An alteration in composition resulting from the impacts. In this case the purpose of the experiment was accomplished, as the impact changed the composition of the basalt as it would do to an asteroid with a surface impacted by micrometeorites;

• A change in texture caused by the craters formed by the impacts, giving it a more porous surface more similar to a powder. As the surface is roughened surface scattering increases and the intensity of the RB diminishes (Salisbury et al. 1991);

• An alteration in composition cause by contamination deriving from the LGG.

If the alteration of the spectra is caused by the change in texture, then the scratched basalt should show a similar spectrum to the impacted basalt. The powdered basalt was also analysed to determine if the results are caused by the formation of a powdered, roughened surface on the impact area. To determine if the cause was contamination, a spectrum of the crater on Slab 2 was also taken - a larger projectile would have sufficient energy to vaporise the contaminants, and the alteration should not be present. Figure 6.12 shows the spectra of the non-impacted basalt, Slab 1, Slab 2, scratched basalt and powdered basalt for texture comparison.
Figure 6.12: MIR spectra of Slab 1 (powder impacted), Slab 2 (cratered), scratched basalt, non-impacted basalt, and powdered mix basalt taken on the Oxford FTIR

The more apparent differences between spectra are as follows: TF are only present in powders. It makes sense that only the basalt powder shows this feature at around 12 µm. The RB were suppressed in the basalt powder and completely disappear, giving rise to the TF. The non-impacted basalt, Slab 1 and scratched basalt were averaged from two different scans in order to eliminate any crystal direction effect on the spectra. On a first approach to analysing the results, the scratched basalt seems very different from the impacted basalt but that may be related to the fact that both the scratched scans that were averaged were very different in the first place, possibly because of a direction effect and the fact that the slab was tilted. They should be analysed independently. In the impacted basalt, the RB are suppressed, as would be expected for a surface altered by impact, and also from a smaller grain size effect. In the spectrum for the cratered sample, the RB are also very small; this was not expected. A small roughness effect was expected but not such a great difference. Maybe there is scattering against the walls of the crater and
insufficient signal reached the detector, or maybe there is a deposition of smaller particles on the bottom of the crater.

In order to better understand the differences between the samples, all the spectra were normalized at 7 μm and the two scratched basalt spectra were plotted separately (Figure 6.13). Table 6.3 contains a description and positions of the main features present in the spectra from Figure 6.12 and Figure 6.13.

![Figure 6.13: MIR spectra of Slab 1, Slab 2, non-impacted basalt, scratched basalt in two different directions powdered basalt and basalt grains between 125 μm and 0.4 mm, all normalized to 7 μm](image)

Figure 6.13: MIR spectra of Slab 1, Slab 2, non-impacted basalt, scratched basalt in two different directions powdered basalt and basalt grains between 125 μm and 0.4 mm, all normalized to 7 μm
Table 6.3: Description and position of the main features of the spectra of the samples shown in Figure 6.19

<table>
<thead>
<tr>
<th>Sample</th>
<th>Water feature I (2.7 µm)</th>
<th>Reflectance between water features (2.7 - 6 µm)</th>
<th>Water feature II (6 µm)</th>
<th>CF</th>
<th>RB</th>
<th>TF</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-impacted basalt</td>
<td>Weakest in all the samples</td>
<td>Lowest of all the samples</td>
<td>Very weak together with cratered</td>
<td>Broad at −7.5 µm</td>
<td>High reflectivity with spectra definition, Peaks at 8.1, 8.9, 9.7, 10.2, 11 µm</td>
<td>N/A</td>
</tr>
<tr>
<td>Scratched basalt direction A</td>
<td>Deep</td>
<td>High reflectance but lower than powder, 125 µm-0.4 mm and scratched direction B and non impacted</td>
<td>Deep</td>
<td>−7.6 µm</td>
<td>High reflectivity and enhanced peaks described above</td>
<td>N/A</td>
</tr>
<tr>
<td>Scratched basalt direction B</td>
<td>Very deep</td>
<td>Highest reflectance after 125 µm-0.4 mm</td>
<td>Deep</td>
<td>−7.5 µm very broad</td>
<td>High reflectivity and enhanced peaks described above (less than direction A)</td>
<td>N/A</td>
</tr>
<tr>
<td>Impacted</td>
<td>Deep</td>
<td>Highest reflectance after the powdered sample</td>
<td>Deep</td>
<td>Broad at −7.9 µm</td>
<td>Reflectivity decreases, peaks are not distinguishable, peak at 11 µm becomes dominant</td>
<td>N/A</td>
</tr>
<tr>
<td>Cratered</td>
<td>Deep</td>
<td>Similar to non-impacted</td>
<td>Deep</td>
<td>Very broad at −7.5 µm</td>
<td>Much weaker and broader RB. Peaks not distinguishable</td>
<td>N/A</td>
</tr>
<tr>
<td>125 µm to 0.4 mm</td>
<td>Very deep (deepest)</td>
<td>Very high (highest)</td>
<td>Very deep</td>
<td>Very deep at 7.5 µm</td>
<td>Very high (highest) reflectivity but peaks are not distinguishable</td>
<td>N/A</td>
</tr>
<tr>
<td>Powder</td>
<td>Very deep</td>
<td>Higher reflectivity of the samples</td>
<td>Very deep</td>
<td>7.9 µm</td>
<td>None</td>
<td>12 µm</td>
</tr>
</tbody>
</table>

From the normalised data, it can be seen that H₂O bands increase in depth from the non-impacted basalt, impacted, cratered, powder and 125 µm to 0.4 mm. It is expected that smaller grain sizes increase the depth of the water bands and in this case this seems to happen perhaps because of the roughening of the surfaces with the impacts. The scratched basalt shows very different behaviour from one scan to the other, and, again, this may be cause by the different scratching direction and because the slab was not horizontal. It has an inclination on the surface, which caused a difference in reflectance. The 125 µm to 0.4 mm powder behaviour remains unexplained. The grains were quite coarse, shiny and loose so the porosity effect seems to be stronger showing a much higher reflectance at the wavelengths where scattering domains. It almost behaves like a metal. The CF on this sample is also the strongest and has similar shape spectrum to the scratched spectra.

The reflectance in the volume scattering wavelengths between the water bands at 2.7 and 6.1 µm decreases in the order 125 µm - 0.4 mm powder > scratched basalt direction B > powdered > scratched basalt direction A > cratered > impacted > non-
impacted. This behaviour is consistent with the behaviour of the water absorption bands described above.

CF of all the samples do not vary much from the non-impacted basalt. The exceptions are the impacted basalt and the powder. Powdered samples of the same composition have CF at longer wavelengths as powders than as solid rocks (e.g. Cooper 2002 and Salisbury et al. 1997). Once again there is the possibility that the surface of the slab has been roughened by the impact and behaves similarly to a powder. An increase in the ratio between the amount of pyroxene and olivine and plagioclase would also shifts the CF to longer wavelengths. It is possible that with the impact, the less resistant plagioclase was removed - the ratio between Fe and (Al plus K) would also be higher – resulting in a change in composition to more Fe-rich. Another explanation might be the effect of shock pressure inducing changes in the structure of the minerals, rather than composition. This possibility is discussed below.

The presence of nanophase Fe would have the same effects on the spectra, however, according to some authors, e.g., Pieters (2000) and Hapke (2001b), for the formation of nanophase Fe, reduction through sputtering by irradiation or H⁺ ions from the solar wind is also needed and this experiment does not include a simulation of that effect. It is, therefore, unlikely that the change in spectra is caused by the formation of nanophase Fe. The next section will include SEM results on the composition of the cratered and non-impacted areas in an attempt to investigate if there are indeed changes in composition with the impacts.

The scratched slabs look quite similar to each other at the RB area, only with a difference in brightness. The positions of the RB are the same between them. They are, however, slightly different from the remaining basalt slabs, impacted or not. Most RB are in the same position but their intensities are different depending on wavelength. They seem to decrease equally between the scratched basalt but are
higher at shorter wavelengths compared to longer wavelengths. The changes from
the non-impacted basalt to the impacted and cratered are not only a reduction in
intensities, but at shorter wavelengths, closer to 8 µm, the decrease is more
noticeable. The RB become asymmetric with a higher peak at 11.5 µm.

In the powdered samples, there seems to be the formation of a TF at 12 µm as
described above. This is consistent with the presence of feldspar dominating the
spectrum as it normally exhibits a TF at this wavelength. The cause of this might be
that harder minerals resist the grinding process better, being retained on the sieves
with a larger mesh, and the less hard like feldspars are broken and accumulated in
the smaller fractions (Salisbury et al. 1997). This selection process will also be
discussed in the next section.

Because they seem to be quite similar, the spectra of non-impacted basalt,
scratched basalt and grain size 125 µm to 0.4 mm were plotted on Figure 6.14 for
better comparison. The results were normalised at 7 µm.
Figure 6.14: MIR spectra of non-impacted basalt, scratched basalt (in two directions) and 125 μm to 0.4 mm

The sample with grain size 125 μm to 0.4 mm is very similar to the scratched basalt. This is expected as it represents an increase in roughness of the surface. There are no other major changes, other than the intensity of the RB, which indicates that there are no differences in composition.

The effects of shock on basalts, including the resulting changes in their spectra, have been studied by several authors. Kieffer et al. (1976) studied the effects of impact shock on basalts from an impact crater and compared them with samples shocked in laboratory conditions using a high velocity gun under vacuum conditions. Wright et al. (2011) further studied the effects of impact shock on the Thermal Infrared Spectra (TIR) of basalts and summarised the different classes of effects described in Kieffer et al. (1976) into a table (Table 6.4) The following discussion is
based on the work of Kieffer et al. (1976), Johnson et al. (2007), Wright et al. (2011) and references therein:

Table 6.4: Classes of shocked basalt and summary of petrography (Wright et al., 2011 adapted from Kieffer et al., 1976).

<table>
<thead>
<tr>
<th>Class</th>
<th>Shock Pressure (GPa)</th>
<th>Brief Petrographic Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>&lt;20</td>
<td>Grains are fractured</td>
</tr>
<tr>
<td>2</td>
<td>20-40</td>
<td>Plagioclase transformed to maskelynite</td>
</tr>
<tr>
<td>3</td>
<td>40-60</td>
<td>Plagioclase transformed to glass of plagioclase composition</td>
</tr>
<tr>
<td>4</td>
<td>60-80</td>
<td>Plagioclase converted to vesiculated glass; pyroxene melted on crystal edges and/or heavily fractured</td>
</tr>
<tr>
<td>5</td>
<td>&gt;80</td>
<td>Plagioclase and pyroxene completely melted</td>
</tr>
</tbody>
</table>

The spectra of un-shocked basalt taken by Johnson et al (2007) and Wright et al (2011) show similarities with the spectrum from this work, with main features at 9 and 11 μm, resulting from Si-O vibrations in Ca-rich plagioclase, plus contributions from olivine and pyroxene. Their results show a shift in the CF to longer wavelengths with increasing pressure, which also happens here (in particular with the impacted basalt, see Table 6.3). Wright et al. (2011) compared the Thermal Infrared spectra of class 2 shocked (20 – 40 GPa) and un-shocked basalts, with particular attention to the features of labradorite (a type of plagioclase), which turns to maskelynite (a glass of plagioclase chemistry, but not crystallography). In type 2 shocked basalt, the features of labradorite (CF at 8.1 μm, followed by a sharp slope, absorption at 9 μm and emission at 10 μm) seem to disappear.

In the results from our work, the CF seems to be influenced by the presence of other minerals, however a shift to longer wavelength is especially clear in the impacted basalt sample (Figure 6.19 and Table 6.3). The slope after the CF also appears flatter and the sharp feature at 9 μm is reduced, as is the broader feature above 10 μm. The entire spectrum is moving closer in appearance to that of the basalt powder – as discussed earlier, this may be because the surface has taken on the more porous texture of a powder. The 10 μm 'hump' is also reduced in intensity
in the cratered basalt; if this feature is, in part, caused by the presence of plagioclase, then it may be that we are seeing the effect of the plagioclase to maskelynite transition in the impacted and cratered samples. At this stage, it is not possible to confirm the definite presence of maskelynite without further tests, although it appears that melting has not occurred because of the absence of a sharp 9.5 μm Si-O feature that occurs in glasses.

6.5 SEM analysis results from cratered and non-impacted area

To complement the investigation of the effect of impacts by shock on the mineralogy of a sample, composition maps were taken by the FIBSEM, in the Department of Physical Sciences of the OU, from an area inside the crater on Slab 2 and a non-impacted area outside but close to the crater. The reason why the cratered sample rather than the impacted sample was chosen resides in the characteristics of the SEM. It required a relatively flat area for the mapping and the impacted area was too rough. Instead, a flattened area inside the crater was used in this investigation. Figure 6.15 shows the composition maps of an area outside the crater and one inside the crater of Slab 2. Because there is great interest in ascertaining if there are differences in the quantities of Fe after impact, this was one of the elements for which data were acquired. One of the hypotheses for a change in compositions might also be that the impact, in the same way as grinding, destroys the less resistant minerals. The less resistant mineral that is present in sufficient quantities for a clear result is plagioclase. For this reason, data for concentration of Ca were also taken.
It can be seen in Figure 6.15 that there is a major increase in the amount of Fe inside the crater at the same time as there is a decrease in Ca. Evidence for the possibility of contamination from the LGG is not seen. If the contaminant is for Fe it would not affect the results of Fe and Ca and if it was Fe, it would be visible in areas outside the crater as well.
These results are consistent with the possibility that, with a high speed impact, even when the impactor is small, generates enough energy to vaporise less resistant minerals. Besides the presence of SMFe, an alteration in the ratios of certain minerals at the surface of an asteroid cause by an impact as seen in this chapter, would modify the spectrum of that asteroid in variable amounts which would make the identification of this asteroid as a parent body of a meteorite more difficult.

6.6 Conclusions

The purpose of this work was to test an impact technique that would help analyse the effects of impact shock on basalt samples. If successful, meteorite samples could be used with the purpose of aiding the establishment of links between meteorites and asteroid parent bodies.

There is some (tentative) evidence provided by spectroscopy that supports the possibility that impact changes the sample composition, either the alteration of plagioclase to maskelynite, or by vaporization of less resistant materials. The results for the FIBSEM further sustain this possibility. However there are other effects that may have given rise to the differences between impacted and non-impacted samples, such as surface roughness or contamination. In order to draw more firm conclusions about whether impact experiments provide a useful analogue for space weathering processes on asteroids, additional experiments need to be carried out. Only a limited number of samples was examined in these experiments, and results should be regarded carefully. Additional tests with basalts and other terrestrial rocks should be done, as well as with other types of meteorite.

The light gas gun is a good technique for the tests although it is vulnerable to contamination and it requires relatively large sample which might not be possible with meteorites. Another option for the study of the effects of impact shock is to use a Van de Graaff (VdG). The VdG has the advantage of requiring very small samples
as a target and it has the possibility of bombarding the target continuously for quite a long time (days or even weeks), as opposed to the LGG which shoots one projectile at a time. On the other hand, the VdG does not reach velocities as high as the LGG. It does not generally reach velocities above 3 kms$^{-1}$ and it requires the projectiles to be conductive. Fe particles are generally used but these would strongly interfere with the amount of Fe found in the target, which is one of the results of high speed impacts (nanophase iron). Copper would also affect the results in the same way. Some work has been done by Hillier et al. (2001) to develop conducting silicate particles that can be accelerated in the VdG and therefore can provide more reliable results.
7 Conclusions and Further work

The technique of UV-Vis microspectroscopy (MSP) is one that has many applications in pharmaceutical and forensic investigations, where samples are mostly analysed using transmission mode, i.e., spectral signatures are characterised following transmission of radiation through a sample. UV-Vis reflectance spectroscopy, where a powdered sample is analysed in bulk, using a target of around 1 mm in diameter, is also a commonly-found technique, and has special application for comparison with remotely-obtained spectra from asteroids.

The aim of this study was to apply the micro technique in reflectance mode, to establish whether the MSP was a suitable technique for UV-Vis spectroscopy of extra-terrestrial materials, specifically samples that are too small for analysis by the bulk technique, i.e., cometary grains. If the technique was found to be appropriate, by analysis of minerals and meteorites, and comparison of results with those obtained from conventional spectroscopy, it would then be applied to study of cometary samples from comet 81P/Wild2 collected by the Stardust spacecraft. Part of the investigation process involved study of grains from HED meteorites, comparing spectra with those from their potential parent body, Vesta. In addition, a study of the effects of shock impact was carried out, using a light gas gun (LGG) on basalts, as a starting point to the determination of whether or not this was a useful analogue for space weathering of asteroid surfaces.

7.1 Operation of the MSP

In the first chapter of the thesis, I describe the type of extra-terrestrial materials available for study, and the formation processes that they have experienced. In Chapter 2, I give a broad introduction to spectroscopy, particularly electronic spectroscopy, without entering into a detailed discussion of the atomic processes that cause the spectra (which was thought to be beyond the scope of the thesis).
Chapter 3 is an extensive analysis of the performance and operation of the MSP. The tests performed included the characterization of the dark current, the stability of the light source, the sensitivity of the instrument at all wavelengths and homogeneity of the diffuse white reflectance standard provided by the manufacturer. I found several problems with the system, all of which would impact on collection and interpretation of data:

**Light source:**

I found by investigating the drift of the dark scan and reference scans with time that the system only stabilised after the light source (a xenon lamp) had been switched on for ~ 4 hours. This finding had a significant effect on the way in which I then had to carry out my experiments. Figure 3.9 also shows that there are emission lines between 450 and 500 nm and above 700 nm that might be related to the light source, and which might influence interpretation of the results if data were improperly corrected.

**Reference homogeneity:**

The reference standard is only suitable for wavelengths above ~ 250 - 350 nm (Figure 3.10) and, even at the lowest of magnifications, is very heterogeneous, which causes great variation in the reference scan used for the corrections of the spectra. For this reason, it is advised that 20 reference scans should be taken in order to minimise the effects of heterogeneity in the results. For this to be possible, the spectrum of a sample has to be taken as a reference scan and the corrections have to be carried out in a spreadsheet, rather than using the software supplied by the manufacturer.

**Focussing:**

The optics of the microscope rely on mirrors, rather than lenses, and so the optical image of a sample is very poor. This led to problems with the correct focussing of
samples, even over the small area of the aperture (generally 10 x 10 μm, but 2 x 2 μm for the Stardust grains). Results of tests made on the reference scan show that there is a change in spectra with different focussing, which might be related to an illumination problem, or to the geometry of the instrument which might be capturing reflected light from sources other than the sample.

7.2 Application of the technique to minerals commonly found in meteorites

A series of scans was taken on minerals commonly found in meteorites in order to ascertain, in the first instance, if the instrument were suitable for the study of minerals, rocks and meteorites (Chapter 3). The results were also compared, when possible, with results from other authors, either taken by the RELAB spectrometer or the UNIWIN instrument (Cloutis et al. 2008).

Results varied greatly with the samples studied, even from grain to grain. It was found that the ability to focus the MSP correctly on the sample significantly influenced the spectra. If the surface of a grain was rough, such that it was not possible to bring to a single focus within the area of the aperture, the resulting spectrum was essentially featureless and flat, which might be evidence for an illumination issue.

Another concern was the size of the sample relative to the field of view. At the 10x magnification that was used for most of the observations, the aperture of the spectrometer was 10 x 10 μm. It had been assumed that as long as the grains being analysed were larger than the aperture, there would be no influence on the spectra from the surrounding area, and that results would be valid. What was found, however, was that if the sample under consideration did not fill the whole field of view of the spectrometer, then light scattered into the detector from the substrate supporting the sample could have a significant effect on the resulting spectrum. In
the samples analysed in Chapter 3, the influence is from the quartz slide on which
the mineral grains were supported. Quartz is transparent to UV light, so it was
assumed that any reflection from the slide would be minimal and would not interfere
with the sample analysis. What was found, though, was that the spectrometer did
pick up a signal from the quartz, possibly from specular reflection from the flat
surface of the slide. This had an effect of ‘diluting’ the signal from the sample, giving
a lower signal to noise ratio.

It was also observed that, when the grains are very dark, the spectrum is flat and
featureless. This occurs even when, with the naked eye, the sample is almost white.
This was evident in the case of olivine (see Appendix A). It seems like a
combination of the fact that the grains are extremely small with the inability to
correctly focus the microscope on the sample and the influence of the quartz slide
are the cause for the featureless, very dark spectra.

When comparing data from different instruments, it was noticeable that results
varied from instrument to instrument. The geometry of the instrument was less
significant in influencing the final spectrum than the size and shape of the sample. It
was apparent that if the grains were flat (and so bright and well-focussed) and
covered most of the field of view, data from the MSP were comparable to those
taken at the RELAB facilities and UNIWIN.

7.3 Application to Stardust samples and meteorites

Once the MSP had been characterised, and the most appropriate experimental
conditions had been determined, the technique could be applied to meteorites and
cometary dust. Chapter 5 gives the results from analysis by the MSP of the UV-Vis
spectra of two grains collected from comet Wild 2 by the Stardust mission.
Unfortunately the grains were covered in aerogel which made the analysis difficult.
Furthermore, as predicted by the results from Chapter 4, the gold foil into which the
grains were pressed, greatly influenced the spectra: even at the highest magnification of the MSP, the grains did not fill the field of view of the microscope, and so the signal observed was dominated by reflectance from the gold foil. Corrections were made in an attempt to minimise the effect of the aerogel and the gold foil, but these are only qualitative, since we could not measure the thickness of the aerogel, and could not be sure how much light was scattered into the MSP from the gold foil. Despite this, results indicated that the two samples analysed were assemblages of low- and high-calcium pyroxene and olivine, possibly with some magnetite, and were not single mineral grains.

There has only been one previously published analysis of extra-terrestrial materials by MSP, and that was a study which compared spectra from the Allende carbonaceous chondrite with spectra from IDPs (Bradley et al. 1996). This comparison was also included in Chapter 4: spectra from the Murchison and Allende carbonaceous chondrites were recorded and compared with the published data, and also with the Stardust grains. The chondrites were very dark to look at, and produced dark, flat and featureless spectra, similar to those recorded by Bradley et al. 1996), who concluded that the spectra were influenced by the presence of carbonaceous material in the samples.

Chapter 5 however shows that the technique, when the conditions described above are met, can be applied in the field of planetary sciences. Spectra from three HED meteorites are compared with results from RELAB and also spectra taken of their possible parent body (asteroid 4 Vesta). The spectra show a great similarity between results from other laboratories, and also from the asteroid, confirming the utility of the MSP under the correct circumstances.
7.4 Investigation of the effect of impacts by shock in the spectra of basalt

Space weathering is a possible cause for the discrepancies between the spectra of meteorites and their possible parent bodies (asteroids) (e.g. Clark et al, 2002) and one of the main factors of space weathering is the impact of micrometeorites on the surface of the asteroid. Previous work attempted to duplicate the effects of space weathering by laser irradiation (Yamada et al. 1999 and Sasaky et al. 2001) and ablation (Brunetto and Strazzulla 2005 and Brunetto et al. 2007). In this work, the effects of impacts by shock were investigated, using a Light Gas Gun (LGG) to shoot powders or grain of basalt onto basalt. Because, at the time, it looked as if it would be difficult to obtain useful information from spectra recorded by the MSP for comparison of impacted and non-impacted materials, spectra were recorded at a longer wavelength, using an FTIR system at the University of Oxford (Chapter 6). Results show that there is a difference between the spectra of impacted and non-impacted areas, although the reasons are not clear. Reasons for this difference include the effect of roughening of the surface by the impact, where the impacted surface behaved as a fine-grained sample; contamination by residue in the LGG; the formation of SMFe by the impact observed by other authors (e.g. Pieters et al. 1993); or a change in the mineralogy within the impacted area. SEM results support the last-mentioned effect, with a high increase in the concentration of Fe inside the impacted area and a depletion of Ca. These results are consistent with the hypothesis that the impact vaporised the minerals that are less resistant, such as plagioclase, compared with ferromagnesian minerals. This effect may also occur at the surface of the asteroids, causing a change in the spectra of the body. Only a few samples were used in this study, not enough to reach any definite conclusions. Further experiments must be conducted.
7.5 Future work

Results from the study of minerals and meteorites have shown that the MSP can be used to record spectra from individual mineral grains and from very small amounts of powdered meteorite, but that the experimental conditions under which valid data can be acquired are very specific. Currently, results can only be interpreted by comparison of the data with results from similar samples that have been measured on other instruments. If the MSP is to be a useful analytical tool, then a more systematic study of minerals and mixtures of minerals must be carried out. It would also be very useful if experiments were made using mixtures of different minerals with different ratios to determine the influence of composition on the results and complement the investigation with the analysis of the spectra of powders and slabs of rocks. The analysis of spectra from the HED meteorites was encouraging, however, other meteorite types should also be analysed and their spectra compared to those of asteroids considered possible parent bodies.

The effect of impact shock on the spectra of basalt slabs as described in Chapter 6 revealed some interesting results with some form of composition change on the impacted areas. Nevertheless only two samples were analysed, which is insufficient for more concrete conclusions. More impacts using the LGG should be made using different types of targets but, ideally, a VDG should also be used on the same samples and for varied periods of time if non-metal projectiles can be used.

In summary, the investigations described in this work are of value for planetary sciences and set a solid stepping stone on the use of new techniques in the field, especially for the analysis of samples that are too small to be analysed by more conventional techniques. However, further investigations should be made and the method should be applied to a greater range of samples.
References


Spectral Range 44,000-4,000 Cm$^{-1}$.


Appendix A - Database of Mineral Compositions and UV-Vis Spectra

A selection of the most common minerals in meteorites was collected and used in this work. Table A1 lists the minerals studied; their compositions, as measured by SEM, are given in the relevant sub-sections of the results section.

Table A1: Minerals studied by UV-Vis Microspectroscopy

<table>
<thead>
<tr>
<th>Mineral Group</th>
<th>Name</th>
</tr>
</thead>
<tbody>
<tr>
<td>Silicates:</td>
<td></td>
</tr>
<tr>
<td>Olivine</td>
<td>Fo₀ (Fayalite)</td>
</tr>
<tr>
<td></td>
<td>Fo₃₀</td>
</tr>
<tr>
<td></td>
<td>Fo₅₀</td>
</tr>
<tr>
<td></td>
<td>Fo₇₀</td>
</tr>
<tr>
<td></td>
<td>Fo₁₀₀ (Forsterite)</td>
</tr>
<tr>
<td>Pyroxene</td>
<td>Rhodonite</td>
</tr>
<tr>
<td></td>
<td>Wollastonite</td>
</tr>
<tr>
<td></td>
<td>Diopside</td>
</tr>
<tr>
<td></td>
<td>Enstatite</td>
</tr>
<tr>
<td></td>
<td>Augite</td>
</tr>
<tr>
<td>Feldspar</td>
<td>Albite</td>
</tr>
<tr>
<td></td>
<td>Nepheline (feldspathoid)</td>
</tr>
<tr>
<td></td>
<td>Anorthite</td>
</tr>
<tr>
<td></td>
<td>Orthoclase</td>
</tr>
<tr>
<td>Phosphates</td>
<td>Apatite</td>
</tr>
<tr>
<td>Sulphates</td>
<td>Pyrrhotite</td>
</tr>
<tr>
<td>Oxides:</td>
<td>Spinel group</td>
</tr>
<tr>
<td></td>
<td>Spinel</td>
</tr>
<tr>
<td></td>
<td>Magnetite</td>
</tr>
<tr>
<td></td>
<td>Hibonite</td>
</tr>
<tr>
<td>Carbonates</td>
<td>Dolomite</td>
</tr>
<tr>
<td></td>
<td>Aragonite</td>
</tr>
<tr>
<td></td>
<td>Siderite</td>
</tr>
<tr>
<td></td>
<td>Magnesite</td>
</tr>
<tr>
<td>Phyllosilicates</td>
<td>Montmorillonite</td>
</tr>
<tr>
<td></td>
<td>Saponite</td>
</tr>
<tr>
<td></td>
<td>Lizardite</td>
</tr>
</tbody>
</table>

Table A2 is a compilation of the most obvious spectral features present in the minerals analysed in this study, with the transitions that they have been assigned to be based on crystal field theory (e.g. Burns 1970).
Table A2: Summary of UV-Vis results from analysis of minerals by the OU MSP

<table>
<thead>
<tr>
<th>Mineral</th>
<th>Feature</th>
<th>Possible Assignation</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Olivines</strong></td>
<td>320 nm</td>
<td>unknown</td>
</tr>
<tr>
<td><strong>Pyroxenes</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Enstatite</td>
<td>300-340 nm</td>
<td>Fe-O charge transfer (CT) Fe₂⁺</td>
</tr>
<tr>
<td>Augite</td>
<td>270, 295, 300 nm</td>
<td>Unknown</td>
</tr>
<tr>
<td></td>
<td>520, 620, 680 nm</td>
<td>Fe-O charge transfer (CT)</td>
</tr>
<tr>
<td>Diopside</td>
<td>330 nm</td>
<td>Unknown</td>
</tr>
<tr>
<td></td>
<td>600-700 nm</td>
<td>Ti⁴⁺ or Fe-O</td>
</tr>
<tr>
<td>Wollastonite</td>
<td>300 nm</td>
<td>unknown</td>
</tr>
<tr>
<td></td>
<td>670, 720, 780 nm</td>
<td>Unknown (system artefact?)</td>
</tr>
<tr>
<td>Rhodonite</td>
<td>300 nm</td>
<td>Mn³⁻-O CT</td>
</tr>
<tr>
<td></td>
<td>410 nm</td>
<td>Mn³⁺ Oct Coord. M1 site</td>
</tr>
<tr>
<td></td>
<td>550 nm</td>
<td>Spin Forb. Fe²⁺ M1 site</td>
</tr>
<tr>
<td></td>
<td>700 nm</td>
<td>unknown</td>
</tr>
<tr>
<td><strong>Feldspars</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Nepheline</td>
<td>280 nm</td>
<td>Fe²⁺-O CT</td>
</tr>
<tr>
<td>Anorthite</td>
<td>270 nm</td>
<td>Fe²⁺-O CT</td>
</tr>
<tr>
<td>Albite</td>
<td>400-500, 700 nm</td>
<td>unknown</td>
</tr>
<tr>
<td><strong>Oxides</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spinel</td>
<td>270 nm</td>
<td>unknown</td>
</tr>
<tr>
<td></td>
<td>320, 550-700 nm</td>
<td>unknown</td>
</tr>
<tr>
<td>Magnetite</td>
<td>270 nm</td>
<td>Fe²⁺-O CT</td>
</tr>
<tr>
<td></td>
<td>305, 350, 620-640 nm</td>
<td>unknown</td>
</tr>
<tr>
<td>Hibonite</td>
<td>270 nm</td>
<td>Fe²⁺-O CT</td>
</tr>
<tr>
<td></td>
<td>340 nm</td>
<td>Ti⁴⁺/Fe²⁺-O CT</td>
</tr>
<tr>
<td>Pyrrhotite</td>
<td>270 nm</td>
<td>Fe²⁺-O CT</td>
</tr>
<tr>
<td><strong>Carbonates</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aragonite</td>
<td>270, 490 nm</td>
<td>unknown</td>
</tr>
<tr>
<td>Dolomite</td>
<td>270, 300, 340 nm</td>
<td>Ti⁴⁺/Fe²⁺-O CT</td>
</tr>
<tr>
<td>Magnesite, Siderite</td>
<td>270, 300, 340 nm</td>
<td>Tetrah Fe³⁺/Octah Fe³⁺ CT</td>
</tr>
<tr>
<td><strong>Phyllosilicates</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lizardite</td>
<td>270 nm</td>
<td>unknown</td>
</tr>
<tr>
<td>Montmorillonite</td>
<td>350 nm</td>
<td>unknown</td>
</tr>
<tr>
<td></td>
<td>260 nm</td>
<td>Fe²⁺</td>
</tr>
<tr>
<td>Saponite</td>
<td>300 nm</td>
<td>Octah Fe</td>
</tr>
<tr>
<td></td>
<td>500, 620 nm</td>
<td>Fe³⁺/Fe²⁺ CT</td>
</tr>
</tbody>
</table>
Silicates

Olivines - \((\text{Ca, Mg, Fe})_2\text{SiO}_4\)

Figure A1: Image taken by the MSP of an agglomerate of grains of Fo0. The black square in the middle is the aperture of the spectrometer and is 10x10 μm in size.

a)
Figure A2: Visible spectra of synthetic olivines of known composition a) spectra from 250 to 800 nm; b) same spectra from 250 to 500 nm; c) same spectra from 500 to 800 nm. Spectra were averaged from spectra taken in different locations on the sample.
Figure A3: Visible spectra of synthetic olivines Fo0, Fo30, Fo50, Fo70 and Fo100 taken on the MSP compared with data from the same samples taken by RELAB (note, though, for the most magnesian samples, the OU data are for Fo100, whilst those for RELAB are for Fo90). The OU spectra (thicker lines) were averaged from spectra taken in different locations on the sample.
Figure A4: Visible spectra of synthetic olivines Fo0, Fo30, Fo50, Fo70 and Fo100 taken on the MSP compared with data from the same samples taken at UNIWIN by Cloutis (2008) (note, though, UNIWIN samples are labelled Ol x(sample number) Fa.y(composition). The compositions are not exactly the same between the two sets of data because no equivalent to the OU compositions was found on the UNIWIN database. Data downloaded from http://psf.uwinnipeg.ca/Sample Database.html
Figure A5: Reflectance spectra of a series of natural olivine grains, comparing data taken by the RELAB system with those acquired at UNIWIN (Cloutis et al., 2008). Data downloaded from http://psf.uwinnipeg.ca/Sample_Database.html

Table A.3: Composition of the pyroxenes analysed in this chapter. The composition analysis was performed using a Scanning Electron Microscope

<table>
<thead>
<tr>
<th></th>
<th>F (wt %)</th>
<th>Na (wt %)</th>
<th>Mg (wt %)</th>
<th>Al (wt %)</th>
<th>Si (wt %)</th>
<th>Ca (wt %)</th>
<th>Mn (wt %)</th>
<th>Fe (wt %)</th>
<th>O (wt %)</th>
<th>Ti (wt %)</th>
<th>Cr (wt %)</th>
<th>B (wt %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wollastonite</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>24</td>
<td>34.8</td>
<td>-</td>
<td>-</td>
<td>41.2</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Diopside</td>
<td>0.4</td>
<td>7.2</td>
<td>0.3</td>
<td>24.7</td>
<td>17.1</td>
<td>trac</td>
<td>7.9</td>
<td>42.4</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Enstatite</td>
<td>-</td>
<td>18.4</td>
<td>0.3</td>
<td>26.2</td>
<td>0.2</td>
<td>-</td>
<td>9.5</td>
<td>45</td>
<td>0.4</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Augite</td>
<td>0.6</td>
<td>10</td>
<td>3</td>
<td>20.1</td>
<td>8.3</td>
<td>-</td>
<td>2.8</td>
<td>54.0</td>
<td>0.45</td>
<td>0.21</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>
Figure A6: Images taken by the MSP of grains of pyroxenes. a) diopside, b) enstatite, c) augite, d) rhodonite. The black square in the middle is the aperture of the spectrometer and is 10 x 10 μm in size.
Figure A7: Visible spectra of pyroxenes a) from 250 to 800 nm; b) from 250 to 500 nm; c) from 500 to 800 nm. Spectra were averaged from spectra taken in different locations on the sample.
Figure A8: Spectra of a) the pyroxene augite taken by the MSP and the spectra of augite of different grain sizes taken in the RELAB facilities; b) the pyroxene enstatite taken by the MSP and the spectra of augite of different grain sizes taken in the RELAB facilities. Spectra of enstatite are offset for clarity. Exact composition of the minerals used by RELAB is unknown. Spectra taken on the MSP were averaged from spectra taken in different locations on the sample.
Pyroxenes – (Ca, Mg, Fe)SiO$_3$

Figure A9: Images taken by the MSP of four grains of diopside a) diopside 1, b) diopside 2, c) diopside 3, d) diopside 4. The black square in the middle is the aperture of the spectrometer and is 10 x10 μm in size. The corresponding spectra are in Fig. A10.
Figure A10: Spectra taken on the MSP of four grains of diopside with the same composition (images in Fig A9)

Figure A11: Spectrum from diopside 2, and the average of spectra from all four diopside grains (shades of red) shown on the same scale as five clinopyroxene grains analysed by UNIWIN (shades of blue). The UNIWIN grain compositions are: Px018 – En47.4Fs3.4Wo48.4; Px009 – En38.3Fs9.7Wo50.5; Px039 – En29.6Fs2.8Wo50.3; Px038 – En11.5Fs35.3Wo53.8; Px021 – En4.5Fs43.7Wo52.1
Figure A12: Data for a suite of high-calcium pyroxenes analysed by both RELAB (lines in green) and UNIWIN (lines in blue). The composition of the grains is given in the legend to Fig A11. Data from (UNIWIN public database; http://psf.uwinipeg.ca/Sample_Database.html) and RELAB database
Figure A13: Spectrum of wollastonite taken on the MSP (in black) compared with that acquired by the RELAB system (RELAB sample-id PP-EAC-006-PYX011; in blue). The steep drop-off below 350 nm in the RELAB data is possibly a system artefact.

Table A4: Composition of the feldspars analysed in this chapter. The composition analysis was performed using a Scanning Electron Microscope.

<table>
<thead>
<tr>
<th>Composition</th>
<th>Na (wt. %)</th>
<th>Al (wt. %)</th>
<th>Si (wt. %)</th>
<th>Ca (wt. %)</th>
<th>O (wt. %)</th>
<th>K (wt. %)</th>
<th>Fe (wt. %)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Albite</td>
<td>An&lt;sub&gt;1.9&lt;/sub&gt;Ab&lt;sub&gt;38.1&lt;/sub&gt;</td>
<td>8.8</td>
<td>10.4</td>
<td>31.8</td>
<td>0.3</td>
<td>48.7</td>
<td>-</td>
</tr>
<tr>
<td>Anorthite (bytownite)</td>
<td>An&lt;sub&gt;7.3&lt;/sub&gt;Ab&lt;sub&gt;21.3&lt;/sub&gt;</td>
<td>1.8</td>
<td>17.4</td>
<td>22.4</td>
<td>12</td>
<td>46.4</td>
<td>-</td>
</tr>
<tr>
<td>Orthoclase</td>
<td>Ab&lt;sub&gt;18.5&lt;/sub&gt;Or&lt;sub&gt;91.1&lt;/sub&gt;</td>
<td>1.7</td>
<td>9.8</td>
<td>30.6</td>
<td>-</td>
<td>46.5</td>
<td>11.1</td>
</tr>
<tr>
<td>Nepheline</td>
<td>12.4</td>
<td>16.9</td>
<td>21.3</td>
<td>-</td>
<td>44.6</td>
<td>4.2</td>
<td>Trace</td>
</tr>
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</table>
Figure A14: Images taken by the MSP of grains of feldspars. a) albite, b) anorthite, c) nepheline. The black square in the middle is the aperture of the spectrometer and is 10 x 10 μm in size.
Figure A15: Visible spectra of feldspars a) from 250 to 800 nm; b) from 250 to 500 nm; c) from 500 to 800 nm. Spectra taken on the MSP were averaged from spectra taken in different locations on the sample.
Figure A16: Spectra of the feldspars (a) anorthite (bytownite An78.7) and (b) albite (An1.9) taken by the MSP compared with data from the RELAB and UNIWIN facilities. The same plagioclase samples were measured by both RELAB and UNIWIN: Albite – PLG103 (An0); Anorthite (bytownite) – PLG111 (An78). Spectra taken on the MSP were an average of spectra taken in 5 separate locations on a single grain.
## Oxides and Sulphides

Table A5: Composition of the oxides and sulphide analysed in this chapter. The composition analysis was performed using a Scanning Electron Microscope.

<table>
<thead>
<tr>
<th></th>
<th>Mg  (wt%)</th>
<th>Al  (wt%)</th>
<th>Si  (wt%)</th>
<th>Ca  (wt%)</th>
<th>Ti  (wt%)</th>
<th>Fe  (wt%)</th>
<th>Ce  (wt%)</th>
<th>O   (wt%)</th>
<th>Cr  (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spinel</td>
<td>17.3</td>
<td>37.1</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>44.8</td>
<td>0.7</td>
</tr>
<tr>
<td>Magnetite</td>
<td>0.4</td>
<td>-</td>
<td>0.8</td>
<td>0.4</td>
<td>-</td>
<td>75.5</td>
<td>-</td>
<td>22.9</td>
<td>-</td>
</tr>
<tr>
<td>Hibonite</td>
<td>1.2</td>
<td>42.8</td>
<td>0.2</td>
<td>4.6</td>
<td>1.5</td>
<td>3.4</td>
<td>2</td>
<td>43.3</td>
<td>-</td>
</tr>
</tbody>
</table>

Figure A17: Images taken with the MSP of oxides and sulphides (a) spinel, (b) magnetite, (c) hibonite, (d) pyrrhotite. The black square in the middle is the aperture of the spectrometer and is 10 x10 μm in size.
Figure A18: Visible spectra of oxides a) from 250 to 800 nm; b) from 250 to 500 nm; c) from 500 to 800 nm. Spectra taken on the MSP were averaged from spectra taken in different locations on the sample.
Figures A12. Reflectance of various soil samples across different wavelengths:

(a) Sp Av and Sp122r. Spectra were developed from spectra taken in 1997 and 1998 on a slope.

(b) MT Av and MAG103r. Spectra were developed from spectra taken in 1987 on a slope.
Figure A19: Spectra of oxides and a sulphide taken by the MSP and in the RELAB facilities: (a) spinel, (b) magnetite, (c) hibonite and (d) pyrrhotite. Spectra taken on the MSP were averaged from spectra taken in at least 5 different locations on a single mineral grain.
Figure A20: Images taken by the MSP of grains of hibonite; the associated spectra are shown in Fig. A21: a) hibonite 1, b) hibonite 2, c) hibonite 3, d) hibonite 4, e) hibonite 5. The black square in the middle is the aperture of the spectrometer and is 10 x 10 μm in size.

Figure A21: Spectra taken on the MSP of different grains (images in Figure A20) of hibonite compared with results of the same mineral taken by RELAB. Sample hibonite RELAB 1 is 0-45 μm particulate.
Carbonates

Table A6: Composition of the carbonates analysed in this chapter. The composition analysis was performed using a Scanning Electron Microscope

<table>
<thead>
<tr>
<th></th>
<th>Cr (wt%)</th>
<th>C (wt%)</th>
<th>Mg (wt%)</th>
<th>Ti (wt%)</th>
<th>Ca (wt%)</th>
<th>Fe (wt%)</th>
<th>O (wt%)</th>
<th>Sr (wt%)</th>
<th>Mn (wt%)</th>
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<tbody>
<tr>
<td>Siderite</td>
<td></td>
<td>13.4</td>
<td>0.9</td>
<td></td>
<td>10.3</td>
<td>25.3</td>
<td>48.3</td>
<td></td>
<td>1.7</td>
</tr>
<tr>
<td>Magnesite</td>
<td></td>
<td>12.9</td>
<td>2.6</td>
<td></td>
<td>33.8</td>
<td>0.5</td>
<td>49.8</td>
<td>trace</td>
<td>0.5</td>
</tr>
<tr>
<td>Dolomite</td>
<td>0.3</td>
<td>11.3</td>
<td>8.8</td>
<td>8.5</td>
<td>12.4</td>
<td>9.6</td>
<td>49.3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Aragonite</td>
<td></td>
<td>13.8</td>
<td></td>
<td></td>
<td>34.8</td>
<td></td>
<td>50.8</td>
<td>0.6</td>
<td></td>
</tr>
</tbody>
</table>

Figure A22: Images taken with the MSP of grains of carbonates. a) siderite, b) magnesite, c) dolomite, d) aragonite. The black square in the middle is the aperture of the spectrometer and is 10 x 10 μm in size.
Figure A23: Visible spectra of calcite (a) from 300 to 800 nm, b) from 300 to 500 nm at four wavelengths (300-500 nm). Spectra taken on the NPP were averages of spectra taken from at least five different coatings on single mineral grains.
Figure A23: Visible spectra of carbonates a) from 250 to 800 nm; b) from 250 to 500 nm; c) from 500 to 800 nm. Spectra taken on the MSP were averages of spectra taken from at least five different locations on single mineral grains.
Figure A24: Spectra of the carbonates aragonite, dolomite, magnesite and siderite taken by the MSP and in the RELAB facilities. Exact composition of the minerals used by RELAB is unknown. Spectra taken on the different sample by the MSP were averages of spectra taken from at least 5 different locations on a single mineral grain.

Phyllosilicates

Table A7: Composition of the clays analysed in this chapter. The composition analysis was performed using a Scanning Electron Microscope.

<table>
<thead>
<tr>
<th></th>
<th>Na (wt%)</th>
<th>Mg (wt%)</th>
<th>Al (wt%)</th>
<th>Si (wt%)</th>
<th>K (wt%)</th>
<th>Ca (wt%)</th>
<th>Ti (wt%)</th>
<th>Fe (wt%)</th>
<th>O (wt%)</th>
<th>Cl (wt%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lizardite</td>
<td>-</td>
<td>25.6</td>
<td>0.3</td>
<td>21.8</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>41.9</td>
<td>0.2</td>
</tr>
<tr>
<td>Montmorillonite</td>
<td>1.1</td>
<td>1.9</td>
<td>9.3</td>
<td>24.9</td>
<td>0.3</td>
<td>1</td>
<td>0.3</td>
<td>4.7</td>
<td>40.3</td>
<td>-</td>
</tr>
<tr>
<td>Saponite</td>
<td>0.6</td>
<td>11.5</td>
<td>3.2</td>
<td>19.9</td>
<td>-</td>
<td>1.3</td>
<td>-</td>
<td>8.4</td>
<td>36.2</td>
<td>-</td>
</tr>
</tbody>
</table>
Figure A25: Images taken with the MSP of grains of clays. (a) lizardite, (b) montmorillonite, (c) saponite. The black square in the middle is the aperture of the spectrometer and is 10 x 10 μm in size.

Reflectance plot showing the wavelength (nm) versus reflectance values for montmorillonite, lizardite, and saponite.

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Figure A26: Visible spectra of clays a) from 250 to 800 nm; b) from 250 to 500 nm; c) from 500 to 800 nm. Spectra taken on the MSP were averages of spectra taken in at least 5 different locations on the sample.
Figure A27: Spectra of montmorillonite and saponite taken by the MSP and in the RELAB facilities. Exact composition of the minerals used by RELAB is unknown. Spectra taken on the MSP were averages of spectra taken in at least 5 separate locations on the sample.