VUV Emission and Absorption Spectroscopy with a Multichannel Near Normal Incidence Spectrometer

A thesis for the degree of Master of Science

submitted to

Centre for Laser Plasma Research
School of Physical Sciences

by

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I hereby certify that the material, which I now submit for assessment on the programme of study leading to the award of Mater of Science is entirely my own work and has not been taken from the work of others save and to the extent that such work has been cited and acknowledged within the text of my own work

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Abstract

An experiment for vacuum UV emission and photoabsorption spectroscopy using a normal incidence multi-channel spectrometer is described. Two detection configurations were used in the experiments, one a Channel Electron Multiplier Array with Photo Diode Array readout (CEMA/PDA), the other a directly illuminated back thinned Charge Coupled Device (CCD).

Emission spectra of plasmas of a number of low and high Z target materials were recorded covering the majority of the vacuum UV spectral region (35 to 200 nm). These experiments were designed mainly to study the nature of the emission (line versus continuum) and its dependence on atomic number. The use of two detection configurations allowed for inter comparisons to be made on resolution, efficiency and dynamic range of the two multi-channel detector and spectrometer systems.

Experiments were also performed on photoabsorption by helium gas. The spectra thus obtained were used to help characterise the system’s application to absorption studies around the He principal series limit at ~ 50 43 nm.

In addition, for the first time the instrument was used to study photoabsorption using the dual laser plasma technique (DLP). Results on the relative photoabsorption cross sections of barium and barium ions using a samarium continuum back-light are presented which indicate that the instrument is well suited to absorption experiments in the range of 35 to 80 nm where second order of the continuum is mainly absent and the grating is being used close to its blaze wavelength (80 nm).

Finally, the work concludes with details on the design, construction and preliminary testing of a vacuum-optical beamline which should both improve the aperture matching ability of the instrument and allow spatially resolved studies of the laser plasma light source to be made. The spectrometer with the additional spherical mirror was modelled by means of simple geometric optic calculations as well as with ray tracing software in order to estimate improvements in throughput.
Chapter 1. Overview of laser produced plasmas for spectroscopy.

The following chapter outlines the general properties of plasmas. The focus of this chapter is on laser-produced plasmas, their evolution and a number of the main atomic processes dictating their behaviour. The laser plasma as a light source for spectroscopy and most especially as a continuum light source for photoabsorption studies is discussed. Other applications to date of laser produced continua in e.g., photoelectron spectroscopy are listed.

1.1 Introduction to laser produced plasmas

A plasma is a quasi-neutral gas of charged and neutral particles. Plasmas are dynamic systems for which restoration of neutrality is the dominant physical process. A plasma is said to exhibit collective behaviour, i.e., particles in a plasma which are charged will interact by the Coulomb force [Carroll et al., 1981]. Plasmas can be formed by any method that can couple enough energy into a material to permit ionization of the atoms and continue this input of energy to overcome the natural recombination process that occurs thereafter. One method of plasma creation involves the use of intense laser light interacting with a material, a so-called laser produced plasma.

When a high power pulsed laser is focused to an irradiance of greater than $10^8$ W cm$^{-2}$ onto a solid material, a short duration, high density and high temperature plasma is formed. The ability to provide a controlled environment for the study of ionized species is principally the reason why the laser produced plasma has been extensively used in atomic spectroscopy. The laser plasma allows for highly ionized atomic species and refractory elements to be studied, something which is not readily possible with other methods [Costello et al., 1991]. Ion densities of $10^{15}$ – $10^{18}$ cm$^{-3}$ and effective temperatures in the range of 10 eV – 1 keV (1 eV = 11605 K) are readily achieved with tabletop lasers. At such temperatures, laser plasmas emit copious amounts of vacuum UV (VUV) and extreme UV (XUV) radiation with a duration comparable to that of the exciting laser pulse. The following sections outline the initiation and evolution of laser produced plasmas and discusses some of the physical processes occurring within the plasma plume.
For collective behaviour to dominate in a plasma, the extent of the Coulomb force on an ionized species must be comparable to the extent of the plasma. A parameter that quantifies the effective extent of the Coulomb force around a charged species is the Debye length \( \lambda_{\text{Debye}} \) (m). It can also be viewed as a measure of the shielding effect of the species surrounding a charged particle. Using Poisson’s equation and the fact that the distribution of energies of the plasma species can be modelled by a Boltzmann distribution, it can be shown that,

\[
\lambda_{\text{Debye}} = \left( \frac{e_0 k_B T}{n_e e^2} \right)^{1/2} \quad (\text{SI units}) \quad [\text{E 1-1}]
\]

where \( T \) (K) is the electron temperature, \( n_e \) (m\(^3\)) is the electron number density, \( e \) (C) is the charge on the electron, \( k_B \) (J K\(^{-1}\)) is Boltzmann’s constant and \( e_0 \) (8.8542 * 10\(^{-12}\) F m\(^{-1}\)) is the electric permittivity of free space [Hughes, 1975].

When the Debye length is comparable to the size of the plasma, collective behaviour results in the plasma exhibiting oscillations at some frequency \( \omega_{\text{plasma}} \) (radians s\(^{-1}\)) such that,

\[
\omega_{\text{plasma}} = \left( \frac{n_e e^2}{m_e e_0} \right)^{1/2} \quad (\text{SI units}) \quad [\text{E 1-2}]
\]

where \( m_e \) (kg) is the mass of an electron [Hughes, 1975]. Similarly, the various ion stages have their own associated frequencies. These natural frequencies determine the response of the plasma to radiation incident on it.

The dispersion relation for electromagnetic waves travelling through a plasma is given by,

\[
\omega^2 = \omega_{\text{plasma}}^2 + c^2 k^2 \quad [\text{E 1-3}]
\]

where \( c \) (m s\(^{-1}\)) is the speed of light in vacuum, and \( k \) (\( k = n - ik \), where \( n \) and \( k \) being the optical constants of the material in question) is the propagation constant describing the material's effect on electromagnetic waves and is in general an
imaginary number. Considering the case were \( \alpha_{\text{plasma}} < \omega, k \) has a zero imaginary component which results in the electromagnetic wave propagating without losses (ignoring other absorption effects). When \( \alpha_{\text{plasma}} > \omega, k \) has a non-zero imaginary component and thus the electromagnetic wave does not propagate and is reflected by the medium. The electron density when \( \alpha_{\text{plasma}} = \omega \) has reached the critical density at which the electromagnetic wave is reflected back from the plasma.

\[
 n_{\text{critical}} = \left( \frac{\varepsilon_0 m \omega^2}{e^2} \right) \quad \text{(SI units)} \quad [\text{E 1-4}]
\]

### 1.2 Evolution of laser produced plasmas.

The initial stages of plasma formation differ for the case of conductors and insulators. When the electromagnetic field of laser radiation impinges on a metallic target surface, it penetrates only to a very small depth. This is commonly referred to as the skin depth \( \delta \) (m) and has a magnitude, which is a fraction of the wavelength \( \lambda \) (m) being used.

\[
\delta = \sqrt{\frac{\pi \varepsilon_0 \mu_0 \sigma}{\lambda}} \quad \text{(SI units)} \quad [\text{E 1-5}]
\]

Where \( \mu_0 (4\pi \times 10^{-7} \text{ H m}^{-1}) \) is the permeability of free space and \( \sigma (\Omega^{-1} \text{ m}^{-1}) \) is the material's electrical conductivity. The strength of the electric field is such that it interacts with the target material’s free electrons and in turn heating, evaporation and vaporisation occur rapidly. The plasma thus formed is relatively cold at this stage and in the shape of a thin sheet close to the target’s surface. The root mean square (r.m.s.) electric field \( E \) (V m\(^{-1}\)) is related to the irradiance \( \phi \) (W m\(^{-2}\)) by the following relationship.

\[
E_{\text{r.m.s.}} = 19.4\sqrt{\phi} \quad \text{(SI units)} \quad [\text{E 1-6}]
\]

In the case of insulators, the laser photon energy is lower than the ionization potential of the material. Despite this, free electrons can be produced by e.g., multiphoton ionization. Furthermore, easily ionized and free electrons exist at defects and impurities and ionization may proceed via other non-linear effects e.g. Stimulated
Brillouin Scattering (SBS) Once free electrons are produced by whatever generation process, laser radiation is efficiently absorbed and a thin plasma sheet forms.

Following this pruning stage, absorption of the laser light by inverse bremsstrahlung occurs. This is a process whereby a photon is absorbed by an electron in the presence of a neutral atom/ion resulting in the electron being raised from a lower continuum level to a higher one, thereby increasing its kinetic energy. Inverse bremsstrahlung provides the dominant process for laser light absorption.

In a laser produced plasma an electron density gradient exists and therefore the plasma frequency as given by $\omega = \sqrt{\frac{e^2 n_e}{\mu m}}$ is also a function of position within the plasma. When the incident laser radiation reaches the region in the plasma plume where $n_e = n_{\text{critical}}$, it is reflected back towards the laser. At this point, the radiation is no longer reaching the target surface and therefore plasma production stops. Because the existing plasma material is been continuously driven away from the target surface, the density gradient will reduce to a level were the laser radiation can again reach the target's surface. This process repeats itself for the duration of the laser pulse. After the termination of the excitation laser pulse the plasma continues to expand outwards away from the target surface adiabatically and cools rapidly.

1.3 Atomic processes in plasmas

In a typical plasma, the range of neutral and ionized species present results in a large number of possible interactions. Quite often, when considering specific plasmas, a small number of these interactions provide the dominant mechanisms for energy losses and gains. One property, which makes the modelling of such a system easier, is the fact that in many laboratory plasmas, local thermodynamic equilibrium (LTE) prevails where collisional excitation and ionization are balanced by their inverse processes – namely, collisional decay and recombination and are the dominant processes within the plasma. In laser plasmas with intermediate electron densities (typically $n_e < 10^{20}$ cm$^{-3}$), in general the system is better represented by the

---

1 Bremsstrahlung German word for 'breaking radiation'
collisional-radiative (CR) equilibrium model [Colombat and Tonon, 1973] The relative significance of particular processes resulting in radiative cooling of a plasma plume is strongly dependent on the atomic number of the target material as well as degree of ionization of the plasma. These processes can be grouped together into three main groups, bound-bound transitions, bound-free transitions and free-free transitions. Each of these occurs with the difference in energy being compensated by a collision or a photon event. Only photon generating and loss events are considered in the following discussion.

As already stated inverse-bremsstrahlung provides the dominant mechanism in the evolution and growth of a plasma. Its inverse process, i.e., bremsstrahlung is important because it is one important mechanism by which continuum emission results. Bremsstrahlung refers to the process whereby an electron makes a transition from a higher free electron energy state to a lower one, with the emission of a photon of energy equal to the difference between the two energy levels \([E \ 1-7]\). The important interactions are those, which occur between electrons and ions, since those between electron-electron and electron-neutral species, do not provide an effective conversion of electron kinetic energy to continuum emission.

\[
X^\text{**} + e^-(T_1) \leftrightarrow X^\text{**} + e^-(T_2) + h\nu_{\text{continum}} \quad T_1 > T_2 \quad [E \ 1-7]
\]

One of the most important processes, which takes place in plasmas – at least in spectroscopic studies – is radiative recombination and its inverse process of photo-ionization. In radiative recombination, the electron combines with an ion with the emission of a photon possessing energy equal to the energy difference between the initial and final states. These processes can occur in single as well as in multiple steps. The resulting emission or absorption spectrum has a continuum appearance with a fairly sharp low-frequency cut-off. This cut-off is known as the recombination limit, and is the minimum energy required to ionize the atom from the bound state. Recombination radiation is generally the dominant mechanism for continuum emission. A single step photoionization process (proceeding from left to right) / radiative recombination process (proceeding from right to left) is represented by
Another process that is of interest in the area of spectroscopy, is photo-excitation and its inverse process of photo-de-excitation. In this process, an electron makes a transition from a lower bound state to an upper bound state with the absorption of a photon. The resulting spectrum exhibits line structure. In the absence of perturbations the transitions gives rise to a spectral line whose profile depends upon the spontaneous lifetime of the upper state, and on the distribution of velocities of the emitting atoms, which causes a distribution of Doppler shifts. Collisions, electric and magnetic fields all perturb the initial and final states of the emitting atom and the resulting spectral line profile therefore contains valuable information on the conditions within the plasma [Hughes, 1975].

\[ h\nu_{\text{line}} + X \leftrightarrow X^+ \]  

1.4 Laser produced plasma light sources.

Shortly after the development of the Ruby laser by Maiman [1960] the study of intense laser–matter interaction was born. Early measurements of the spectra emitted by laser produced plasmas showed them to be intense sources of both line [Fawcett et al, 1966] and continuum emission [Ehler and Weissler, 1966]. Since then, laser produced plasmas have proven themselves to be both efficient and versatile sources for the study of emission and absorption spectroscopy of highly ionized atoms.

1.4.1 Line emission dominated plasmas.

Spectra of elements in the periodic table lighter than iron and nickel were the focus of special attention during the 1960’s and 1970’s, the two decades when most of the analysis of the solar spectrum retrieved by space based instruments took place. This was due to the fact that these studies displayed previously unidentified lines which could only be explained by higher ion stages being present in the sun’s corona. These new observations motivated studies of higher ion stages along iso-electronic sequences in these lighter elements. Experiments on fusion necessitated studies being
formed on the heavier elements in the periodic table. Impurity elements in tokamak* plasmas resulted in characteristic spectra, which proved to be a useful diagnostic for the estimation of electron densities and temperatures in these plasmas.

Line dominated emission from plasmas are important for the following reasons:

- They provide a testing ground for theoretical calculations.
- They increase our understanding of atomic structure.
- In addition, they provide a diagnostic tool for the measurement of plasma parameters, i.e., temperatures and densities etc.

Laser produced plasmas are thus a useful laboratory light source in which the ionization distribution may be somewhat tuned by adjusting the on target irradiance. It is well known that both ion stage distribution and the average electron temperature $T_e$ (eV) are dependant on the target irradiance $\phi$ (W cm$^{-2}$).

The average electron temperature for incompletely ionized targets of medium to high atomic number elements can be approximated by,

$$T_e = 5.2 \times 10^{-6} Z^{1/5} (\lambda^2 \phi)^{3/5} \quad 10^{11} \geq \phi \geq 10^{13} \text{ W cm}^2 \quad [ \text{E 1-10} ]$$

where $\lambda$ (µm) is the wavelength of the excitation laser and $Z$ is the atomic number of the parent material [Colombant and Tonon, 1973]. This approximation follows from the application of the collisional-radiative (CR) model to laser produced plasmas.

Using [E 1-8] the average ion stage can be approximated to,

$$Z \equiv \frac{2}{3} (AT_e)^{1/3} \quad T_e \geq 30 \text{ eV} \quad [ \text{E 1-11} ]$$

---

*Tokamak fusion devices rely on the magnetic confinement of a high temperature plasma in a "doughnut" shaped vessel. They represent one potential method for the implementation of a fusion energy source.
where $A$ (a.m.u) is the atomic mass number of the parent material and $T_e$ (eV) is the average electron temperature [Colombant and Tonon, 1973]. If a Nd YAG laser ($\lambda = 1064$ nm) is focused onto aluminium to an irradiance of $\sim 10^{12}$ W cm$^{-2}$, the average electron temperature ($T_e$) would be $\sim 142$ eV. The plasma thus formed would contain aluminium ions with an average ion stage of $\sim 10$ times ionized.

It is also well known that the ionization distribution varies with time (i.e., plasma cools and recombination takes place). Hence, by incorporating both time and space resolution in a spectrometer system, it is possible to choose, to a reasonably high degree of selectivity, a single ion stage [Costello et al., 1991].

### 14.2 Continuum emission dominated plasmas

The line-free continuum emission from a laser produced plasma was first observed and noted by Ehler and Weissler [1966]. In these studies, a 0.4 J Q-switched Ruby laser was focused onto a tantalum ($Z = 73$) target with a pulse duration of $\sim 40$ ns. Both a 0.5 metre normal incidence and a 2.0 metre grazing incidence spectrometer were used to record the resulting emission spectra, therefore giving a combined wavelength range between 5 and 400 nm with some overlap between the two instruments. A broad band continuum was observed with two principal intensity maxima (at $\sim 170$ nm and $\sim 50$ nm). Other high atomic number targets, tungsten ($Z = 74$) and platinum ($Z = 78$), were also studied in terms of their emission spectra, and these showed similar line-free continua with some small line structure being present on the platinum spectra at $\sim 25$ nm.

The use of rare-earth elements for generation of continua was first reported by Carroll et al. [1978, 1980]. In these investigations, the authors noted that with appropriate targets, laser-produced plasmas can provide clean essentially 'line-free' continua suitable for photoabsorption experiments in the range 4 – 200 nm. The rare-earth elements investigated were the group from samarium to ytterbium. It was observed that continua from these elements had time duration's which were essentially the same as the excitation laser source (in this case a Q-switched Ruby laser with pulse duration $\sim 25$ ns) in the XUV region and rising to in excess of 100 ns in the VUV normal incidence region.
O'Sullivan et al [1982] did further work on the absolute spectral irradiances of laser-produced continua in the range 115 to 200 nm. This work showed that the reproducibility of the continuum was as good as or better than the reproducibility of the excitation laser (typically better than 5%).

In recent years, a lot of interest in laser produced plasmas has been focused on the generation of ultra short pulses of continuum light. These studies have been motivated by the possibility of generating short duration x-ray pulses for pumping short wavelength lasers and for time-resolved studies of materials. In work performed by Meighan et al [1997] using ‘tabletop’ lasers, ‘line-free’ continuum emission in the XUV was observed from laser produced plasmas of high atomic number elements. The targets were irradiated by 248 nm laser pulses of 7 ps (FWHM) duration at a power density of ~ $10^{13}$ W cm$^{-2}$. A streak camera with a time resolution of ~ 100 ps recorded the resulting XUV emission. Both the coupling optics and the Mylar® thin film filter used yielded an instrument bandpass of 85 eV centred at 145 eV. Results on the broadband XUV emission from a lead plasma yielded an upper limit of FWHM of ~ 750 ps. More recent experiments [Meighan et al, 1998] have shown that XUV pulse durations of 100 to 200 ps can be obtained.

It is well established that the presence of a weak pre-plasma either formed by an actual pre-pulse or by ASE can dramatically increase the VUV and x-ray emission from the main plasma [Tom and Wood, 1988, Cobble et al, 1991 and Theobald et al, 1994]. The main laser pulse interacts with a gaseous plasma instead of the solid target material, and so the interaction volume within the short duration laser main pulse is larger in the former case. The interaction with the plasma, i.e., the absorption of laser radiation, the rate of the various atomic relaxation processes, and therefore the plasma emission will change with the changing interaction volume [Steingruber et al, 1996, Teubner et al, 1992, and Kieffer and Chaker, 1994].
1.4.3 The origin of rare-earth continua.

The extent and ‘line-free’ content of the continuum emission from rare-earth plasmas stems from three processes, recombination radiation, bremsstrahlung radiation and unresolved line radiation. The relative contribution to the continua from each of these three principal sources depends on the degree of ionization of the plasma as well as the atomic number of the parent material. The recombination component of the continuum scales as $\xi_{\text{average}}^4$, while the bremsstrahlung component scales with $\xi_{\text{average}}^2$, where $\xi_{\text{average}}$ is the average ion stage which is present in the plasma [O'Sullivan, 1983]. When the plasma is sufficiently hot (electron temperatures greater than ~ 50 eV) most of the outer electrons have been stripped away from the atoms and therefore recombination radiation will provide the dominant continuum emission process in the XUV. In contrast, bremsstrahlung contributes a relatively small amount (~ 20\%) to the overall continuum intensity making it’s greatest contribution at longer wavelengths (i.e., VUV) and generally lower ion stages i.e., cooler plasmas.

However, the latter discussion does not explain the key aspect of the continuum i.e., the lack of any significant structure due to discrete transitions within the distribution of ions in the plasma even at high resolution studies [Gohill et al., 1986], [Orth et al., 1986] ($\Delta\lambda = \sim 0.05$ nm in both cases). The elements which proceed the rare-earth’s and those immediately after show strong line emission and therefore factors must exist other than the large atomic number and highly charged species present which contributes to pure continuum emission from these rare-earth plasmas. In order to understand this observation one must consider the following argument.

In low ion stages of the elements $62 \leq Z \leq 74$ (samarium to tungsten) the 5s and 5p sub-shells are filled while the occupancy of the 4f sub-shell varies with atomic number and ion stage. With increasing ionization the binding energy of 4f electrons increases more rapidly than that of 5s or 5p electrons so that 4f / 5s or 5p level crossing occurs at about six times ionized species and 4f / 5s at about twelve times ionized species. Consequently all the elements of interest contain 4f electrons in the ground state for each ion stage generated in the laser plasmas under discussion here.
It is the presence of f electrons in both the upper and lower states involved which give rise to very large number of lines which are generally not possible to resolve (hence the term 'Un-resolved Transition Array' or UTA).

Figure 7.7  Total number of lines in LS coupling predicted for 4d-4f transitions in the rare-earth and immediately proceeding elements. Broken curve, number of predicted lines when (4f, 5p) and (4f, 5s) degeneracy is included [Carroll and O'Sullivan, 1982]

The proximity of 4f, 5s and 5p binding energies gives rise to a band of overlapping complex configurations with variable numbers of 4f, 5s and 5p electrons. Furthermore, if the plasma is optically thick (e.g., pure metal plasma) then the photoabsorption cross sections for these more intense lines will be greater and so the lines will be attenuated and merge into the underlying continuum [Carroll and O'Sullivan, 1981].

From this brief qualitative analysis, it can be seen that the continuum from rare-earth elements under the conditions normally present in a laser produced plasma is not the result of pure bremsstrahlung and recombination processes but that it
contains a very large number of possible transitions which give the appearance of a pure continuum band even at high spectral resolution. The presence of a few lines on the spectra (mostly at wavelengths greater than ~100 nm) of the rare-earth's is due to simpler electronic configurations in low ion stages and/or a configuration from which only a very small number of permitted transitions are allowed (e.g. $4d^{10}(^{1}S) \rightarrow 4d^{9}nl^{(1}P)$ arrays, see figure 1-2 and 1-3).

The extent of the continuum and position of the characteristic lines have been illustrated in figure 1-2 by O'Sullivan [1983] and summarizes work done by Carroll et al. [1980] on continuum emission of elements $62 \leq Z \leq 74$ in the wavelength range of 200 to 4 nm.

It can be seen that a number of trends are evident. Considering the long wavelength portion of the figure 1-2, two principal sets of characteristic lines appear superimposed on the continuum. The first of these sets located at ~100 nm gradually increases in both intensity and extent as the atomic number of the target material increases. The other set of lines located at wavelengths greater than ~180 nm for samarium rapidly moves to shorter wavelengths as the atomic number of the target material increases and overtakes the 100 nm group at the element ytterbium. In tungsten this latter group is evident at wavelengths greater than ~125 nm.
Figure 1-2
Extent of the continuum emission as a function of Z, the atomic number. It should be noted that most of the line emission from ion stages with $\xi \leq 6$ and can be eliminated by increasing the plasma temperature. The shaded region denotes line emission present. The resulting power density at focus was estimated by consideration of the abundance of ion stages to be $\sim 2 \times 10^{11} \text{ W cm}^{-2}$ [O'Sullivan, 1983].

Figure 1-3
(a) Approximate intensity distribution in rare earth continuum (samarium $Z = 62$ to ytterbium $Z = 70$) in the 20 to 200 nm region. (b) Region where line emission is superimposed on rare earth continuum. Lines in central region are relatively weak and scarcely effect the continuum character of the emission even at moderate dispersion $\sim 0.12 \text{ nm mm}^{-1}$ [Carroll et al, 1980].
1.5 Other applications of laser-produced continua.

The laser plasma light source has proven itself efficient in the production of continuum radiation. A distinction between applications requiring broadband continua and those requiring narrow band XUV or VUV continua can be made. Laser-produced plasma light sources for proximity lithography require narrow band continua above the silicon K-edge (~ 0.674 nm) but of short enough wavelength for efficient absorption by the photo-resist (i.e., generally between ~ 1.0 and ~ 1.4 nm) [Celliers et al., 1995]. Therefore, choice of targets here generally include L-shell emitters such as Ne-like ions (Fe, Z = 26, Co, Z = 27, Cu, Z = 28, Zn, Z = 29) or M-shell emitters such as Ni-like ions (Xe, Z = 54). On the other hand, XUV sources needed for microscopy require narrow band emission confined to the water window (4.376 nm to 2.332 nm)†. In this case, sources using target materials such as silicon (Z = 14) or carbon (Z = 6) are generally used. Some applications requiring broadband continua and using rare-earth elements are briefly discussed below.

1.5.1 VUV reflectivity measurements

The study of the interaction of light with a material has long been recognized as a fundamental method of determining the electronic structure of matter. The number of applications requiring efficient optics for the low wavelength regime is increasing and therefore accurate determinations of the optical constants across the entire range from x-rays to VUV is important. The optical constants are macroscopic quantities, which describe the electrical permittivity of the medium. One of the techniques of determining the optical constants from raw reflection data is by use of the Kramers Kronig integrals. A prerequisite of an accurate Kramers Kronig analysis is the possession of reflectivity data over a wide spectral range.

In work performed by French et al. [1989], a cylindrical samarium target.

† The water window: Wavelength region between the K absorption edges of carbon (4.376 nm) and oxygen (2.332 nm). In this range, the contrast between a biological sample and the surrounding water is high.
(Z = 62) was illuminated by a focussed 500 mJ, 8 ns (FWHM) Nd YAG laser operating at a repetition frequency of a 20 Hz. The focused laser had an estimated spot size of ~ 100 μm in diameter corresponding to an irradiance of approximately 10^{12} \text{W cm}^{-2}. The VUV light from the laser-produced plasma was focused by a spherical mirror onto either a sample surface or an iridium reference mirror. The spot size of this focused broad band VUV light was estimated to be 3 mm in diameter. A normal incidence spectrometer utilising an 1800 lines mm\(^{-1}\) grating was used to observe the reflected light in the region of 5 to 40 eV. The use of the iridium reference mirror allowed for the absolute reflectivity of the samples to be measured with a quoted accuracy of ~ 5 %. A Kramers Kronig analysis of this reflection data yielded values for the optical constants for the samples under consideration.

15.2 VUV radiative transfer standards.

Radiometric standards can be classified as those sources of calculable spectral radiant power – primary standards, and those sources with stable spectral radiant power, which can be controlled by a few source parameters. These latter sources are calibrated against primary standards and serve as transfer source standards – secondary source standards [Kuhne and Wende, 1985].

Radiometric techniques for wavelengths are well established with calibrated transfer sources for λ ≥ 100 nm and transfer detectors for λ ≤ 5 nm. Below ~ 100 nm, however, in general the problem of accurately determining spectral radiant quantities of sources, e.g., radiance (W m\(^{-2}\) sr\(^{-1}\)) and spectral radiance (W m\(^{-2}\) sr\(^{-1}\) nm\(^{-1}\)) is not yet well solved. Detectors with known sensitivity can only be used for measuring the spectral radiance of sources if the spectral transmittance of the instrument has accurately been measured. Ideally, the instrument should be calibrated by use of a primary source standard such as light from a synchrotron insertion device, which has been passed through an accurately calibrated monochromator. Unfortunately, this is not always possible for such reasons such as cost and convenience. Also, optical instruments degrade with time due to contamination of reflecting surfaces and detectors and therefore the ability to calibrate the instrument in-situ would be a great advantage [Bibinov et al, 1997].
Laser produced plasmas serve as effective sources for XUV and VUV radiative transfer standards. The emission is both intense (typically $10^{23}$ photons cm$^{-2}$ sr$^{-1}$ s$^{-1}$ nm$^{-1}$ @ 120 nm) and reproducible (better than 5% typically) [Carroll et al, 1983]. The fact that this radiometric standard uses a source, which is both small and simple to implement means that it can offer a non-obtrusive in-situ means of calibration in various plasma experiments.

### 1.5.3 VUV photoelectron spectroscopy

Laser-produced plasmas have recently been used as an alternative to synchrotron based sources for X-ray photoelectron spectroscopy (XPS). Research in Japan by Ohchi et al [1996] has shown promising results using a grazing incidence based monochromator with toroidal optics combined with a Wolter mirror in the study of laser ablated silicon surface using photoelectron spectroscopy. To detect the energy of the photo-generated electrons, a hemispherical electron energy analyzer was used. A pinhole between the monochromator and the Wolter mirror was used in order to select the x-rays with wavelength of 5.24 nm. The Wolter mirror was designed and positioned to demagnify the monochromatized x-ray source to 1/32. With a pinhole diameter of 500 μm, it resulted in a spot size of approximately 20 μm in diameter. Photoelectrons from the sample were collected by a retarding lens, analyzed by a hemispherical electrostatic electron analyzer and detected by a position-sensitive MCP. The energy resolution of the analyzer was estimated to be ~2 eV.

### 1.6 Advantages of laser produced plasma continuum light sources

The inherent properties of rare-earth plasma continua are important considerations when choosing an ideal light source for an application. There are certainly disadvantages as with any source but by careful attention to system design and geometry some of these can be reduced to an acceptable level, i.e. uniformity of...
emission, shot to shot reproducibility etc. The basic property of being 'tabletop' means that some of these disadvantages are much reduced when considering cost, size and commercial availability of complete set-ups. In applications requiring an unpolarized source of XUV and VUV radiation with high single shot brightness, a laser plasma continuum source is ideal [Costello et al., 1991] Where polarization is critical, a high degree of polarized XUV can be obtained by using a 45° multilayer reflectors

1.6.1 High spectral irradiance

An important parameter of light sources of all types is the spectral radiance per pulse and can be expressed in units of photons cm$^2$ sr$^{-1}$ nm$^{-1}$ s$^{-1}$. This summarises a number of properties such as intensity, directionality and spectral content. Whereas the peak x-ray irradiance from laser-driven plasma source can be up to 4 orders of magnitude higher than that of synchrotron light sources and have a duration up to 200 times longer, the repetition frequency is much lower, typically 4 to 5 orders of magnitude [Costello et al., 1991]. This reduces the average power of the LPP source which (even at a repetition rate of 100 Hz) can be lower than bending magnet beam lines and orders of magnitude below insertion devices. Further, the LPP is a point source while light from say an insertion device is collimated. The laser produced plasma continuum has a peak spectral irradiance per pulse which is a factor of $10^3$ higher than that of the B-R-V (Ballofet-Romand-Vodar)\(^1\) source [Ballofet et al., 1961] which was one of the few laboratory continuum sources available before the advent of the laser produced plasma light source [Carroll et al., 1980].

1.6.2 Low line structure content and good reproducibility

The ideal continuum light source for spectroscopic studies would be a uniform point source and would have a broad band continuum spectral content and would be perfectly reproducible from shot to shot. The laser plasma continuum meets somewhat with this ideal if the target is chosen and prepared carefully. Work has

\(^1\) The original B-R-V source consists of three electrodes – one of which is of material with high atomic number - separated by a small distance in a vacuum. A high voltage stored in capacitor is discharged across the intervening gap between the electrodes thus producing a plasma which emits continuum radiation
been carried out in this respect by Kuhne [1982] and it has been shown that shot to shot reproducibility can be better than ~5% for part of the normal incidence spectral region (40 to 80 nm). This compares to a value of better than ~1% for synchrotron light sources and 25–40% for B-R-V based sources [Kuhne, 1982].

16.3 Temporal and spatial plasma properties.

The small size and potentially short duration of laser plasma continuum emission make it an exceptionally good back-lighter in experiments such as the dual laser plasma experiments (DLP). At DCU, a predominantly tungsten target is illuminated with 1064 nm radiation from a Nd:YAG laser with a duration of ~17 ns. The resulting plasma has a continuum emission duration in the XUV comparable to the laser pulse width [Collins et al., 1993] and thus allows time resolved experiments to be carried out without the need for detector gating. Because of the small size, the continuum source acts as a point source, which is generally easier to model in ray tracing software such as Shadow†. The LPP continuum source also has increased uniformity in the angular sense when compared to the synchrotron source [Carroll et al., 1980].

† Shadow Ray tracing software using Monte Carlo techniques for the modelling of VUV, XUV and soft x-ray optical systems. Available from www.xraylith.wisc.edu/shadow/
1.7 Dual laser plasma (DLP) technique.

The structure of the atom is such that inner shell electron binding energies generally result in energy transitions in the VUV normal incidence region and XUV part of the spectrum. Therefore, in order to investigate these transitions in detail, both a way of generating continuum at these wavelengths and a means of detecting the spectral signal are needed.

The laser produced plasma is ideal for the production of charged species as the relative abundance of certain ion species can be controlled to some degree by changes in the power density, i.e., by changing the on-target focusing conditions or altering the laser pulse energy. If the system set-up allows for temporal—by control of the inter laser delay—as well as spatial resolution—i.e., by the use of apertures, slits and/or toroidal optics—then a high degree of ion stage differentiation can be obtained.

![Diagram showing general arrangement for dual laser plasma (DLP) experiment](image)

*Figure 1-4 Diagram showing general arrangement for dual laser plasma (DLP) experiment*
1.7.1 History of the DLP technique.

The first attempt to develop a dual laser plasma technique for photoabsorption studies was made by Carillion et al. [1970] who used aluminium for both the continuum emitting plasma as well as for the sample plasma. Although the emission from aluminium is rich in line structure, the authors found that the underlying weak continuum emission present could be used for a continuum light source between the stronger line emissions. This restricted them to time and space studies of the laser generated plasma plume within these 'line free' windows (~ 98 nm). The results showed that inverse bremsstrahlung was the dominant mechanism in the hot dense region of the aluminium plasma plume (i.e., at short inter laser time delays) and also that photoionization was evident at moderate distances (~ 0.3 mm) where the plasma was cooler and thus recombining.

The application of rare-earth continua to photoabsorption studies was first performed by Carroll et al. [1977]. In these experiments, the authors studied the photoabsorption spectra of single ionized lithium Li⁺. Using a beam splitter, a Ruby laser was split into two beams, the first of which was focused on a cylindrical tungsten target and the other onto a planar lithium target. The experimental geometry was such that it resulted in spatially resolved spectra with a magnification of ~ 20. The 1s² ¹S₀ → 1snp ¹P₀ principal series of Li⁺ was recorded up to the n = 7 member. The singly charged ion Li⁺ is iso-electronic to helium and thus shows analogous doubly excited hollow atom states to helium but at a much higher energies due to the greater nuclear charge present. In the work by Carroll et al. [1977] the doubly excited 1s² ¹S₀ → 2snp ¹P₁ series – which was first observed 13 years previously in He by Madden and Codling [1965] with the use of a synchrotron radiation source – was observed for the first time in a He like ion.

One of the main advantages of using the DLP technique is that it allows for the study of photoabsorption in extended iso-nuclear and iso-electronic sequences.
1.7.2 Recent XUV DLP experiments on lithium

The application of the dual laser produced plasma technique to the study of ‘hollow lithium’† was recently performed by Kiernan et al. [1994]. On moving from helium to lithium the addition of a single electron makes possible the formation of triply excited states in which all three electrons reside in the L (or higher) shell leaving the K shell unoccupied. Electron-electron effects are dominant in this Coulombic four-body system, thereby offering the possibility of studying both direct and resonant double photoionization. The experimental arrangement consisted of a grazing incidence type spectrometer with a 1200 groove mm⁻¹ grating and a CEMA/PDA detector combination resulting in a resolution of ~1500 before de-convolution. The absorbing column was produced by focusing the output from a flashlamp pumped dye laser (2.5 J with pulse duration of 0.5 µs FWHM) onto the lithium target with a 22 mm line focus. The photoabsorption spectrum in the vicinity of the 1s²2s (²S) → 2s²2p (²P) triply excited transition was studied, and the position, width and profile index of the main resonance were measured.

† A hollow atom or ion is one in which at least one inner-shell is empty.
Figure 1-5 (a) K-shell absorption spectrum of the dye laser produced lithium plasma at a inter plasma delay of 750 ns corresponding to maximum ground state neutral lithium. Indicated on the plot are the strongest lines of valence excited Li and ground state Li$^+$ in this energy region. The weakness of both lines is a good measure of the purity of the ground state neutral lithium in the absorbing sample (b) Photoabsorption spectrum of atomic lithium in the energy window of the main 1s$^2$2s ($^2S$) $\rightarrow$ 2s$^2$2p ($^2P$) transition. The broken line represents a Fano profile fit to the restored data following removal of instrument broadening [Kennedy et al, 1996]

1.7.3 Normal incidence VUV DLP experiments

In work carried out by Jannitti et al [1985], the photoabsorption spectrum of beryllium was measured in both the normal incidence and grazing incidence regions by use of the dual laser plasma technique. To produce the two plasmas a 10 J, 15 ns (FWHM) laser was split into two parts with a variable path difference between the two beams thus allowing a variable delay between the two plasmas. The continuum
The experiments used copper. A spherical cylindrical lens focuses the laser onto the beryllium target, producing a line focus with dimensions of ~100 μm wide and up to 10 mm in length. In this way, the spreading velocities of the streaming plasma along the line of sight were minimized, and the absorbing length is maximized.

The experiments utilized a 2 meter normal incidence spectrometer system with a 1200 line mm⁻¹ spherical grating. The dispersed VUV radiation was incident on a scintillator coated face plate and imaged by two high speed objectives (NA = 0.75) external to the vacuum system on an intensified photo diode array (~0.0195 nm pixel⁻¹). The absorption spectrum close to the ionization limit of Be⁺ was studied. The photoionization cross sections of the outer electron for transitions of the type 2s → ep and 2p → ed were measured at threshold. The values thus obtained were

\[ \sigma = (1.5 \pm 0.1) \times 10^{-18} \text{ cm}^2 \text{ and } \sigma = (2.5 \pm 0.5) \times 10^{-18} \text{ cm}^2, \text{ respectively} \]

**Figure 1-6** Solid curve, experimental absorption coefficient of the resonance series \( 1s^22s^2S \rightarrow 1s^2np^2P^0 \) with \( n = 5 \text{ to } 8 \) of Be⁺ at a distance 0.6 mm. From the target surface and ~70 ns laser delay. Dashed curve, convolution of the computed coefficient with the instrument function. In the inset, the best fit of the instrumental function (solid curve) with a Voigt profile (dashed curve) is shown [Jannitti et al, 1985].

**Figure 1-7** Solid curve experimental absorption coefficient of the series \( 1s^22p^2P^0 \rightarrow 1s^2nd^2D \) with \( n = 6 \text{ to } 8 \) of Be⁺ versus wavelength at 0.6 mm distance from target. Dashed curve, convolution of the computed coefficient with the instrument function [Jannitti et al, 1985].
In some further experiments also done by Jannitti et al [1986], the photoabsorption spectrum of B⁺ was studied in the range 40 to 170 nm. The experimental set-up used was similar to one used in the Be⁺ experiments of Jannitti [1985]. The length of the absorbing plasma was increased using a cylindrical lens, up to about 18 mm. The contribution of the absorption from B III was minimized by controlling both the power density on the boron target and the delay between laser pulses. The photoabsorption spectrum of B⁺ was found to be optimized at a position of 1 to 1.5 mm from the surface of the boron target with a 80 ns inter laser delay. The values for the photoionization cross-section at threshold thus obtained were 

\[(1.7 \pm 0.6) \times 10^{-18} \text{ cm}^2 \] for the series \(2s^2 1S \rightarrow 2snp 1P^0\), and 

\[(4.0 \pm 0.5) \times 10^{-18} \text{ cm}^2 \] for the series \(2s2p 3P^0 \rightarrow 2snd 3D\).

Figure 1-8. The absorption coefficient \(k\) for (a) the series \(2s2p 3P^0 \rightarrow 2snd, ns 3D, 1S\) and (b) the series \(2s^2 1S \rightarrow 2snp 1P^0\) and for \(2s^2 1S \rightarrow 2pnd, ns 1P^0\). Solid line: experiment. Dotted line: best fit [Jannitti et al, 1986].

1.8 Concluding remarks.

In this chapter the fundamental physics of laser produced plasmas has been discussed in terms of the interaction of the laser radiation with the target material and the resulting plasma formation and primary emission processes within such plasmas. A number of applications of laser plasma broadband VUV continuum sources have been discussed. The properties of such sources include; ease of production, good
reproducibility, 'line-free' structure, high peak brightness, large spectral range, short duration, ease of source alignment and small extent. A sample of recent photoabsorption experiments in the XUV and normal incidence VUV region which utilize such sources for the backlight were briefly described.
Chapter 2. Experiment set-up.

This chapter outlines the experimental system configurations used in the spectroscopic investigations reported in chapter 3. A brief outline of the theory of the Rowland circle mount is considered and the important parameters describing the spectrometer performance such as linear dispersion and resolution are summarised. Associated instrumentation such as a Channel Electron Multiplier Array (CEMA), Photo Diode Array (PDA) and back thinned charge coupled devices (CCD) are described in terms of mode of operation and characteristics.

2.1 Overview of experimental arrangement.

The core experimental system is based on a near normal incidence spectrometer which has a focal length close to one metre. Throughout the studies two image detection systems were used. These consisted of a Galileo™ Channel Electron Multiplier Array with an EG&G™ Photo Diode Array readout and an Andor Technology™ back thinned CCD. The longest possible wavelength the instrument can reach mechanically when equipped with a grating with 1200 grooves mm⁻¹ is 325 nm at the centre of the detector plane. However as the grating is blazed at 80 nm the efficiency is observed be poor above ~ 120 nm.

A laser plasma light source was coupled to the spectrometer system by a vacuum-optical beam-line. A glass capillary array (GCA) was positioned between the plasma target chamber and the spectrometer’s slit. This reduces the amount of plasma debris that can enter the spectrometer and also allows for differential pumping across the GCA needed for experiments on gaseous absorption while still holding a vacuum of < 10⁻⁶ mBar in the spectrometer chamber.

Nearly all VUV and XUV spectrometers employ a configuration developed by H.A. Rowland at the end of the last century. Whereas the detection of short wavelengths has had major advancements in recent years by development of channel electron multiplier arrays† (CEMA) [Oschepkov et al, 1960, Goodrich and Wiley, 1960.†

† Original conceived by Farnsworth in 1930, the continuous dynode electron multiplier was implemented concurrently by groups in the USSR, USA and UK in the 1960’s.
1962, Adams and Manley, 1965, 1966] and back thinned CCD arrays [Janesick et al, 1985], the Rowland circle mount has not been changed in any significant fashion since it’s early development. The other significant developments in the field of XUV and VUV spectrometer design have been the use of toroidal optics [Tondello et al, 1975] which allow better spatial imaging of the source, holographic gratings which have resulted in more uniform groove spacing and the possibility of aberration corrected gratings and vari-spaced gratings which produce focal planes devoid of curvature and hence suitable for readout by flat detectors (e.g. CCDs).

2.2 Rowland circle normal incidence spectrometer mount

Diffraction gratings have their origins in the early 1800’s where they were first used to disperse light by T. Young and later by J. von Fraunhofer. Since then, several attempts were made to produce quality gratings for scientific studies. It wasn’t until the late 19th century, when H.A. Rowland began producing gratings with sophisticated ruling engines, that the ruled diffraction grating made an impact on the field of spectroscopy [Jobin Yvon™, 1988]. It is now over 100 years since H.A. Rowland discovered the possibility of using a concave grating placed tangentially on a circle in order to diffract and focus light which had it’s source on this circle. The diffracted spectrum is focused onto this circle which is referred to as the Rowland circle.

The associated theory may be developed using Fermat’s Least Time Theorem. There are many variations on the Rowland circle mount (i.e. Eagle, Grasshopper etc.) and each of these have relative advantages and disadvantages depending on the application. The parameters which govern the choice of mount are principally the spectral range to be covered, the resolution required, degree of astigmatism that can be tolerated as well as limitations placed on the movement of the entrance and exit arms of the spectrograph [Samson, 1967 and Namoka 1959, 1961].
2.2.1 Spherical grating equation.

Spherical grating spectrometers combine the dispersive characteristics of a grating with the focusing characteristics of a spherical surface. When limits on the movement of the entrance and / or exit arm of these spectrometers are placed, various geometries and mount are defined.

The basis of the following discussion lies in the analysis of the path length difference between two different beams reflected from two different points on a curved reflecting surface. These are assumed to originate from a point source, which is situated on the Rowland circle. If this path difference equals an integral number of wavelengths then constructive interference occurs at this point. No rigorous treatment of the optical path function is given. A more detailed discussion can be found in the work done by Namioki [1959].

If we simplify the calculations by assuming that the source and diffracted spectrum lie in the x-y plane — as in the case of the Acton™ VM 521 spectrometer (see figure 2-1), then the optical path function can be written as,

\[
F = |AP| + |BP| + \frac{kw\lambda}{d} = |IA| + |IB| + \frac{kw\lambda}{d} + \Delta \quad [\text{E 2-1}]
\]

where \( k \) is an integer and by definition is the spectral order of the diffracted image at wavelength \( \lambda \), \( w \) is the inter groove distance of the grating, \( d \) is the distance between \( I \) and \( P \) on the grating surface and \( \Delta \) is the aberration optical path function and contains coefficients which describe the defocusing, astigmatism, first and second type of coma, and others which describe the spherical aberration. \( \Delta \) can also be thought of as a perturbation on the simple optical path lying in the x-y plane and can be expressed in terms of \( Y \) and \( Z \) coefficients as follows,

\[
\Delta = FY + D \frac{Y^2}{2R} + A \frac{Z^2}{2R} + C_1 \frac{Y^3}{2R^2} + C_2 \frac{YZ^2}{2R^2} + S_1Y^4 + S_2Y^2Z^2 + S_3Z^4 + ... \quad [\text{E 2-2}]
\]

where \( R \) is the radius of curvature of the grating, \( D \) is a coefficient which characterises defocusing, \( A \) characterises astigmatism, \( C_1 \) characterises the 1st type
coma or classical coma, $C_2$ characterises the 2\textsuperscript{nd} type coma and $S_1$, $S_2$, and $S_3$ characterise spherical aberration.

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{concave_grating_diagram.png}
\caption{Diagram showing parameters used in the description of the concave grating. Point A represents the position of the entrance slit while point B represents the position of the exit slit or detector plane.}
\end{figure}

In order to obtain the location of the spectrum, the terms $F$ and $D$ are equated to zero. This will give the locus of the spectrum. The residual values for $A$, $C_1$, $C_2$, $S_1$, will give the remaining aberrations present in the image spectrum. These aberrations degrade spectral resolution, spatial resolution and / or signal-to-noise ratio. The most significant are astigmatism, coma, spherical aberrations and defocusing. The optical path function $F$ represents the permissible path lengths. Coma is the result of the off-axis geometry and is seen as a skewing of rays in the dispersion plane enlarging the base on one side of the spectral line. Spherical aberrations are the result of rays emanating away from the centre of an optical surface failing to find the focal point as
those from the centre. Astigmatism is characteristic of an off-axis geometry. It tends to have the effect of representing a point source as a line on the focal plane.

Expressing this aberration optical path function in terms of polar co-ordinates one can obtain [ Jobin Yvon™, 1988 ],

$$\Delta = Y\left[\sin(\alpha) + \sin(\beta) - kn\lambda\right]$$

$$+ \frac{Y^2}{2} \left[\frac{\cos^2(\alpha)}{l_A} - \frac{\cos(\alpha)}{R} + \frac{\cos(\beta)}{l_B} - \frac{\cos(\beta)}{R}\right]$$

$$+ \frac{Z^2}{2} \left[\frac{1}{l_A} - \frac{\cos(\alpha)}{R} + \frac{1}{l_B} - \frac{\cos(\beta)}{R}\right]$$

$$+ \frac{Y^3}{2} \left[\frac{\sin(\alpha)}{l_A} \left(\frac{1}{l_A} - \frac{\cos(\alpha)}{R}\right) + \frac{\sin(\beta)}{l_B} \left(\frac{1}{l_B} - \frac{\cos(\beta)}{R}\right)\right]$$

$$+ \frac{YZ^2}{2} \left[\frac{\sin(\alpha)}{l_A} \left(\frac{1}{l_A} - \frac{\cos(\alpha)}{R}\right) + \frac{\sin(\beta)}{l_B} \left(\frac{1}{l_B} - \frac{\cos(\beta)}{R}\right)\right]$$

$$+ Y^4[\ldots] + \ldots \quad [\ E 2-3 ]$$

where \( l_A, \alpha \) and \( l_B, \beta \) are the polar co-ordinates of \( A \) and \( B \), and \( Y^4 \) and higher order terms can be considered negligible. By equating the coefficient of \( Y \) ( i.e \( F \) ) term to zero in equation 2-3 one obtains the fundamental grating equation,

$$\sin(\alpha) + \sin(\beta) = \frac{m\lambda}{d} \quad [\ E 2-4 ]$$

where \( m \) is the spectral order and \( d \) is the inter groove distance.

### 2.2.1.1 Dispersion.

The angular dispersion, \( d\beta/d\lambda \), defines the angular separation between two wavelengths for a particular spectral order \( m \), inter groove spacing \( d \) and diffracted angle \( \beta \) and can be expressed as.

$$\frac{d\beta}{d\lambda} = \frac{m}{d\cos(\beta)} \quad [\ E 2-5 ]$$

Linear dispersion, \( d\lambda/dx \), defines the extent to which a spectral interval is spread out across the focal plane of a spectrometer and therefore is associated with
an instrument's ability to resolve fine spectral detail. Linear dispersion perpendicular
to the diffracted plane at a central wavelength $\lambda$ can be expressed as,

$$\frac{d\lambda}{dx} = \frac{d\lambda}{d\beta} \cdot \frac{d\beta}{dx} \quad [E\ 2-6]$$

$$\frac{d\lambda}{dx} = \frac{d\cos(\beta)}{mR} \quad [E\ 2-7]$$

where $d$ is the inter groove spacing, $m$ is the spectral order and $R$ is the effective
focal length of the concave grating. In the case of a McPherson™ normal incidence
type mount, the angle of incidence does not remain constant throughout the
wavelength range. In this case it is the deviation angle (i.e. the angle subtended by
the entrance slit and centre of the detector plane, $D_v = \alpha - \beta$) which remains constant
at 15° and therefore equation 2-10 must be modified to account for this. If one
substitutes $\alpha = 7.5^\circ + \varphi$ and $\beta = 7.5^\circ + \varphi$ into the general grating equation where $\varphi$
is the grating's angle of rotation, the linear dispersion perpendicular to the diffracted
plane can be expressed as,

$$\frac{d\lambda}{dx} = \frac{d\cos(7.5)\cos(\varphi)}{mR} \quad [E\ 2-8]$$

By equating the $D$ term in equation 2-2 to zero one obtains the condition for
focusing on the tangential focal plane, with the possible values for $\alpha$ and $\beta$ given by
the following.

$$\frac{\cos^2(\alpha)}{I_A} - \frac{\cos(\alpha)}{R} + \frac{\cos^2(\beta)}{l_B} - \frac{\cos(\beta)}{R} = 0 \quad [E\ 2-9]$$

Considering the last equation and it's solutions, it can be shown those if point
$A$ is on the Rowland circle then point $B$ will also be on the Rowland circle. If the
polar co-ordinates of point $A$ and $B$ are substituted in the terms for the aberrations
then it is observed that only the $C_1$ term in equation 2-2 vanishes. Therefore, on a
Rowland circle there is no 1st type coma; only astigmatism, 2nd type coma and
spherical aberration.
2.2.1.2 Wavelength and order considerations.

Because of the inherent structure of a grating there will exist multiple orders with relative intensities determined by the coating used and the blaze angle as well as the groove shape. The presence of these multiple orders degrades the performance of the spectrometer and it is sometimes necessary to incorporate filters to limit the bandpass of the system. For the same angle $\alpha$ and at the same angle of observation $\beta$ several wavelengths are superimposed. These are all wavelengths for which $m\lambda/d = \text{constant}$ or equally $m\lambda = \text{constant}$.

2.2.1.3 Resolving power.

Resolving power $R$ is a theoretical concept and is given by,

$$ R = \frac{\lambda}{\delta\lambda} \quad [\text{E 2-10}] $$

where $\delta\lambda$ is the wavelength separation of two peaks which are just resolved and $\lambda$ is the wavelength of the spectral region of interest. Applying this concept to the use of gratings, it can be shown that [Samson, 1967],

$$ R = knW_g \quad [\text{E 2-11}] $$

where in this case $W_g$ is the width of the grating illuminated with the incident radiation, $k$ is the spectral order that the instrument is being used in and $n$ is the groove density of the grating being used. Using equation [E 2-11] to define the geometry of a system with the separation of $\alpha$ and $\beta$ fixed it can be shown [Samson, 1967] that the theoretical resolution can be expressed as,

$$ R = \frac{W_g (\sin \alpha + \sin \beta)}{\lambda} \quad [\text{E 2-12}] $$

The theoretical resolving power of an instrument is defined principally by the width of the grating illuminated, the central wavelength to be resolved and geometry of the system used. According to the modified Rayleigh’s criteria, two spectral peaks are considered resolved if the ratio of minimum intensity (at the position between the peaks) to that of the maximum intensity of either peaks is less than $8/\pi^2$. 

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Although the theoretical resolving power of the spectrometer predicted by equation 2-12 is approximately 30000, the experimentally measured resolving power as measured by Acton™ is approximately 5700 @ 80 nm when using 10 μm entrance and exit slit widths and represents the maximum resolution achievable. The difference in theoretical and experimental values is primarily due to optical aberrations present and the use of entrance and exit slits of finite width.

2.2.1 4 Grating efficiency

The concentration of a limited region of the spectrum into any order other than the zero order is performed by using grooves which have a defined angle to the incident light. Blazed gratings are manufactured to produce maximum efficiency at designated wavelengths by appropriate selection of groove geometry. Blazed grating groove profiles are calculated for the Littrow condition where the incident and diffracted rays are in auto-collimation (i.e., \( \alpha = \beta \)). At the blaze wavelength, \( \lambda_{\text{blaze}} \), \( \alpha = \beta = \omega = \text{blaze angle} \) and,

\[
2 \sin(\omega) = mn\lambda_{\text{blaze}}
\]

where \( n \) is the groove density.

In the case of the Bausch & Lomb 1200 groove mm\(^{-1}\) grating which is blazed at 80 nm in first order the required angle is 2.75°. The principle is to rule the grating so that the reflecting elements are tilted with respect to the grating surface. The grating efficiency curve shifts so that the maximum efficiency is not at the zero order. Efficiency in higher spectral orders usually follows the first order efficiency curve. For a grating blazed in first order the maximum efficiency for each of the subsequent higher orders decreases as the order \( k \) increases. The efficiency also decreases the further off-Littrow the grating is used.

2.2.2 Acton spectrometer mount

The spectrometer used throughout these investigations was an 1 m focal length Acton™ model VM 521 using a 1200 lines mm\(^{-1}\) spherical concave holographic
grating with an iridium coating. The optical mount system used is based on a Rowland Circle mount in the general McPherson™ normal incidence configuration [Tang et al, 1991]. This mount is ideal for normal incidence work as both the entrance slit and exit slit or, in this case, the focal plane are fixed in space.

The principle of the McPherson™ mount is that the grating is constrained to move along the bisector of the angle subtended by the entrance slit and centre of exit plane from the centre of the grating while simultaneously the grating is rotated about a vertical axis tangent to its centre. This movement is achieved by using a cam driven stage which is machined to give the correct combination of movements for each wavelength. Fine focus adjustments to the movement can be made through the use of two micrometers external to the vacuum system.

It can be shown that the diffracted spectrum will only be focused for one value of $\phi$, where $\phi$ is the grating rotation. If the grating is rotated the diffracted spectrum will no longer be focused on the Rowland circle and therefore a linear motion is needed. This linear motion can be approximated by considering figure 2-2.

$$x + GM_0 = R \quad \text{cos}(\phi) \equiv \frac{GM_0}{R}$$

Therefore,

$$x \equiv R[1 - \cos(\phi)]$$

[ E 2-14 ]
Figure 2-2  Schematic showing rotational ($\varphi$) and translational ($x$) movement of the grating in the McPherson mount when grating is scanned from $\lambda$ to $\lambda + \Delta \lambda$.

With this off-Rowland circle type mount, the translation of the grating causes the shift of the optical axis from the centre of the grating to the slits. The displacement of the optical axis is sufficiently small to be considered negligible.

Although the limits of the grating movement allow for a possible wavelength range between 30 and 325 nm, in practice it has been observed that measurements are confined to the region where the CEMA detector and grating are both efficient. The grating efficiency profile falls off gradually in the long wavelength region while the CEMA detection efficiency falls off rapidly at wavelengths greater than ~180 nm. The use of a back-thinned CCD detector does not extend this range as the efficiency of this detector also falls off similarly. In absorption studies, the effect of order over lapping at wavelengths greater than ~75 nm means that studies are confined to the wavelength range where second order continuum emission does not saturate the measurement.
The only modifications made to the commercial VM 521 spectrometer for this project were the replacement of the existing flanges to facilitate a penning gauge to be positioned closer to the main chamber of the spectrometer and also to house an inspection window which allows viewing of alignment HeNe laser on the grating.

2.2.2.1 Aperture ratio and acceptance angle.

The aperture ratio and acceptance angle, are principally defined by the external coupling system used. Without any spatial slits or apertures and with no GCA separating the source from the spectrometer, the instrument operates as a f/10 instrument. The acceptance angle in the horizontal plane (dispersed plane) is 5.7° while the acceptance angle in the vertical plane is 3.2 °. In the case of experiments carried out in this thesis, a glass capillary array (GCA) was used to prevent plasma debris contaminating the spectrometer chamber and also to allow for a pressure differential between the target and spectrometer chambers. The GCA used had specifications which are outlined in the appendix VIII, the most important of which is the thickness of the plate (3 mm) and the pore diameter (50 μm) as well as exposed diameter (6 mm). With the limited exposed diameter, it can be shown that only single reflections within the capillaries are possible. The distance between the entrance slit and the GCA plays an important part in determining the acceptance angle of the combined system. The GCA was located approximately 212 mm from the entrance slit in experiments with direct coupling of the target chamber to the spectrometer, the acceptance angle was 17 mrads (assuming an extended source geometry). This implies that the maximum illuminated width of the grating will be 34 mm, and the theoretical maximum resolution achievable will be down proportionally. However, in reality grating aberrations will diminish the actual resolution achievable well below the grating width limit.

2.2.2.2 Bausch & Lomb™ concave grating.

The holographic concave grating used in the Acton™ VM 521 is one provided by Bausch & Lomb™. It has 1200 lines mm⁻¹ and a blaze angle of 2.75 °, which results in a first order blazed wavelength of 80 nm. The grating has
dimensions of 56 mm in height (in groove direction) and 96 mm in width and is located by a kinematic mount which facilitates interchange of gratings without realignment. The coating used is iridium, which yields a good reflectivity throughout the wavelength range used. Iridium is ideal for efficient reflection coatings at short wavelengths as it has one of the highest electron densities of all the metals [Samson, 1967]

2.2.2.3 Entrance and exit slit assemblies.

The exit slit assembly was removed and replaced by an adapter flange to hold the CEMA intensifier to allow for the multi-channel detection of the spectrum [Doyle, 1995]. The entrance slit has a vertical baffle, set at 3 mm. The entrance slit width used was generally 50 μm (dial setting), although investigations with other slit widths were performed in order to find the optimum slit width for each of the detectors used. The slit is adjustable from 5 to 3000 μm with a quoted accuracy of 10 μm [Acton™]. The slit assembly has stops which prevent the slit jaws from touching and therefore it is not possible to completely close the slit.

2.2.2.4 Spectra drive controller

The wavelength range was scanned using a precision lead screw external to the vacuum system. This was connected to a stepper motor and in turn connects to a drive controller provided by Acton™. This permits computer as well as manual wavelength selection. In the initial commissioning of the instrument the bellows assembly was found to be leaking and was therefore changed. This assembly was replaced with one which was slightly too long and therefore the dial reading on the spectral drive controller no longer represents the actual wavelength at the centre of the focal plane. Because the equation governing the translational motion of the grating (equation 2.6) is not linear, the offset is not exactly constant throughout the wavelength range. This was confirmed using carbon and aluminium spectra lines and identifying features so that a reference spectrum was obtained for the 30 nm to 220 nm spectral range.
2.2 2.5 Aberrations.

Since the spectrometer is a near normal incidence type with a 15° deviation angle there is very little astigmatism present in the final image. Most of the important work for this thesis was done using an Andor™ back-thinned CCD array detector used in full vertical binning mode over a 15 mm height. This procedure yielded no appreciable degradation in resolution when compared to a spectra recorded by a single horizontal row of pixels thus indicating the very low astigmatism and good angular alignment of the detector with the spectral image.

2.2.6 Stray and scattered light

Stray and scattered light are kept to a minimum by use of a number of baffles in the entrance and exit arms of the spectrometer. Also the use of a GCA limits the extent of the grating illuminated thus reducing the possibility of light reflecting off the grating mount itself. Stray light also results from non-perfect planeity of the reflecting surfaces. A holographic grating, which is produced by use of the interference of monochromatic radiation, does not have significant periodic imperfections in the groove profile or spacing and so generally does not suffer from ghosts as much as ruled gratings [Lerner and Theranon, 1988].

2.2.7 Order overlapping and efficiency curves

The efficiency curve of a grating shows a characteristic sharp drop off in the low wavelength limit with a lesser fall off in the high wavelength limit centred approximately around the blaze wavelength in first order (80 nm in this case). The groove shape determines the amount of radiation concentrated into a given spectral order. As a result of the optical interference inherent in the operation of a grating there exists order overlapping in which light at λ/m appears at the λ location, where m is the spectral order. Depending on the experimental study, this may be an important consideration and may require the use of filters to reduce the radiation from this higher wavelength.
2.2.2.8 Resolution.

The groove separation determines the angular separation of the diffracted beams from two different wavelengths. The theoretical maximum achievable resolution is directly proportional to the width of the grating illuminated. This maximum resolution is never physically attainable due to effects such as detector resolution and entrance slit widths used as well as aberrations present in the optics of the system. The choice of entrance slit width should be chosen so that the bandpass of the instrument is minimized and the throughput is maximized.

2.2.2.9 Wavelength and pixel position relationship

With the removal of the exit slit assembly and its replacement with a multi-channel detector such as the back thinned CCD or CEMA / PDA variety, a range of wavelengths may be viewed simultaneously in a single image collected from the detection electronics. Depending on the geometry involved in the spectrometer design, the diffraction may or may not be linear across the field of view. Under these circumstances, it is common to take reference spectra of such an element that has well-documented line structure (e.g., Kelly, 1987) in this region each time the spectral range is changed. Although the rotational movement of the grating is a linear function with respect to required wavelength, this is not true for the translational movement, and therefore the offset between dial reading and actual centre wavelength can be expected to be not entirely constant. With the use of a planar detector, the linear dispersion will not be constant across the observed range and for an accurate calibration, the points should be fitted with a quadratic expression of \( \lambda(p) \) where \( \lambda \) is the wavelength and \( p \) is pixel number for each change of spectral region.

2.3 Galileo™ Glass Capillary Array (GCA).

Use of glass capillary arrays GCA in VUV and x-ray applications have increased greatly in the last twenty years. They are efficient in transmitting VUV radiation and show focusing properties even at low wavelengths e.g., XUV and soft x-rays [Chapman et al., 1991]. They are also used in applications of gas flow collimation, in-line filtration or as calibrated leak sources [Tebo, 1994], [Wilkins et
GCAs are sometimes referred to as passive MCPs. This term comes from the fact that the bulk of the early development work was concentrated on MCP with production techniques spilling over to GCAs.

The manufacturing process starts in much the same way as MCPs. Single channels are drawn as solid-glass fibres having two components, a core glass, soluble in a chemical etchant, and a lead glass cladding which is not soluble in the core glass etchant. The fibre is then drawn out by processes much like that used in the production of optical fibres and multiple fibres are stacked together in a hexagonal structure. The array of fibres is again drawn out and stacked and the process repeated. The blocks are sliced into wafers with the channels running normal to the surface. The wafers are then ground and polished to the desired dimensions. The solid core is now etched thus creating the pore-like structure with differences in pore diameters of less than 6% [Wiza, 1979], [Lucatorto et al., 1979]. The uniformity of pore dimensions and the fact that these structures can have open areas up to 60% permits GCAs to be used as VUV and x-ray windows. They can sustain a pressure differential across their faces (the controlled leak rate typically allows for 2 to 3 orders of magnitude in pressure differential for the GCA used here) and also are able to prevent contamination of spectrometers from material used in studies.

A Galileo™ GCA having a pore diameter of 50 μm and a length of 3 mm was used in the experiments. It was housed in a custom made flange which allowed connection to 40 KF vacuum flanges at either end. The length of the housing was kept small in order to ensure that the GCA could be located close to the target chamber thereby maximizing the instrument's acceptance angle. It was held in place on a Viton® o-ring by an aluminium plate with three nylon screws which were positioned so that the GCA could be angularly aligned. The GCA was set off centre so that it could be rotated to provide a fresh area following a build up of any plasma debris. An aluminium disk with a 6 mm bore was used to mask the unused section of the GCA. The full specifications of the GCA are contained in appendix VII. A schematic of the GCA housing is shown in figure 2-13.
In order to align the GCA, its mount was coupled to the target chamber and an alignment HeNe laser was centred on the aperture. It was found that the HeNe laser could be aligned normally to target chamber side by minimising the location of the multiple reflections which occur in the rear glass plate. This was important since the GCA had only limited angular movement. The GCA was then placed in the housing and the nylon screws were gently tightened. A screen was placed a number of meters away and the spectral pattern intensity and symmetry was maximised by a process of tightening and loosening the three screws.

In order to estimate the transmission of the GCA for VUV radiation at 50 nm it is assumed that no interference effects occur and the diffraction can be considered separately from geometrical aperture effect of the capillary array. Considering a point like source at a distance, $u$, from the front face of the GCA (length $l$, pore diameter $d$ and effective diameter $D$) with uniform angular radiance. By using geometric optics, the rays that are emitted from the source can be divided into four groups:

1. Rays that are blocked by the non open area of the GCA
2. Rays that pass directly through without any reflections
3. Rays that are reflected an odd number of times from the capillary walls
4. Rays that are reflected an even number of times from the capillary walls

The rays that pass directly through without reflections can be considered collimated as there will be a maximum divergence angle $D_{max}$ which will depend on the $dl$ ratio. The intensity on a screen placed a distance $v$ from the output face of the GCA will display a Gaussian type distribution. Rays which are reflected an odd number of times will tend to result in a focused intensity distribution. The extent of the focus will depend on the $u/v$ ratio. The depth of focus will be approximately $2d$. Rays which are reflected an even number of times can be considered to result in a diffusive intensity distribution with no simple intensity distribution.

If these arguments are applied to the optical system being used, it can be shown that $D_{max} = 17 \times 10^{-3}$ radians. The GCA holder was manufactured with a 6 mm
unmasked diameter centred on the optic axis. This enabled the use of an o-ring for sealing and GCA angle alignment as well as the possibility of rotating the GCA to provide a fresh section. Therefore, the maximum angle that a ray from the source can strike the capillary walls is $18 \times 10^{-3}$ radians. This results in a maximum number of reflections of 0.17 and so only single reflections need to be considered. The reflectivity of SiO$_2$ glass at these large incidence angles ($\sim 18 \times 10^{-3}$ radians) and relatively large VUV wavelengths (~50 nm) can be assumed to be close to unity. In reality, the laser plasma has non-point like extent and so rays that originate off-axis can still possibly pass through the GCA although they do not fulfil the above criteria.

2.4 Galileo™ Channel Electron Multiplier Array (CEMA)

A single stage 12° bias angle channel electron multiplier array (CEMA) consisting of a Micro Channel Plate (MCP) with an VUV detection enhancement coating of MgF$_2$ coupled to a fibre face plate with a P20 phosphor coating was used for preliminary work. The CEMA assembly was constructed by Galileo™ and was attached to the exit arm of the spectrometer with the front face of the Micro Channel Plate (MCP) lying on the focal plane of the instrument. Applications for CEMA based intensifiers and detectors include scanning electron microscopes, residual gas analysers, image tubes and time-of-flight mass spectrometers where they are used to detect VUV, XUV, soft x-rays, neutrons or charged particles.

2.4.1 Basic construction

The construction of a single stage CEMA intensifier is shown in Figure 2-3. The MCP has an active diameter of ~25 mm and is 0.5 mm thick. It consists of pores which are 10 µm in diameter and have a centre to centre spacing of 12 µm resulting in an open area of ~55%. The pores have an electrically semi-conductive and secondary electron emissive layer. A single pore acts in the same way as a photomultiplier tube but using a continuous dynode structure of the photomultiplier tube. When a photon or charged particle is incident on the entrance face of a microchannel, secondary electrons are generated. Due to the bias applied, these secondary electrons are accelerated down the channel towards the output end. When these electrons strike
the wall, additional secondary electrons are released. This process is repeated continuously along the micro-channel and therefore the process amplifies the initial event yielding up to $10^4$ electrons at the output. Higher gains are possible if two or more of these micro-channel plates are stacked together usually in opposing angles resulting in a so-called chevron pair MCP stack. The limit is set by ion feedback problems, which result in saturation and noise. Normally the electrons are detected by use of a phosphor-covered faceplate, which converts the kinetic energy of the electrons to visible light. This visible light is usually coupled by the use of a fibre faceplate to a detector such as a photo diode array (PDA).

In applications requiring the detection of VUV radiation at the long wavelength extreme some form of electron emissive enhancement coating on the input face of the MCP is needed. Un-coated MCPs generally have quantum efficiencies of ~ 10% in the VUV with an upper wavelength limit of ~ 130 nm after which the quantum efficiency rapidly decreases. The MCP described here has a MgF$_2$ coating which extends this upper limit to ~ 180 nm.
Figure 2-3  
*Schematic of a channel electron multiplier array assembly (CEMA) showing actual micro channel plate (MCP) and fibre face plate with phosphor coating. Also shown is photodiode array PDA detector for readout*

The early part of the construction of a MCP is very much like that of a glass capillary array (GCA). The main difference is the thickness of the plate is usually smaller having a thickness of $< 1$ mm. Through further processing of the micro-channel walls, a silicon dioxide rich secondary electron emissive layer is formed on an electrically semi-conductive layer. Finally, a thin metal electrode is vacuum deposited on both input and output faces to electrically connect all channels in parallel.
2.4.1 Characteristics.

The gain of a single stage MCP can be expressed as,

\[ G = \left( \frac{AV}{2aV_0^{1/2}} \right)^{4V_0^{1/2}} \]  

where \( V \) is the channel voltage, \( V_0 \) is the initial energy of an emitted secondary electron \(-1 \text{ eV}\), \( a \) is the length to diameter ratio of a MCP pore and \( A \) is a proportionality constant \(-0.4\). For the Galileo\textsuperscript{TM} 6025-FM MCP, \( a = 40 \) and \( V = 1000 \text{ V} \) which results in a gain of \(-3 \times 10^4\). The gain was measured by Galileo\textsuperscript{TM} to be \(2.6 \times 10^4\) for the same conditions. Efficiency for the MCP is generally about 10 to 15\% for radiation in the VUV region of the spectrum. Due to the inherently high electron gain, this combination still produces a very sensitive detector. The detection efficiency falls away rapidly above 200 nm.

![Figure 2-4: Gain of a single stage MCP as a function of plate voltage \(V_0\)](image)

The limiting spatial resolution of a MCP is generally characterised by the modulation transfer function (MTF). This parameter brings together three important parameters of an imaging system. That is, resolution, contrast and sharpness of the resulting image. Using the Nyquist criterion, the limiting resolution of a MCP expressed in line pairs mm\(^{-1}\) is proportional to \(1/2\Delta p\) where \(\Delta p\) is the centre to centre...
spacing of the micro-channels in mm. The emerging electrons are then accelerated across a gap by a positive potential and proximity focused onto a phosphor coated face plate with a visible array detector readout or direct electronic readout with the use of a single anode plate, thus providing photon/particle counting.

Other factors which may limit the resolution of a CEMA are the size of electron bunch at the position of the phosphor and the size and the quality of the fibres in the faceplate used for image transfer. Considering the first of these limiting factors it may be shown that the theoretical spot size $d$ (μm) can be given by [Cromer et al, 1985],

$$d(\mu m) = \frac{s(\mu m)}{\sqrt{V_{gap}}} \quad [E\,2-16]$$

where $s$ (μm) is the inter gap spacing and $V_{gap}$ (V) is the inter gap potential. The CEMA used in the experiments considered here had an inter gap spacing of 1.2 mm and an operational inter gap voltage of 4 kV which resulted in a theoretical spot size of ~19 μm. There is no magnification by the fibre face plate (fibre centre-to-centre spacing ~10 μm) so the limit in resolution of the complete detector set-up was theoretically due to the PDA with a pixel size of 25 μm (ignoring other resolution degrading effects) which should have yielded an instrument function of 3 pixels FWHM.
In practice the instrument function was a factor of >2 worse @ 5 pixels FWHM and so the electron bunch spot was obviously significantly larger than 19 μm. Problems in the detector towards the end of experiments limited the inter gap voltage to 2.5 kV which resulted in a theoretical spot size of ~60 μm. In this case the CEMA was the limiting element in terms of resolution.

If the capacitance of the MCP can be kept to a minimum, nanosecond gating is possible with such intensifiers using basic image PDA or CCD/ICCD detectors. With electric fields of the order of 10 kV cm\(^{-1}\) and pore lengths of 0.5 mm the transit time for an electron pulse can be reduced to less than 500 ps without too much temporal spread of the electron bunch [Galileo\(^{TM}\), 1995].

2.4.2 Noise contributions in CEMA intensifiers.

Although CEMA intensifiers have excellent spatial and temporal resolution properties and unrivalled efficiency in the XUV wavelength region, they are inherently noisy with significantly higher background noise than CCD based detectors [Fraser et al, 1987]. Cosmic rays, thermal emission, field emission, ion feedback and internal radioactivity and material outgassing are all sources of noise in
CEMA based detectors In view of the very large internal surface areas of channel plates (~ 200 cm² for each cm² of intensifier area) it is reasonable to suppose that the time-dependent noise source is related to the out-gassing of the plates, whereby ions or neutrals released from the channel walls subsequently initiate avalanches. Therefore, exposure to atmosphere should be kept to a minimum and after each exposure, the MCP should be thoroughly out-gassed before operation in order to protect the intensifier. Once the intensifier has been out-gassed, the principal source of noise is the random variations in avalanche gain due to the inherently high gain and statistical process of electron gain. Also, the contribution to noise from radioactivity in the CEMA components has been shown to be significant. Radioactive impurities such as potassium ⁴⁰K and rubidium ⁸⁷Rb present in the lead glass which makes up the MCP and its casing and also in the ceramic supports are estimated to account for ~ 22% of all background noise. Cosmic ray flux is not a major background noise contribution and is estimated to account for ~ 8% of all background noise [Fraser et al. 1988, 1987]
2.5 Andor Technology™ back thinned CCD.

Experiments were also performed using a back thinned CCD borrowed from Queens University Belfast (QUB). The sensitivity of a CCD imager to light is determined by system noise levels and quantum efficiency. Quantum efficiency measures the sensor's efficiency in the generation of electronic charge from incident photons. Electron-hole pairs are produced by photons in the region from 400 to 1100 nm for front illuminated CCD. This spectral window is set by the depth of penetration of the photons into the chip and the band gap energy respectively. Light normally enters the CCD through the gates of the register. These gates are made of very thin amorphous silicon, which is reasonably transparent at long wavelengths but becomes opaque at wavelengths shorter than 400 nm. In order to extend this lower wavelength limit further so that VUV and XUV photons can be detected, the radiation must enter and pass through as little amorphous silicon gate structure as possible. This is not possible with front illumination as the gate structure is necessary for electrical connections and also because ideally the image electrons should be stored away from the interface layer.

It is possible to chemically etch the back face of a CCD after production so that the charge storage areas are effectively exposed to incoming radiation. There is thus no extra structure that the photons need to pass through before being absorbed in the vicinity of the CCD charge storage potential wells and generating charge carriers. For any given wavelength, the amount of charge stored is proportional to the intensity of the illuminating radiation. This results in an ideal detector for imaging and spectroscopy studies.

2.6 EG&G™ photo diode array (PDA).

The detector used to detect the visible output image from the CEMA intensifier was a linear photo diode array with 1024 pixels. The detector, Model 1432, was supplied by EG&G Princeton Applied Research™. The detector interface Model 1471A optical multi-channel analyser (OMA) was also supplied by EG&G Princeton Applied Research™. The size of the PDAs pixels are 25 μm * 2.5 mm with 25 μm centre to centre spacing. It has a quoted spectral sensitivity range of 200
to 1100 nm. The detector is thermally bonded to a Peltier cooler which can lower the detector temperature and therefore reduce thermally generated dark counts. With no water cooling the Peltier cooler can operate down to 5 °C. With water cooling it is possible to extend this to -19 °C. When the cooler is used for temperatures below room temperature, a dry nitrogen flow is needed in order to prevent condensation from forming inside the detector head. The analog signal is converted to a digital format using a 16 bit (with 1 bit allocated for an overflow flag) analog to digital converter (ADC) in the OMA and is available for interrogation by a computer using the GPIB interface provided [EG&G Princeton Applied Research™ literature]. See Doyle, 1995 for more details on software and interfacing.

The detector consists of an array of reverse biased photodiodes, each diode behaving like a charged capacitor. When incident light (200 to 1100 nm) enters the depletion region of the p-n junctions it has the effect of creating charge carriers – electron hole pairs – and thus decreases the effective dielectric property of the depletion layer. During each scan, the shift registers and FET switches cause the photodiodes to be successively connected to the input of the detector amplifier. Each successive junction voltage potential defines the integrated photon flux on the addressed pixel. The resulting signal, following buffering and amplification is sent to the controller interface for digitising and data processing. The output video signal is directly proportional to the incident light flux.

2.7 CCD and PDA characteristics.

2.7.1 Dynamic range of CCD and PDA detectors.

The dynamic range of a detector is the next most important parameter after the quantum efficiency. The dynamic range of a CCD imaging system is defined as the ratio of CCD saturation signal to the readout noise. A high dynamic range is more desirable especially in spectroscopic studies as it allows very weak emission lines to be seen in the same frame as very strong lines and therefore the higher intensity lines do not saturate the detector.
2.7.2 Noise contributions in CCD and PDA detectors.

The sources of noise in a CCD or PDA imaging system are predominantly from three areas. These are photon noise, preamplifier noise, and dark current noise. Photon noise, also known as shot noise, arises from the discrete nature of light production. The total number of photons emitted by a steady source over any time interval varies according to a Poisson distribution. The charge collected by a detector exhibits the same Poisson distribution, so that the noise is related to the square of the signal. Photon noise is unavoidable and is always present in imaging systems. Preamplifier noise, also called readout noise, is generated by the on-chip output amplifier. This noise can be reduced to a few electrons by design.

Dark noise current also referred to, as thermal noise is the dominant mechanism for noise in CCD and PDA based imaging systems. The electrons are generated by thermal processes and therefore cooling the detector reduces this dark current and hence the dark noise level. The back thinned CCD imaging system used in the experiments carried out for this thesis had three stages of thermoelectric cooling by which it is possible to cool the detector to a temperature of $-90 \, ^\circ C$. At these temperatures the detector performance rivals that of the more expensive liquid nitrogen cooled CCDs with a quoted dark current of 40 electrons pixel$^{-1}$ sec$^{-1}$ at a temperature of $-20 \, ^\circ C$ (Andor Technology\textsuperscript{TM} InstaSpec IV technical manual). The PDA also had a thermoelectric cooler which allowed for the detector head to be cooled to $-40 \, ^\circ C$ (see page 57) resulting in a quoted dark current of 100 electrons pixel$^{-1}$ sec$^{-1}$ (EG&G\textsuperscript{TM} 1453A operating manual).

With the exception of a few full frame images, the back thinned CCD measurements were taken using full vertical binning of the frame pixels. This increases the SNR (signal to noise ratio) of the detection system but can reduce resolution of spectroscopic data if image aberrations are present or if the detector is tilted with respect to the lines.

Several sources of noise are present when the signals stored by a PDA or CCD are read out. They can be classified as either random noise or fixed pattern noise. The fixed pattern noise in principle can be removed either by the electronics or
by later data processing. Random noise arises from the electronics (primarily the input section of the pre-amp circuit) and from the statistical nature of the arrival of photons at each pixel. In general, reset noise $n_{\text{reset}}$ (also referred to as kTC noise) will dominate the signal and may be expressed as,

$$n_{\text{reset}} = \left( \frac{1}{q_e} \right) k_BT \cdot \sqrt{C_{\text{pix}}} \]  

where $q_e$ (C) is the charge on an electron, $k_B$ (1.381*10$^{-23}$ J K$^{-1}$) is Boltzmann’s constant, $T$ (K) is the absolute temperature and $C_{\text{pix}}$ (F) is the capacitance of a single pixel element [Howes and Morgan, 1985]. The total reset noise is larger due to the multiple readout switching and therefore the $C_{\text{pix}}$ should be replaced by $2*(C_{\text{pix}} + C_{\text{video line}})$ where $C_{\text{video line}}$ is the video line to clock capacitance. Considering the PDA used here with $C_{\text{pix}} = 2$ pF, $C_{\text{video line}} = 1.2$ pF and $T = 288$ K, the reset noise is approximately 997 electrons and is by far the dominant noise component within the detector. The noise due to the preamplifier $n_{\text{preamplifier}}$ can be expressed as,

$$n_{\text{preamplifier}} = \left( \frac{C}{q_e} \right) 4k_BT \Delta B \cdot (k_e/gm) \]  

where $\Delta B$ (Hz) is the bandwidth of operation, $k_e/gm$ is the channel resistance of the input FET and $C_{\text{input}}$ (F) is the capacitance at the preamplifier input which is composed of the series combination of $C_{\text{video line}}$ (F), $C_{\text{preamplifier}}$ (F) and $C_{\text{stray}}$ (F). In the case of EG&G™ PDA where $\Delta B = 10$ MHz, $T = 288$ K, $C = 10$ pF and $k_e/gm = 100$ the noise contribution from the preamplifier will be approximately 250 electrons over the entire detector array.

The dark current shot noise can be expressed as,

$$n_{\text{dark}} = \sqrt{\frac{I_{\text{dark}} t_f}{q_e}} \]  

where $I_{\text{dark}}$ (A) is the equivalent dark current and $t_f$ (s) is the readout time. Typically, the equivalent dark current will be 1 pA and so a 30 ms readout time will
result in a dark noise of approximately 433 electrons averaged over the entire array. All of the above noise sources combine to give an effective total noise \( n_{\text{total}} \) which is superimposed onto the actual detected image.

With the Galileo™ CEMA / EG&G™ PDA detector system the dominant type of noise present in spectral signals is fixed pattern noise (FPN) which can be reduced considerably by subtraction a background noise signal from the spectral signal. Also it is worth pointing out that the PDA described in this work had a dc offset count (\( \sim 45 \) average counts) which is necessary for correct operation of the ADC and is present on all signals recorded. Random fluctuations in all components of the detection system (e.g., CEMA gain, laser energy etc.) can be reduced by averaging a number of shots (typically 10).

2.8 JK Lasers™ ruby laser

A JK Lasers™ ruby laser operating in Q-switched mode was used to generate the plasma in most experiments reported in this thesis. The laser was of the single oscillator type and provided \( \sim 1 \text{ J} \) (\( @ 1.85 \text{ kV capacitor charging voltage} \)) with a pulse length of 35 to 45 ns at a wavelength of 694.3 nm. The ruby rod consists of a crystal of aluminium oxide \( \text{Al}_2\text{O}_3 \) in which some of the \( \text{Al}^{3+} \) ions were replaced by \( \text{Cr}^{3+} \) ions. The lasing medium was pumped by a Xenon flash-lamp. A Pockels cell combined with a polarizer was used to provide Q-switching. The Pockels cell is normally biased to prevent lasing and the high voltage is removed to allow for an intense pulse to be generated.
The laser used in the initial experiments was a Spectron Laser Systems™, Q-switched Nd:YAG SL803 laser featuring a two stage oscillator/amplifier configuration. This system yielded an output pulse energy of ~ 690 mJ in ~ 15 ns @ 1064 nm and was used for producing the continuum plasma. Also used was a Q-switched Nd:YAG SL404 laser which did not have an amplifier stage. This laser was predominantly used for the production of the absorbing target plasma and had an output pulse energy which was measured to be ~ 170 mJ with a pulse duration of 15 ns @ 1064 nm.
The lasing medium consists of a rod of crystalline yttrium-aluminum-garnet (YAG, $\text{Y}_2\text{Al}_5\text{O}_{12}$) doped with neodymium (Nd$^{3+}$) ions. The crystal is pumped to a state of population inversion by a flash-lamp close coupled in a diffusion cavity. A Pockels cell/polarizer combination is used to inhibit normal lasing action until a Q-switch trigger pulse is received. This can be either internally or externally triggered. A Stanford pulse delay generator with 5 → 15 V step-up switches is used to fire all flashlamps and Pockels cells in synchronisation for laser pulse timing on a nanosecond timescale.

2.10 Target chamber and vacuum system

For the study of electromagnetic radiation at wavelengths, less than 200 nm it becomes necessary to work under vacuum conditions due to the significant attenuation of radiation by atmospheric gases. The spectral region $\lambda < 200$ nm is generally referred to as the vacuum ultra violet (VUV). Below $\lambda \approx 0.2$ nm the atmosphere becomes transparent again and therefore wavelengths of about 0.2 nm can therefore be regarded as the lower wavelength limit of the VUV. One can further divide this spectral range into sub regions depending on the type of spectrometer required. For example, the region between about 30 nm and 200 nm has been termed the normal incidence region, and the region between about 3 nm and 30 nm is quite...
often referred to as the grazing incidence region. It must be stressed that no formal definitions of spectral ranges in the VUV exist and the regions do overlap somewhat.

![Diagram of spectrometer arrangement](image)

**Figure 2-8**  *Schematic of direct coupled spectrometer arrangement showing main dimensions between optical components*

### 2.10 Vacuum pumps

A number of vacuum pumps were used throughout the experiments to pump the spectrometer and target chambers. The target chamber was pumped using a turbo mechanical and rotary pump combination. The spectrometer was pumped using a turbo mechanical pump, with the ion pump taking over once a pressure of $10^{-5}$ mBar was reached. A full schematic of the vacuum arrangement used is shown in Figure 2.9.
Typical operating pressures were $9.7 \times 10^{-7}$ mBar in the spectrometer chamber and $7 \times 10^{-6}$ mBar in the target chamber. The pressure differential between the spectrometer and target chamber was primarily across the GCA. It is possible to increase this differential pressure e.g., for gaseous absorption studies, of up to $\sim 10^2$ for steady state and higher for small periods of time (i.e., less than $\sim 3$ mins). If a CEMA based intensifier is being used, it is important that the pressure in the spectrometer is never allowed to increase beyond $2 \times 10^{-6}$ mBar. With the use of the back thinned CCD, no limitation is imposed, but the pumps require a minimum of $10^{-5}$ mBar at the inlet ports for correct operation and to prevent back streaming of oil from the rotary pumps.

2.10.2 Vacuum component design

The target chamber consisted of an aluminium cube of edge 126 mm with each face drilled to provide holes of 68 mm in diameter. This resulted in a chamber
with vacuum connections on all six sides. The target chamber was supported by an
in-house constructed base facilitating attachment of the TPH060 turbo pump and
pressure gauges. The entrance window consisted of a 6 mm thick BK7 glass plate
held in place by a collar with an o-ring seal between the plate and the target chamber.
A dual target holder flange suitable for connection to the target chamber was
designed and constructed (see figure 2-12). The flange allows translational (both
across the optic axis and vertically) as well as rotational movements of the target
holder. The vacuum seal on each of the target holder bars was made through the use
of a Wilson seal. This resulted in a target chamber which was vacuum compatible
with the spectrometer with or without the GCA in place. A GCA holder was
designed and built (see figure 2-11) which allowed the GCA to be both rotated and
positioned close to the target chamber. The holder incorporated a mask which
allowed an unused section of the GCA to be rotated into place when plasma debris
built up. The holder was 50 mm in length and facilitated connection to 40 KF flanges
at either end.

In order to couple the back thinned CCD camera to the spectrometer an
intermediate flange was constructed. This was constructed from aluminum and had
an o-ring seal on one face. The thickness of this flange was machined to 57 mm so
that the focal plane of the spectrometer would be coincident with the front face of the
detector. Replacement flanges for the spectrometer were constructed which allowed
the attachment of a pressure gauge close to the spectrometer’s chamber and
incorporation of a window which made it possible to view the grating. The ability to
view the alignment HeNe laser footprint on the grating with the system under
vacuum proved crucial in the alignment of the system.
Figure 2-10  Schematic of adapter flange for use with Andor Technology™ InstaSpec IV back thinned CCD. Dotted line indicates spectrometer’s focal plane.

Figure 2-11  Schematic of GCA holder.
Figure 2-12 Schematic of target holder flange showing the two target holders. Centre to centre spacing between target holders 47 mm. Maximum movement possible in x-axis direction 15 mm. Hole size for continuum target 30*22 mm. Hole size for absorber target or spatial slit/aperture 30*20 mm.

Figure 2-13 Schematic of CEMA/PDA detector arrangement [Doyle, 1995]
Chapter 3  Results and discussion.

Experiments carried out with the use the Acton™ normal incidence VUV spectrometer are described here. Various system configurations and detectors were used during these experiments and are highlighted at the beginning of each relevant section. The results include emission studies of plasmas with line or continuum dominated spectra, VUV photoabsorption by helium gas, and also photoabsorption studies with the dual laser plasma (DLP) technique. The chapter concludes with results on spatially resolved YBCO (YBa$_2$Cu$_5$O$_{7.8}$) plasma VUV emission.

3.1  Line dominated laser plasma spectra  carbon and beryllium oxide

Line dominated spectra are useful for characterising a spectrometer instrument in terms of wavelength calibration, resolution, sensitivity and dynamic range. In addition they are useful in plasma diagnostics, e.g., to estimate peak plasma temperature by observing the peak ion stage present in the spectrum. If the wavelength resolution is such that spectral lines display their true line width (i.e., instrument function width is much less than typical plasma line width) then estimates of the electron densities can also be made. Line dominated plasmas also serve to optimize detector alignment and focusing.

3.1.1  Carbon emission spectra

Carbon is an ideal reference spectrum for use in the VUV region. It has strong discrete features, which can have narrow line widths under appropriate plasma conditions and which can make it possible to estimate the system wavelength resolution. Most lines of the CI to CVI spectra have been classified [Kelly, 1987 and NSRDS-NBS 60] and hence accurate estimates of the plasma ion stages present and also some plasma parameters (e.g., electron temperature) can be made. Because the known lines extend well into the UV and VUV spectral regions this allows estimates of the degree of order overlapping present and combined grating and detector efficiency over the wavelength range to be made. Figures 3-1 and 3-2 show the observed multiple laser shot (corresponding to the accumulation of ten spectra from ten successive laser shots) emission spectrum of a polyethylene laser produced.
plasma with both the Andor™ back thinned directly illuminated CCD and the Galileo™ CEMA / EG&G™ PDA detectors. Both spectra show strong line dominated emission with very little underlying continuum. In the case of the CEMA / PDA detector the width of the narrowest spectral lines observed as measured at full width half maximum (FWHM) is 6 to 7 pixels (linear dispersion: 0.021 nm pixel⁻¹), while with the improved spatial resolution with the back thinned CCD means that the same spectral lines are much narrower at about 2 to 3 pixels (linear dispersion: 0.022 nm pixel⁻¹).

From a comparison of figure 3-1 and 3-2, it can be seen that the relative intensities of the emission lines are different for the two detector arrangements. This can be attributed to different plasma conditions such as on target irradiance and position of the plasma relative to the spectrometer’s optical axis as well as differences in detector efficiency. The CEMA / PDA polyethylene emission spectrum of figure 3-1 shows improved signal at wavelengths greater than about 120 nm than that obtained with the use of the back thinned CCD detector.

All wavelengths mentioned are actual calibrated values unless otherwise indicated. The spectrometer’s dial was found to be ~5.9 nm (approximately constant over the 30 to 325 nm range) lower than actual centre wavelength.
Figure 3-1 Single shot polyethylene (C₄H₈) plasma emission spectrum observed with the Acton™ 1 metre normal incidence spectrometer in the spectral range 20 to 220 nm. A Galileo™ CEMA based intensifier with an EG&G™ PDA detector was used to record the spectral signal. The spectrometer slit width was set at 50 μm. Laser Ruby ~ 1 J, 25 – 35 ns (FWHM) @ 694 3 nm.
Figure 3-2. Single shot polyethylene (C,H) plasma emission spectrum observed with the Acton™ 1 metre normal incidence spectrometer in the spectral range 20 to 220 nm. An Andor Technology™ back thinned CCD was use to record the spectral signal. The spectrometer slit width set at 50 μm. Laser : Ruby ~ 1 J, 25 – 35 ns (FWHM) @ 694.3 nm.
3.1.2 Beryllium oxide emission spectra

The laser produced plasma emission from beryllium oxide is dominated by intense oxygen lines in the normal incidence VUV region. The near complete absence of beryllium spectral lines in this region makes this material ideal for observing the pure oxygen spectrum. As a surface contaminant, oxygen lines are usually present in spectra of most metal plasmas, especially for the first few laser shots and the ability to identify these impurity lines in subsequent recorded spectra is important. The single shot sensitivity of the PDA / CEMA detector arrangement for the study of emission spectra in the wavelength range 30 to 220 nm is clearly seen in figure 3-3 where the spectral signal is well above the underlying noise. Most of the lines were identified using reference tables (Kelly, 1987 and NSRDS-NBS 60) and correspond to transitions in ion stages from O+ to O4+. Second order lines begin to dominate the spectrum at wavelengths greater than ~100 nm while third order lines begin to dominate at wavelengths greater than ~150 nm. There are a significant number of lines which could not be identified despite attempts at matching with transitions in lower ion stages of beryllium or the impurity element of carbon. Their origin remains an open question.

The oxygen emission spectrum is such that most of the observed spectral lines are the result of a number of close lying transitions which are not resolved in first order with the instrument used. However some of the more intense lines are seen up to fourth order and the increased resolution results in some of the components of such lines being resolved.

The order overlapping present in spectra recorded on the Acton™ instrument is also illustrated in figure 3-3. The poor efficiency of the grating at wavelengths below about 48 nm means that second order light is only significant at wavelengths greater than approximately 95 nm and third order light greater than approximately 170 nm. The decrease in efficiency of the grating at wavelengths greater than 100 nm and the decrease in intensity of the plasma emission results in very few first order lines observable at wavelengths greater than 95 nm.
Figure 3-3. Single shot beryllium oxide emission spectrum observed with the Acton™ 1.0 metre normal incidence spectrometer in the spectral range 20 to 220 nm. A Galileo™ CEMA intensifier with an EG&G™ PDA readout was used to record the spectral signal. The spectrometer slit width was set at 50 μm. Laser: Nd : YAG, ~ 690 mJ, 15 – 25 ns (FWHM) @ 1064 nm.
3.2 Continuum dominated laser plasma spectra: samarium and tungsten

The continuum-dominated emission from high atomic number targets has already been discussed in terms of importance in photoabsorption studies and other applications such as photoelectron spectroscopy and VUV reflectivity measurements. The common requirement of these applications is not so much a 'line free' but more a reproducible continuum. Studies on the extent of the continuum observable with the Acton™ VM-521 spectrometer as well as structure, flux and reproducibility are discussed in the following sections.

3.2.1 Samarium continua dominated emission spectra

A samarium (Sm, Z = 62) plasma, formed using a cylindrical shaped target with diameter 6 mm was used for continuum generation in most of the experiments described here. Samarium is a lanthanide group metal which has a silvery-white appearance. It is relatively stable in air but oxidises on contact with moisture. It is quoted as being harmful and noxious and hence may involve limited health risk if ingested or if it penetrates the skin.

Laser produced samarium plasmas generally exhibit line free continuum emission over the range 4 to 200 nm with weak line emission superimposed on the continuum mainly around 125 nm. This means it is an ideal backlight for use in normal incidence VUV absorption experiments. It was also chosen for these experiments because in the spectral region of interest for resonant absorption by helium gas (~ 50 nm) the intensity of the samarium plasma emission has a maximum within the normal incidence wavelength range [Carroll et al., 1983].

A full scan of the samarium spectrum throughout the range of the Acton™ VM 521 spectrometer was performed using the Andor™ back-thinned CCD detector. The output from the back-thinned CCD reached $2.5 \times 10^4$ counts @ 45 nm for the accumulation of 10 shots with the spectrometer slit width set at 50 μm, corresponding to $2.5 \times 10^3$ counts pixel⁻¹ per laser shot or ~ 13% of pixel saturation.
Figure 3-4  Schematic spectrum of samarium showing the calculated approximate position of the strongest transition arrays [O'Sullivan, 1983].

Figure 3-5  Samarium continuum emission observed by the Acton™ VM 521 normal incidence spectrometer in the range 20 to 220 nm. An Andor Technology™ back thinned CCD was used to record the spectral signal. The spectrometer slit width was set at 50 μm. Laser Ruby ~ 1 J, 25 – 35 ns (FWHM) @ 694.3 nm.
3.2.2 Tungsten continuum dominated emission spectra.

The XUV continuum from a laser produced tungsten plasma is ideal for use in photoabsorption experiments as it has a reproducible and reasonable 'line free' structure. In the longer wavelength normal incidence VUV region the number of discrete lines present increases and arise from transitions within lower ion stages of the tungsten plasma as well as impurities present such as oxygen. Although the presence of these lines can degrade the quality of photoabsorption measurements taken with such continua in practice the overall structure is reproducible. Hence it tends not to affect transmission ($I/I_0$) measurements. Although samarium has less visible line structure superimposed on the underlying continuum and the overall intensity is greater in the normal incidence VUV region, samarium is a softer material and therefore the position of the target must be changed more frequently than that of tungsten. Also the fact that the samarium target used in the experiments described here was of a cast form and hence was not perfectly cylindrical, when both targets are rotated to provide a fresh surface the tungsten target will deviate less from its previous position.
Figure 3-6  Tungsten continuum emission as observed by the Acton™ VM 521 normal incidence spectrometer in the range 20 to 220 nm. A Galileo™ CEMA based intensifier with an EG&G™ PDA detector was used to record the spectral signal. Spectrometer slit width set at 50 μm Continuum plasma excitation laser Nd YAG ~ 690 mJ, 15 – 25 ns (FWHM) @ 1064 nm

3.2.1 Accuracy and precision of mechanical movement of grating

In order to quantify the reproducibility of the wavelength scanning mechanism for the Acton™ VM 521 spectrometer a series of spectra were recorded over the range 30 to 200 nm in steps of 15 nm with the grating moved over the full range in between scans. Inspection of the pixel position of spectral lines showed that there were no relative peak shifts thus indicating that the scanning mechanism was indeed reproducible to within the pixel dimension of 25 μm. Also initialising the stepper motor drive (i.e. turning the drive off and on) had no effect on the pixel positions over the entire range. Although manual movement of the scanning mechanism to zero order (dial position ~ 9933) resulted in a shift in the pixel position of a number of lines by one pixel, this shift was not observed with all lines.
and therefore is more likely to do with changes in line structure and profile due to source fluctuations.

This shift of less than a single pixel (~ 0.02 nm) confirms the reproducibility quoted by Acton™ of 0.005 nm. The dial reading no longer determines the centre wavelength position accurately and there is an offset which varies slightly over the complete range of the instrument. Although the equations governing the motion of the grating (equation 2-10 and 2-11) result in a non-linear rotation, the mechanical movement consisted of a sine drive which translates the cam driven stage linearly. In the range 20 to 220 nm the offset was measured to be 5.899 ± 0.022 nm. It is therefore noted that so long as the detector arrangement is not interfered with, calibration spectra need only be taken once. The removal and replacement of the PDA detector can result in an offset of the centre pixel up to ~ 40 pixels.

The Galileo™ CEMA / EG&G PDA detector arrangement displayed over a small region at the lower wavelength end of the spectra with depressed pixel counts, indicating that a portion of the detector was shadowed. This region extended for approximately 40 pixels from the edge of the detector and was more evident when photoabsorption measurements were taken where the calculation of the relative photoabsorption cross section yielded a high value within this region. As the recorded spectra are usually taken with an overlapped wavelength coverage of ~ 50% or so, these shadowed pixels may be simply ignored.
As can be seen from the graph the actual wavelength is 5.9 nm longer than that of the dial reading.

3.2.2 Wavelength resolution comparisons between the two detectors.

In figure 3-8 the emission spectrum of a polyethylene plasma is shown in the limited wavelength range for both detectors where the improvement in wavelength resolution achievable by the Andor™ back thinned CCD is clearly seen. The vast majority of spectral lines recorded by the Andor™ back thinned CCD are at most 3 to 4 pixels in width (FWHM) whereas with the Galileo™ CEMA / EG&G PDA the spectral lines are at most 6 to 7 pixels in width. The wavelength resolution of either detector is not sufficient to resolve the spectral lines to their natural widths. This is confirmed by comparing second and third order lines to their first order counterparts where no increase measured line width is visible.
Figure 3-8  Single shot polyethylene \( (C_4H_8) \) plasma emission spectra observed with the Acton\textsuperscript{TM} 1 metre normal incidence spectrometer using both (upper two rows) an Andor\textsuperscript{TM} back thinned CCD and (lower two rows) a Galileo\textsuperscript{TM} CEMA/EG&G\textsuperscript{TM} PDA detector. Proceeding from left to right the same region is displayed in first, second and third order.
3.2.3 Quantum efficiency and dynamic range comparisons between the two detectors.

The efficiency of the grating with respect to wavelength has not been investigated as this would require a calibrated source at least of known relative intensity. Therefore any discussion of the quantum efficiency of the detectors must include a discussion on both the efficiency of the grating as well as radiance of the source. The detected spectra from a specific detector arrangement, $I_{\text{Detected}}(\lambda)$ will be a function of the spectral radiance of the laser plasma source, $I_{\text{LPP source}}(\lambda)$, the efficiency of the grating, $G_{\text{Grating efficiency}}(\lambda)$, depending on both the reflectance of the coating as well as blazed wavelength and that of the detector, $D_{\text{Detector quantum efficiency}}(\lambda)$. In addition a geometric factor, $F_{\text{Geometric factor}}(\lambda)$, accounting for the entrance optics of the spectrometer, position and shape of the plasma source is also present.

$$ I_{\text{Detected}}(\lambda) = I_{\text{LPP source}}(\lambda) * G_{\text{Grating efficiency}}(\lambda) * D_{\text{Detector quantum efficiency}}(\lambda) * F_{\text{Geometric factor}}(\lambda) \quad [E\ 3-9] $$

It is evident from emission spectra that the efficiency of the grating determines to a significant degree the overall shape of the continua and relative heights of lines. The short wavelength falloff in efficiency is rapid while the longer wavelength is more gradual. The plasma intensity, $I_{\text{LPP source}}$, tends to peak at shorter wavelengths and compounds this difficulty in determining the efficiency curve of the grating. With most of the line and continuum dominated emission spectra the peak counts are observed to be at $\sim 60$ nm. The relative strength of the spectral lines of polyethylene, aluminium and beryllium oxide in second order tend to be as intense as their first order counterparts principally around $\sim 80$ nm. This suggests that the blaze wavelength of the grating within the Acton™ spectrometer is indeed 80 nm in agreement with the Acton™ specifications.

The quantum efficiency of the two detectors also has an effect on the observed counts for both the continuum and line dominated emission spectra. The Galileo™ CEMA intensifier with the MgF$_2$ coating has a quoted quantum efficiency of 10% when detecting normal incidence VUV photons ($\lambda < 1800$ nm). It is also
evident that the quantum efficiency of the back thinned CCD falls off at lower wavelengths than the CEMA / PDA detector combination. The back thinned CCD displays higher dynamic range and lower noise.

Generally speaking, the description of the system sensitivity is rather crude because many of the factors involved, such as gain and efficiency of the detection device, are strong functions of other parameters. Since the gains of both the MCP and of the phosphor screen are steep functions of applied voltage, and the CCD dark noise also depends on operating temperature, the estimate of the sensitivity is more qualitative than quantitative.

3.2.4 Reproducibility of continuum emission from rare earth plasmas

In figure 3-10 the effect of a large number of shots has on the average intensity of the tungsten plasma continuum at a centre dial wavelength of 30, 40, 50, 60 and 70 nm with and without a 525 μm spatial resolving slit introduced 35 mm from the source is shown. Once the position of the tungsten plasma has been optimized the average counts are seen to gradually increase over several hundred shots and then gradually decrease. The centre of the plateau for each wavelength is observed to tend towards more shots with increasing wavelength. Although the number of data points is limited, the inclusion of the spatially resolving slit has the effect of reducing the slope at which the average counts increase at, while not severely affecting the overall intensity.
Figure 3-8  Reproducibility of continuum dominated emission spectra of a tungsten plasma generated with a Nd: YAG laser, ~ 690 mJ, 15 to 25 ns (FWHM) @ 1064 nm  Number of laser shots averaged for each recorded spectrum  10 Black lines represent data collected without a secondary slit present Red lines represent data collected with a secondary slit present

These observations can be justified by considering the extent of the emitting plasma and the fact that as the focused laser forms a crater increasing in depth with each subsequent laser shot. Initially the crater is likely to confine the plasma keeping it hot and hence bright. However as the crater deepens the hot plasma core becomes increasingly occluded and the detector signal starts to drop. The secondary slit reduces the amount of scattered light entering the spectrometer system and also reduces the sensitivity of the system to fluctuations in source size.
3.3 Helium photoabsorption measurements.

These experiments were carried out with the use of the Andor™ back-thinned CCD and with the target chamber directly attached to the spectrometer entrance slit assembly. The GCA was positioned a distance of 170 mm from the target source and 360 mm from the entrance slit. The total absorption path length defined by the GCA to plasma source had dimensions of 170 mm. A full schematic of the experimental arrangement is shown in figure 3-12. The spectrometer entrance slit width was set at 50 μm throughout the experiments.

The target chamber was connected to the region between the GCA and entrance slit via a 40 KF flexible coupling. A 40 KF gate valve was inserted in this line to enable isolation of the chamber from this region when leaking helium gas into the target chamber. This allowed the small amount of helium gas that leaked through the GCA during an absorption experiment to be pumped out between experiments. Also, it was important not to disturb the alignment of the GCA when the target chamber was vented. This was only possible if there was no pressure difference on either side of the GCA and was achieved by opening the 40 KF value. The spectrometer was pumped using a Balzers™ TPH240 turbo mechanical pump and the pressure was relatively stable at 3.5 * 10^{-5} mBar throughout the experiments.

The laser used to generate the back light continuum source was a JK Lasers™ Ruby laser operating in Q-switched mode. It generated a pulse with a duration of ~ 30 ns (FWHM) at a wavelength of 694.3 nm with an energy of ~ 1 J per pulse. The laser was tightly focused via a plano-convex lens (focal length 105.0 mm, diameter 22 mm) to an estimated spot size of ~ 150 μm in diameter, thus achieving a power density of ~ 3 * 10^{11} W cm^{-2}.

Helium is a much-used gas for absorption measurements. It has well known cross sections in the VUV region of the spectrum. This makes it ideal for characterising a VUV spectrometer instrument. Helium exhibits strong absorption (with a relatively small absorbing column length) by the 1s^2 (^1S) → 1snp (^1P) Rydberg series before the absorption K edge at 50.43 nm.
Samarium was chosen for its relatively 'line free' continuum emission in this region as well as the fact that it is substantially brighter than tungsten in this limited region. In addition, samarium displays an average counts profile which has its broad peak in the 40 to 60 nm region. Although samarium is not completely 'line free' in the VUV, the spectrum below ~ 100 nm is composed of broad structures which are very reproducible with suitable experimental conditions. A measure of the reproducibility of the intensity of the continuum emitting plasma was made by both monitoring the resulting spectrum as well as observation of the average counts of the detector in successive recorded spectra without the absorbing gas present.

The position of the continuum emitting samarium plasma was optimized by maximizing the average counts of the entire PDA detector array while adjusting the target position holder (translation across the optic axis, x) and focusing lens assembly (three degrees of freedom, x, y and z). The output energy of the laser used to generate the continuum emitting plasma depended both on repetition rate as well as delay between experimental runs and so the continuum emission also displayed poor reproducibility. Figure 3-9 displays a number of the recorded continua. Although the shape of the continuum was similar with each shot the intensity varied over a range approximately ~ 20%. Despite attempts at making the continuum emission more reproducible by incorporating a slit between source and spectrometer and also maintaining the delay between experimental runs constant the reproducibility was not improved significantly. Wavelength calibration was performed using the emission spectrum from a polyethylene plasma in a cylindrical rod form. This resulted in an emission spectrum with well identifiable carbon emission lines across the entire detector plane.
Figure 3-9. (upper plot) Reproducibility of continuum dominated emission spectra of a rare earth, samarium (Z = 62), plasma for sum of ten laser shots at the center wavelength of 52 nm used in the helium photoabsorption experiments. All of the spectra are within ~20% of the average counts (lower plot) ratio of recorded spectra indicating good reproducibility of features. Laser used: Ruby, ~1 J, 25 to 35 ns (FWHM) @ 694.3 nm.

Helium gas was taken directly from the laboratory piped supply with a quoted purity of 99.993%. A precision leak valve was used to vary the flow of helium into the target chamber. The target chamber was pumped continuously throughout the experiments using the Balzers™ TPH040 turbo mechanical pump. A relatively stable helium pressure throughout each experiment was achieved by adjusting the leak valve. The range of pressure for each experiment was recorded. Although a small volume of gas leaked through the GCA to the spectrometer side of the GCA, it was found that the amount of helium was so small that it resulted in negligible additional absorption. Therefore the length of the helium absorbing path was well defined as the distance from the continuum plasma source and the face of the GCA.
Figure 3-10  (a) Continuum emission from a samarium plasma, (b) absorption of the continuum emission by helium gas and (c) relative photoabsorption cross section in the wavelength range around the K absorption edge at 50.43 nm corresponding to direct photoionization at the 1s² S limit (Residual helium gas pressure 1.4 * 10⁻⁴ mBar, column length 170 mm) The features circled are the result of changes in backlight emission
The dramatic K absorption edge at 50.43 nm is clearly seen in both first and second order in figure 3-11 (a) and (b). In first order (λ_{centre} = 52 nm), the reciprocal linear dispersion is ~ 0.022 nm pixel⁻¹ at the detector plane and therefore the observed range is from ~ 41.0 nm to ~ 63.3 nm. The helium photoabsorption lines observed are 1s²¹S → 1snp¹P⁰ where n = 2 to 8. In second order the reciprocal linear dispersion is ~ 0.011 nm pixel⁻¹ and therefore the observed range is from ~ 46.9 nm to ~ 57.1 nm. The helium photoabsorption lines observed are 1s²¹S → 1snp¹P⁰ where n = 3 to 9.
Figure 3-11. (Left image) Image showing full vertical binned image of helium photoabsorption, I, with the use of a samarium laser produced plasma backlight in first order 41.8 to 62.2 nm and (right image) in second order 93.8 to 114.2 nm. Parameters: Absorbing length: 170 mm., Residual helium gas pressure: 2.0 - 2.7 mBar in first order, 2.7 - 3.0 mBar in second order. Absorption lines observed \( n = 2 \) to 8 in first order and \( n = 3 \) to 9 in second order.
By calculating the number density of the residual gas species, cross section measurements can be performed on photoabsorption data. This allows valuable information of system performance to be collected specifically on effects such as multi-order and scattered light contributions as well as wavelength resolution of the entire spectrometer and detection system.

The absolute cross section $\sigma$ of a particular species is defined by the following expression,

$$ \ln \left( \frac{I_0}{I} \right) = \sigma nl $$ \[ E\ 3-3 \]

where $\sigma$ (m$^2$) is the absolute photoabsorption cross section, $n$ is the number density of the absorbing species and $l$ (m) is the column length of the absorbing species. The number density can be calculated from the following expression,

$$ PV = nRT $$ \[ E\ 3-5 \]

where $P$ (Pa) is the absolute pressure, $V$ (m$^3$) is volume of the vacuum chamber containing the gas (estimated to be $\sim 3.5 \times 10^{-3}$ m$^3$), $n$ is the number of moles, $R$ is molar gas constant (8.314 J K$^{-1}$) and $T$ (K) is the absolute temperature of the gas. The cross section should not vary as residual gas pressure changes, indicating a linear relationship between $\ln(I_0/I)$ and the column length. Figure 3-12 shows the calculated cross section versus the residual helium pressure at a number of different wavelengths after the K absorption edge.
Figure 3-12. Experimentally calculated relative photoabsorption cross section of helium after K absorption edge versus residual helium gas pressure.

From figure 3-12 it is evident that the slope of the relative photoabsorption cross section, $\ln(I_0/I)$, versus pressure is only linear for a limited portion of the graph. The non-linearity of the graph at relatively high helium gas pressures indicates that the absorption saturates before complete absorbed signal decreases to zero. The saturation should be insensitive to poor detector resolution and is more dependent on both stray light and detector noise. The poor reproducibility of the backlight emission results in discrete features being present in the relative photoabsorption, $\ln(I_0/I)$ plots and a significant offset in baseline signal level.

The principal series running up to the K edge is seen to be largely independent of pressure indicating that the vast majority of data obtained was indeed saturated. The lines do not increase in absorption or increase in width for increasing pressure. Any measurement on the relative photoabsorption cross section for the series is therefore not possible. Stray light, detector resolution and detector noise are all thought to play a part in the saturated signal. Stray light in the form of multiple order light present and also scattered light within the spectrometer system will reduce the saturation limit in photoabsorption experiments. The resolution of the detector
used to record the dispersed spectrum also affects the saturation limit. As the gas pressure increases, the relative photoabsorption cross section of the principal series does not increase substantially indicating that these features saturate at lower residual gas pressures.

The blaze of the grating used in the Acton™ spectrometer is such that it exhibits a significant drop in efficiency at wavelengths greater than ~120 nm. Considering that it is blazed at 80 nm and that second order helium photoabsorption results display strong absorption, it is clear that the underlying continuum is truly weak at ~100 nm and the observed spectrum is principally second order. This is illustrated well in figure 3-12 (b) where the K edge and Rydberg series are clearly seen in second order.

In addition, it is noted that the non-zero step in the second order spectra is due to residual first order light at ~100 nm passing through the gas with only a small amount of residual absorption. The contribution of second order continuum light at ~100 nm is estimated to be ~62% and therefore the weak first order continuum signal prevents photoabsorption studies at wavelengths much greater than ~100 nm.

Also it is observed that the width of the photoabsorption lines do not change with increasing residual gas pressure or become any more resolved in second order where the linear dispersion is effectively doubled. This observation suggests that the photoabsorption features are truly narrow in width and it is not possible to resolve them.

Also the importance of the effect of saturation of photoabsorption measurements is clearly illustrated in figure 3-11 where the data points on the experimentally extracted photoabsorption cross section deviate from a linear relationship with increasing residual gas pressure. This is due to a number of effects such as stray light, presence of multi-order light, detector spatial resolution and detector dynamic range.
3.4 Dual laser plasma barium 5p photoabsorption measurements

In order to characterise the suitability of the system for photoabsorption studies using the dual laser plasma technique, measurements on the 5p photoabsorption spectra of a barium plasma were made. This was the first time that the dual laser plasma technique for photoabsorption was performed on the Acton™ 1 meter normal incidence spectrometer at the DCU laboratory. The experimental arrangement is illustrated in figure 3-13. All measurements were made using the Galileo™ CEMA / EG&G™ PDA detector.

The lasers used to generate the continuum emitting plasma and absorbing plasma were both Q-switched Nd:YAG laser's (15 – 25 ns FWHM @ 1064 nm). The laser used to generate the continuum plasma had an energy of ~ 690 mJ (Spectron™ SL803) while that used to generate the absorbing plasma had an energy of ~ 160 mJ (Spectron™ SL404). The output of the SL404 Nd:YAG laser was focused to a spot size of ~ 500 μm with a plano convex lens of focal length 105 mm and diameter 22 mm in order to generate an absorbing plume of principally neutral barium. The output of the SL803 Nd:YAG laser was tightly focused with a plano convex spherical lens of focal length 100 mm and 22 mm in diameter to a spot size of ~ 150 μm. The burn pattern for the SL803 laser at the lens position was observed to be symmetrical (10 mm diameter) while the burn pattern for the SL404 laser at the target lens position was very elongated (3 * 10 mm²) due to poor internal alignment of the laser. Subsequent investigation showed damage to an output coupler in the SL404 to be responsible for poor beam quality.
Figure 3-13
Schematic of experimental arrangement used in DLP experiments.
Dimensions between optical components are shown in mm. Lens assembly not shown.

Position of CEMA / PDA detection combination
Connected by 1 meter flexible vacuum coupling.

Position of back continuum source
Position of front target

Position of GCA in holder

Grating curvature
Rowland circle

Entrance slit of spectrometer
Action 1 meter spectrometer

Spectron SL300 Nd:YAG laser
160 mJ 15 to 25 ns FWHM @ 1064 nm

Spectron SL800 Nd:YAG laser
690 mJ 15 to 25 ns FWHM @ 1064 nm

To photo diode and fibre optic collecting visible plasma emission
Barium is an alkaline earth metal having an atomic weight of 56. It has a silvery-white appearance and oxidises readily in air. The VUV photoabsorption spectra of atomic and ionic barium have been investigated in work done by Kent et al. [1987] and also by Lucatorto and McIlrath [1980] amongst others. It has been found that the 5p photoabsorption spectra of barium and its ions display a rich variety of lines. Some important physical processes have been first seen in barium e.g., double ionisation anomaly [Brehm and Hofler, 1975]. It also is important for the reason that the compound YBCO (YBa2Cu3O7-δ) is actively being investigated as a high temperature superconducting material. It is also worth noting that the laser plasma ablation technique has proved itself ideal in producing thin films of such compounds.

Barium possesses a closed outermost sub shell and therefore its photoabsorption spectrum should be relatively simple. Instead of what should be a simple Rydberg series converging on just two series limits according to the one electron excitation scheme, \(5p^66s^2^1\text{S}_0 \rightarrow 5p^56s^2^2\text{P}_{1/2,3/2}\) ns, nd the 5p spectrum of Ba I is found to consist of 14 series converging on 12 limits [Lucatorto and McIlrath, 1980], [Kent et al., 1987]. The 5p photoabsorption spectrum of atomic barium is therefore one of the richest inner shell excitation systems. The large number of extra series encountered in the barium spectrum can be pictured as a 'shattering' of the 6s\(^2\) sub shell by 5p excitation [Connerade, 1978]. In effect 6s\(^2\), 5d6s and 5d\(^2\) configurations lie close in energy and exhibit significant configuration mixing allowing photoexcitation to 5p\(^6\)6s\(^2\)nl, 5p\(^5\)6s5dnl and 5p\(^5\)5d\(^2\)nl series from the 5p\(^6\)6s\(^2\) (\(^1\text{S}_0\)) ground state. The corresponding Ba\(^+\) ion-core yields the 12 observed limits [Rose et al., 1978].
Figure 3-14. The 5p sub shell photoabsorption spectrum of Ba I showing how the ordering into Rydberg series is accomplished [Rose et al, 1978]. The range extends from approximately 48 to 79 nm.

In the experiments discussed here, the barium plasma producing laser was defocused and the barium plasma itself was positioned a distance from the optical axis in order to optimize the barium atomic population in the absorbing plasma plume. The wavelength range studied was between 48 and 68 nm and allowed for all of the series limits of the 5p photoabsorption spectrum to be observed within one wavelength setting of the grating.

It was found that the front plasma emission, $I_{\text{front plasma}}$, was of sufficient intensity that it needed to be subtracted from the absorption spectrum signal before the relative photoabsorption cross-sections could be derived. In order to reduce the collection of front plasma emission the barium planar target was rotated to make an angle of $\sim 5^\circ$ with the optic axis of the spectrometer. In this way it was possible to
partially occlude the hot (VUV emitting) core of the barium plasma. The absorption coefficient is given in general by the expression,

\[ k(\lambda) = \exp\left(\frac{I_0(\lambda) - I_{so}(\lambda)}{I(\lambda) - I_{front-plasma}(\lambda)}\right) \]  

[ E.3.4.A. ]

where the spectrum of the background continuum source, \(I_0(\lambda)\), and the absorption spectrum, \(I(\lambda)\), are corrected by the contributions from stray light, \(I_{so}(\lambda)\), during the measurement of \(I_0, I_s(\lambda)\) from both plasmas during the measurement of \(I(\lambda)\) and for the emission, \(I_{front-plasma}(\lambda)\), from the absorbing plasma. The contribution of stray light, presence of higher order light and light entering the spectrometer without passing through the absorbing plasma all result in an additional ‘baseline’ signal on the detector video out signal. Hence, features in photoabsorption measurements will saturate before complete absorption. Stray light was not calculated but assumed to be negligible as the spectrometer had a number of baffles and the illuminated width of the grating was significantly smaller than the actual width due to the aperture defined by the GCA. Due to the poor grating efficiency below ~30 nm, no significant second order light is expected in the 30 to 60 nm spectral range of interest here.

A small rectangular section of barium (approximately 4(w) * 6(h) * 10(l) mm) was cut from the cast ingot which was of spectroscopic purity. The barium sample was clamped in place by a small c-shaped machined holder. Barium oxidises readily when exposed to air thus forming a relatively loose barium oxide layer. This oxide layer was cleaned while under vacuum by firing a few loosely focussed laser pulses onto the barium target. The distance from the continuum plasma and the absorbing plasma was approximately 32 mm along the optic axis.

The CEMA intensifier was operated with a gap voltage significantly less than the recommended 4 kV so that the wavelength resolution achievable was degraded somewhat. This was unavoidable as higher voltages (greater than 2.5 kV) led to arcing across the gap. Although the resulting theoretical electron spot size on the phosphor within the CEMA intensifier (given by equation 2-10) is a factor of 12 higher than with normal operating gap potentials, it is found in practice that this is
not the limiting factor in spatial resolution of such a detector arrangement. Emission spectra of polyethylene and other line-dominated plasmas displayed no significant linewidth increases when using the lower gap potential (perhaps one pixel or so).

The synchronization of the Q-switched lasers was achieved by the use of a multi-channel Stanford Research™ time delay generator. This triggered both the flash lamps and Pockels cells in the two Nd:YAG lasers with a variable time delay between the Pockels cell triggers for the leading absorbing plasma generating laser and that of the following continuum emitting plasma generating laser. The TTL level signals from the delay generator were converted to 15 V level signals suitable for the triggering of the lasers using step-up transistor circuits. The master trigger was provided by the OMA controller and also was used for triggering storage oscilloscopes used to monitor the laser pulse profiles, inter laser pulse delays (with the aid of an optical fiber and fast photodiode, e.g., BPX-65) and also the OMA video signal (see figure 3-13).

A pair of 45° Nd:YAG mirrors located on optical rails were used to direct the two Nd:YAG laser beams onto focusing lenses located outside the target chamber. The collected samarium plasma emission was optimized by maximizing the observed average counts of the entire PDA detector array while adjusting the target position holder (translation across the optic axis, x) and focusing lens assembly (three degrees of freedom, x, y and z). It can be seen from figure 3-10 that the samarium continuum has a reproducibility of ~2% using a Nd:YAG SL803 laser over the restricted wavelength range 48 to 68 nm used in most of the barium photoabsorption experiments.
Figure 3-15. (upper plot) Continuum dominated emission from a samarium plasma produced by a Spectron™ SL803 Nd : YAG laser (~ 690 mJ in 15 – 25 ns FWHM @ 1064 nm) showing good reproducibility and lack of discrete features and also (lower plot) ratio of pairs of I₀ spectra indicating good reproducibility of features.

The barium target was moved inwards towards the optic axis of the spectrometer until it began to occlude the continuum emission which was accepted by the spectrometer. This point was taken to correspond to zero deviation of the target surface from the optic axis of the system. It was then pulled back by ~ 0.5 mm (referred to as Δx) and a series of photoabsorption spectra were taken for various time delays.
In figure 3-16 (a) the continuum emission from a samarium plasma is displayed in the region of interest for the barium 5p photoabsorption spectrum. The continuum is essentially ‘line free’ with the two strongest line features most likely to arise from the lower ion stages of the samarium plasma. The absorption is displayed in figure 3-16 (b) for an inter laser time of 104 ns and $\Delta x = 0.5$ mm with the absorption plasma producing laser de-focussed by 5 mm. In figure 3-16 (c) the front plasma emission is displayed with no target rotation. A dark spectrum was not subtracted from this particular data set and so the relatively large offset from zero counts is not all continuum emission from the barium front plasma alone. Some of the strong emission lines display self absorption and there is also signs of absorption of the continuum emission generated by the barium plasma.
Figure 3-16 (a) Continuum dominated emission from a samarium plasma in the region of interest 48 to 68 nm, (b) barium absorption spectrum ($\Delta t = 104$ ns and $\Delta x = 0.5$ mm), (c) barium front plasma emission only and (d) relative photoabsorption cross section.
The relative photoabsorption of barium is displayed in figure 3-16 (d) for the experimental conditions already given above. The spectrum is characteristic of that produced by the absorption of predominantly neutral barium. However comparing the spectrum with that taken by Lucatorto et al., 1980 it is evident that some Ba\(^+\) and Ba\(^{++}\) is also present. The relatively strong photoabsorption feature (within the region between 51 nm to 57 in figure 3-10 (c) ) at 55 548 nm corresponds to the Ba\(^{2+}\) resonance transition of \(5p^6 \, ^1S \rightarrow 5p^5d^1 \, ^2P_{1/2}\).
Figure 3-17  Photoabsorption of predominantly neutral barium in the wavelength range 48 to 68 nm for various different inter laser time delays. The barium sample was positioned 0.5 mm back from the start of cut-off (see text). The continuum plasma was produced by a Q-switched Nd: YAG (~690 mJ, 15 - 25 ns FWHM @ 1064 nm) focused onto a samarium target. The barium absorbing plasma was produced by a Q-switched Nd: YAG (~160 mJ, ~15 ns FWHM @ 1064 nm)
Figure 3-18  Barium photoabsorption spectra in the range 50 to 57 nm as recorded by Lucatorto and McIlrath [1980]. Upper spectrum Ba, middle spectrum, \( \text{Ba}^{+} \), lower spectrum \( \text{Ba}^{2+} \). Densities and temperatures are the same for all spectra. Indicated wavelengths are in Angstroms.

In figure 3-17 the time resolved spectra show the absorption of predominantly neutral barium around the limits of the 5p photoabsorption. The region around 56.5 nm displays the most intense relative photoabsorption cross-sections which compares to that result of Lucatorto and McIlrath [1980]. The general features of the absorption spectrum agree with the full absorption spectrum as taken by Connerade et al. [1978]. The reduced resolution of the Acton™ VM-521 spectrometer makes it difficult to compare the results directly to those of Kent et al. [1987]. At inter laser delays of ~70 ns the dominant ion stage is observed to be \( \text{Ba}^{2+} \), \( 5p^6 1S \rightarrow 5p^5 2P_{1/2} 5d[3/2] \), at 55.548 nm with few other absorption features present. At this early stage in the plasma formation the high temperature and density facilitates the production of \( \text{Ba}^{2+} \).
Figure 3-19. Graph of relative absorption for the resonance transitions of Ba$^{2+}$: $5p^6 1S \rightarrow 5p^5 5d^1 2P_{1/2}$ at 55.55 nm and neutral Ba at 57.72 and 60.48 nm versus inter laser delay.

In figure 3-19 the relative photoabsorption cross-section is plotted against inter laser delay for identified transitions. The relative photoabsorption cross-section is seen to peak at ~ 500 ns for nearly all the spectral lines. The temporal resolution is determined by the duration of the continuum emitting plasma which is approximately ~ 200 ns in the normal incidence VUV region with the Spectron™ Nd : YAG laser used in the experiment [ Rydén, 1993 ].

The results presented in this section serve to demonstrate the system’s ability to perform relative photoabsorption cross section measurements on a plasma plume using the DLP technique. Use of a secondary slit between the absorbing plasma and continuum emitting plasma would limit the extent of the source emission at the absorbing plasma position as collected by the spectrometer thereby improving spatial resolution. The increased spatial resolution would further improve the selection of an individual ion stage and also help to discriminate against front plasma emission which is predominantly from the hotter core. The experimental arrangement facilitates temporal photoabsorption measurements to be performed on the secondary
plasma plume. It must be pointed out that the duration of continuum emission from rare earth plasmas increases with wavelength and therefore the temporal resolution possible in normal incidence VUV experiments using the above arrangement are limited to approximately 200 ns [Rydnén, 1993]. Use of a gated MCP would of course reduce this to the chosen gate width (typically ≤ 10 ns) and would also serve to reject a substantial amount of front plasma emission.
3.5 \textit{YBa}_{2}\textit{Cu}_{3}\textit{O}_{7.5} emission.}

For a number of years, \textit{YBa}_{2}\textit{Cu}_{3}\textit{O}_{7.5} (YBCO) as a high $T_C$ superconductor has led interest in laser deposition techniques for the generation thin films of such materials. The ion distribution, density and temperature as well as expansion velocity of the plume and presence of ambient oxygen atmosphere are all thought to determine the resulting thin film properties [Mehlman et al., 1993]. Several techniques including optical emission spectroscopy [Fried et al., 1991 and Dyer, 1990], absorption spectroscopy [Geohegan and Mashburn, 1989], laser induced fluorescence [Fried et al., 1994], direct imaging [Wazzan et al., 1996 and Geohegan, 1992] and mass spectroscopy [Wiedeman and Helvajian, 1991] have been used to study the plasma parameters following laser ablation.

The current section is concerned with presenting spatially resolved VUV emission spectra from a laser produced YBCO plasma plume. Spatial resolution was achieved by placing a slit of width 525 $\mu$m, aligned with the spectrometer slit some 47 mm from the plasma source and 335 mm from the entrance slit (see figure 3-15). As an aid to identifying spectral features in the recorded YBCO spectra, the emission spectra of some of the constituent elements in the compound, namely barium (Ba, Z = 56), copper (Cu, Z = 29) and oxygen (in the form of beryllium oxide, BeO) were also recorded. Yttrium was unavailable at the time of the experiment. Emission spectra were recorded using a Spectron™ SL404 Nd : YAG laser ($\sim$ 163 mJ 1064 nm, 15 – 25 ns FWHM) focused to an estimated spot size of 3 mm diameter resulting in an on target irradiance of $\sim$ 1.2 * 10$^8$ W cm$^{-2}$ or a fluence of $\sim$ 2.3 J cm$^{-2}$. The number of laser shots integrated to produce each spectrum record was ten.

The experimental arrangement was similar to that illustrated in figure 3-1 with the continuum emitting target being replaced by a sample holder which allowed for the attachment of the YBCO [supplied by TCD], barium, and copper planar targets as well as carbon and aluminium. For recording oxygen plasma spectra a cylindrical beryllium oxide target was used which also had the secondary purpose of calibrating the spectrometer. Although the laser ablation and deposition of YBCO compounds is usually done in an oxygen rich atmosphere (100 to 200 mTorr), the
following results are concerned with the emission of a plasma expanding into vacuum

The YBCO spectrum displays a continuum dominated structure with a shape partly determined by the grating efficiency profile. The observed emission lines are mostly oxygen up to the fourth ion stage with a small number of the discrete features from barium and copper present also. The intense emission from barium is thought to make up most of the underlying continuum signal. Although not studied here, the high atomic number target yttrium (\(Y, Z = 39\)) is expected to produce predominantly continuum emission. The copper emission displays a low underlying continuum with low average counts with three visible broad features centred at approximately 68, 75 and 85 nm. A small number of oxygen emission lines are also evident in the copper emission spectrum.
Figure 3-20  

Emission spectra of YBCO sample taken with Acton™ 1 meter normal incidence spectrometer in the range 20 to 220 nm without the use of a spatial resolving pre slit. Nd YAG laser ~ 169 mJ, 15 to 25 ns @ 1064 nm was used with an on target irradiance of ~ 12 * 10^8 W cm^2. In addition, shown underneath are individual element spectra of barium, copper and beryllium oxide taken at a tight focus condition.
With no spatially resolving slit the acceptance angle of the spectrometer is such that almost all the emission from the plasma is collected. With the introduction of the slit it was possible to isolate a region of the plasma plume with dimensions of ~580 μm in the horizontal plane. No additional spatial resolution in the vertical plane is obtained. Figure 3-15 shows the arrangement specific to this experiment.

**YBCO planar target**
- Spatial resolving slit
  - Width: 525 μm
  - Height: 15 mm

**GCA**
- Pore diameter: 50 μm
- Pore length: 3 mm
- Clear aperture: 6 mm

**Spectrometer slit**
- Width: 50 μm
- Height: 3 mm

---

Figure 3-21 Schematic of positions of the source, spatial resolving slit, GCA and spectrometer slit used in the recording of spatially resolved emission from a laser produced $\text{YBa}_2\text{Cu}_3\text{O}_y$ plasma.

In order to characterize the spatial resolution of the system with the introduction of the pre-slit the YBCO target was temporarily replaced with a cylindrical tungsten target. The tungsten target was irradiated by ~690 mJ Nd:YAG using a tight focus at ~45° and the resulting continuum emitting plasma was aligned to the optical axis of the spectrometer by maximizing the average counts at a central wavelength of ~39 nm. The spatially resolving slit was now introduced by simple rotation of the slit holder and emission spectra were recorded for various positions of the spatial resolving slit. The position of this slit was set at the location which yielded optimum flux and remained at this optimized position for the remainder of the experiment. The extent of the emission from the tungsten plasma was recorded by moving the target relative to the fixed slit in 0.5 mm increments and recording the average counts at each position. Considering the geometry of the arrangement a maximum dimension of approximately 145 mm in diameter is placed on the...
tungsten plasma emission at a wavelength of $\sim 39$ nm with the portion emitting the most of this emission being somewhat smaller than this dimension.

**Figure 3-22** Average counts of tungsten emission plasma at a central wavelength of $38$ nm versus relative horizontal position of tungsten target with spatial resolving slit $525$ $\mu$m in width at a distance of $47$ mm from tungsten source.

The tungsten target was now replaced with the planar YBCO sample and the position of the plasma emission was optimized (both vertical and horizontal directions) so that maximum average counts were obtained. By moving the YBCO target horizontally the emission from a spatially resolved chord $\sim 580$ $\mu$m was recorded at various distances from the target surface (see figure 3-16). The transition from continuum dominated emission close to the target surface to line dominated emission at the cooler regions further from the target surface is clearly seen in figure 3-17.
Figure 3-23  Spatially resolved YBCO emission spectra at various distances from target surface (2.5, 3.5, 4.5 and 5.5 mm) The spatially resolving slit (of ~525 μm in width) was positioned a distance 47 mm from the source. The GCA to source distance was 170 mm and the spectrometer slit to source distance was 382 mm.
The observed spectra at distances of $\geq 3.5$ mm from the target are seen to be dominated by oxygen lines, OII, OIII, OIV and OV, superimposed on a weak continuum with the most intense being OIV. A limit of 5.5 mm was placed by the target holder movement. The integration time for the PDA detector was set at 30 ms and therefore the spectra are the integrated emission within this time. Some of the more intense lines show signs of self absorption resulting from the optically thick plasma plume at distances close to the YBCO target.
Chapter 4 Further system development.

The present chapter discusses the design, construction and part characterisation of a vacuum-optical beam line which serves to focus the plasma plume onto the entrance slit. The development has the following benefits, (1) One can study emission from various parts of the plasma plume by moving the plasma image across the spectrometer entrance slit, (2) in conjunction with an iris of known diameter, a section of the absorbing plasma plume with known conical geometry may be selected in DLP experiments and (3) an increased throughput is obtained over the arrangement used in experiments to date. Although no spectral data could be obtained using this set-up because the concave mirror available had a surface coating which resulted in an extremely poor reflectivity in the normal incidence VUV range, work to date is included simply to highlight the advantages of such an addition. Estimates of the increase in throughput and spatial resolution are presented using both simple geometric arguments as well as Monte Carlo simulations using the software package ‘Shadow’ developed by Lea et al [1986] at the Center for X-ray Optics, Wisconsin University.

4.1 Introduction

Spectroscopic studies of laser produced plasmas with some form of spatial resolution and/or imaging qualities are important as they can reveal the true nature of laser generated plasmas, i.e. dynamic systems with parameters which vary with space and time (e.g. electron and ion density and temperature gradients). Laser generated plasmas generally have symmetry about the axis perpendicular to the target plane and so simple space resolving pre-slits can isolate in a limited section of the plasma plume to be spectroscopically studied. Furthermore, if an entrance slit with limited vertical extent (i.e. pin hole) is used and a two dimensional array detector plane is positioned at the vertical focus, the resulting system can achieve monochromatic spatial imaging of the emission source with a one-to-one correspondence between points on the source plane and points on the detection plane. As the system works effectively as a pin hole camera, spatial imaging with magnification of the source is possible [Koog et al, 1996 and Richards et al, 1980]
4.2 Addition of VUV focusing optic.

The Acton™ VM 521 spectrometer is inherently a large f-number instrument (approximately f/10 in the horizontal plane when equipped with 96 * 56 mm grating) and so without any apertures or slits the instrument is unable to discriminate between emission from various regions of a plasma. The spectrometer effectively integrates all of the available VUV emission from the plasma which falls within the large acceptance angle of the instrument.

Although this experimental arrangement results in a relatively high throughput of flux for extended sources, it nevertheless means that experimental studies are limited to those that do not require spatial information. With the addition of the GCA the situation is somewhat changed and the acceptance angle of the combined spectrometer system is considerably reduced to an effectively f/60 instrument. As the plasma plume is of small extent and is positioned a distance of ~382 mm from the entrance slit of the spectrometer (a consequence of the need for a GCA and support vacuum components) the combined system does not allow for space resolved studies of the plasma plume without the use of a secondary slit.

Spatial resolution can be important for photoabsorption studies as the dominant ion stage can be selected somewhat by passing the continuum radiation through a specific portion of the plume and onto the entrance slit of a spectrometer. The use of a slit allows for the collection of light from a vertical chord of the plasma plume. Usually slits reduce the amount of light coupled into an optical system considerably and therefore the signal to noise ratio of the recorded spectrum suffers. Hence, it is important that the system be as close to aperture matched as possible.

Using a detector with smaller pixel dimensions would increase the resolution capability of the instrument but the reduced entrance slit width necessary may reduce the spectral signal to a level where un-intensified detectors may be insufficient to provide an adequate dynamic range. Changing the grating to one which has a higher groove density...
would also increase the wavelength resolution achievable but at the expense of the extent of the available spectral range which is determined by the maximum mechanical movement of the grating.

Considering the fact that only a small region of the maximum spectral range is observable for reasons such as choice of the blazed wavelength, order sorting problems and availability of a suitable backlight continuum the replacement of the grating with one which has a higher groove density may be the most appropriate way of increasing the wavelength resolution of the instrument.

4.3 Vacuum-optical design

In designing the vacuum system to couple to the existing spectrometer instrument the main requirements are for a UHV mirror chamber with a mirror holder enabling angle and focusing adjustments to be made external to the vacuum system. A near normal incidence arrangement for the additional optic was chosen to reduce the effect of aberrations. The entrance and exit arms of the mirror chamber were therefore set at 15° between them. A concave focusing optic with focal length of 500 mm and diameter 50.8 mm with an UV enhanced aluminum coating (i.e., aluminum with MgF2) was initially used. It was thought to be feasible to use such a coating in the VUV (λ ≥ 30 nm) even though the coating is more suited to longer wavelength UV radiation and no reflectivity measurements below λ = 200 nm were available. A schematic diagram showing the main components of the vacuum-optical beam line is displayed in figure 4-1.
Figure 4-1: Schematic of system arrangement with attached focusing optic and vacuum system. Dimensions between optic elements are shown in mm.

- F/Number 20
- Diameter 50
- Coating: Alum + MgF2/Iridium

Spherical concave mirror:
- Focal length 500 mm
- Diameter 50.8 mm
- Coating: Aluminum + MgF2/Iridium

Entrance slit of spectrometer:
- Height 3 mm
- Width 50 µm typically

Dimensions:
- 1000 mm
- 35 mm
- 88.5 mm
- 2.18 mm
Spectrometer's concave grating
Grooves / mm 1200
Concave radius 995 4mm
Coating Iridium
Blaze angle 2.75
Blaze wavelength 80μm (First order)
Ruled width 96mm
Ruled height 56mm

Exit of spectrometer
MCP and PDA detector

- Backlight plasma
- Target plasma
- Glass capillary array
- Optional variable aperture
Ideally, the grating should be illuminated to both its full vertical (56 mm in the direction of the rulings) and horizontal extent (96 mm) without overfilling which would cause stray light problems. With the small extent of the laser produced plasma light source and the necessary positioning of such a source a distance from the spectrometer entrance slit some form of aperture matching optics are needed for this criterion to be achieved. The further the source is positioned away from the slit, the smaller the width of the grating that is illuminated. The throughput also decreases significantly and is proportional to $1/r^2$ where $r$ is the distance from the source to spectrometer entrance slit. Although the resolution of the spectrometer is theoretically linearly proportional to the illuminated width, it is found in practice that other factors can effect the limiting resolution such as optical aberrations, entrance slit width and detector spatial resolution. Reduction of the spectrometer slit below a certain value in practice gave no appreciable increase in resolution achievable although the throughput was severely decreased. An entrance slit width of 50 $\mu$m when imaged in the dispersion plane is at best $\sim$ 2 pixels (FWHM) in the case of the back thinned CCD (25 $\mu$m pixel dimension). Therefore considering the fact that an array detector such as this needs greater than 2 pixels to provide an accurate representation of the spectral line the observation can be justified.

With the addition of the focusing mirror, the plasma light can be focused onto the entrance slit of the spectrometer thereby increasing the throughput of the complete spectrometer system compared to that arrangement used in experiments to date. Also, it is possible to study emission from various parts of the plasma plume by moving the plasma image across the spectrometer’s entrance slit.

Even though the very process of achieving a spatially resolved experimental arrangement inherently reduces the aperture matching of the instrument, it is important that the maximum flux from this selected section of the light source be collected by the spectrometer. The GCA, with its intrinsically small acceptance angle limits the extent of the grating which can be illuminated when using a focusing optic. On the other hand, in the direct-coupled system the small size of the emitting plasma is the limiting factor. Although the reflectivity of most materials
(non multi-layer) in the extreme VUV region is poor, the reduced reflectivity is compensated by the fact that the collected VUV radiation is focused onto the spectrometer slit and so even though the flux is reduced the amount of flux entering the spectrometer is increased.

However, after the experiment it became apparent that the available mirror did not possess good reflectivity in the 30 to 80 nm spectral range of the grating and was not sufficient to improve the light gathering ability of the combined system over the direct coupled system. In view of this, an iridium-coated mirror with the same dimensions and focal length is presently on order and this will directly replace the existing mirror.

Two flexible couplings (length 94.5 ± 6 mm) were used at both entrance and exit arms of the mirror chamber (see figure 4-1). These had the primary purpose of allowing for slight misalignments in the vacuum flanges and also served to allow both vacuum arms to be assembled without moving the mirror chamber, target chamber or spectrometer from their aligned positions.

A manipulator which permits alignment of the mirror external to the vacuum system was designed and is illustrated in figure 4-2. It consists of two parts, an aluminum flange which connects to a 100CF flange suitable for connection to the mirror chamber rear port (part A) and a stainless steel plate with three Wilson type seals to allow the three 6 mm diameter silver steel bars which could rotate as well as move in and out (part B). One of these bars is rigidly attached to the gimbals optical mount while the other two connect by the use of small chucks to the fine pitch alignment screws of the mirror mount. Therefore rotation of the bars external to the vacuum system allows for angular deflections of the attached mirror in two orthogonal planes while translation of the third bar when rigidly fixed to the other bars moved the whole mirror assembly towards or away from the plasma source and entrance slit. The aluminum flange had a groove on both sides which facilitated a 4.5 mm o-ring. The 100CF flange makes use of a copper gasket seal to provide a vacuum connection to the mirror chamber. The stainless steel plate is not rigidly connected to the aluminum flange and is able to slide on the o-ring seal thus allowing the mirror to be moved. Horizontal and/or vertical.
(x, y-axis) movement of the mirror assembly is facilitated by 4 drive screws placed at 90° to each other.

![Diagram of mirror manipulator]

**Figure 4-2** Schematic of mirror manipulator with focusing (z-axis), angle and optic axis (x, y-axis) alignment adjustments. The mirror holder facilitates mirrors with 50.8 mm diameter.

### 4.4 Basic flux calculations

Some preliminary calculations were performed in order to estimate the increase in flux with the additional focusing optic incorporated into the system. It has been shown that the spectral radiance emitted from laser produced plasma using Q-switched lasers of ~1 J per pulse is approximately $2 \times 10^5 \text{ J cm}^{-2} \text{ sr}^{-1} \text{ s}^{-1} \text{ nm}^{-1}$ @ 120 nm emitting into $4\pi$ steradians [Carroll *et al.*, 1983]. This in turn corresponds to a flux of $1 \times 10^{23}$ photons cm$^{-2}$ sr$^{-1}$ s$^{-1}$ nm$^{-1}$. The normal incidence VUV duration of a plasma is generally accepted to be a longer than the XUV pulse which lasts for approximately the same duration as the laser pulse. Work carried out by Rydén.
[1993] on the Acton™ VM-521 spectrometer with photomultiplier based detector indicated that continuum emission from tantalum plasmas produced by a Nd YAG laser (~ 1 J, 15 to 25 ns duration FWHM @ 1064 nm) had a time duration of ~ 200 ns (FWHM) in the long wavelength region (λ ≥ 200 nm) of the normal incidence VUV spectrometer. Hence there are ~ 2.42 * 10^{16} photons cm^{2} sr^{-1} nm^{-1} in a 200 ns (FWHM) VUV pulse. Assuming the source has an effective diameter of ~ 1.45 mm at this wavelength, one obtains ~ 4 * 10^{14} photons sr^{-1} nm^{-1}.

### 4.4.1 Original system

The effective acceptance angle of the GCA (defined by the l/d ratio 1/60 radians) is ~ 2/60 radians. The GCA has an effective optical transmission of ~ 30%, principally due to the loss in flux by occlusion of the rays. The entrance slit of the spectrometer is located 212 mm from the exit face of the GCA. Taking the plasma source to be of point-like extent and located 170 mm from the GCA, the source footprint on the spectrometer's slit will be ~ 13 mm in diameter. Therefore, the flux entering the spectrometer through a slit of 50 μm in width and 3 mm in height is ~ 9.8 * 10^{6} photons nm^{-1}.

This flux is incident onto the grating which has an iridium coating with an estimated reflectivity of ~ 20% at the blaze wavelength of 80 nm and therefore the flux focused at the dispersion plane of the instrument with a linear dispersion of ~ 0.83 nm mm^{-1} is ~ 1.6 * 10^{6} photons mm^{-1} or equivalently,

~ 4.1 * 10^{4} photons per pixel,

assuming a detector with pixels 25 μm in width. If an extended source is considered (~ 1.45 mm in diameter), the illuminated area of the grating will in this case be approximately ~ 12 mm in height and ~ 4 mm in width.

### 4.4.2 System with additional focusing optic

In the case of the aperture matched system, the GCA is positioned 218 mm from the source due to the necessary in-line valve. The acceptance angle of the GCA
is therefore $-28 \times 10^2$ radians with the same optical transmission of $-30\%$. The flux incident onto the concave mirror will be $-5.8 \times 10^9$ photons nm$^{-1}$. Considering the reflectivity of the concave mirror to be $-20\%$ @ 80 nm the flux focused on the spectrometer's entrance slit plane is $-1.2 \times 10^9$ photons nm$^{-1}$. Assuming the focus is approximately the same size as the source the slit will have an effective transmission of $-9\%$ and the flux incident on the grating will be $-1.1 \times 10^8$ photons nm$^{-1}$. The grating has a surface reflectivity of $-20\%$ @ 80 nm and so the flux focused at the dispersion plane of the instrument with a linear dispersion of $-0.83$ nm mm$^{-1}$ is $-1.8 \times 10^7$ photons mm$^{-1}$ or equivalently,

$-4.4 \times 10^5$ photons per pixel,

assuming a detector with pixels 25 μm in width. If an extended source is considered (1.45 mm in diameter) the illuminated area of both the spherical concave mirror and grating will be approximately $-28$ mm in diameter.

The increase in throughput is therefore a factor of $-11$ assuming the estimates on reflectivity of the concave mirror are correct and the source can be accurately modeled as a point source. This latter assumption has the effect of increasing the difference between the calculated throughput of the two arrangements. In reality, the laser produced plasma will emit VUV light within a volume which depends on the observation wavelength and focus conditions as well as target material. Experiments on a tungsten target with the use of a 525 μm wide spatial resolving slit indicates that the source size at a wavelength of ~50 nm has dimension which are approximately 1.45 mm in diameter as measured at FWHM of intensity.

4.5 Shadow ray trace results

Shadow is a software suite which uses Monte Carlo$^\dagger$ methods to model ray traces in optical systems predominately used in the XUV and soft x-ray wavelength regions. It was initially written for the UNIX operating system but has been ported to other platforms in recent years. Through successive application of the physical laws.

---

$^\dagger$ Monte Carlo Describes a mathematical technique used in modeling of physical systems where the variable or variables—in this case the source emission features—follow random distributions.
of reflection, refraction and attenuation. Shadow traces the path of individual rays using vector algebra with a minimum use of trigonometric functions. Work was done using Shadow in order to investigate the following topics:

- General spectrometer understanding
- Throughput calculations and comparisons
- Misalignment errors
- Spatial resolution properties with the use of slits and apertures

The laser produced plasma was assumed to be a spherical light source with diameter 1.45 mm estimated from space resolved measurements on a tungsten plasma and uniform radiance with respect to position in source volume as well as angle of observation. Other sizes and types of sources (e.g., point, disc shaped etc.) were used in the initial testing of the ray traced system. A conical source divergence was used with a half angle chosen to ensure that as many rays as possible were conserved while still completely filling the effective aperture defined by the GCA. The maximum number of individual rays which can be traced is 25000 although a number of results of runs with different random number seeds can be summed together to provide better statistics. Although Shadow allows for capillary based optical elements to be included, in practice it was found that the code was more suited to longer length capillaries (e.g., Kumohov optics etc.) and gave inconsistent results when trying to model the GCA directly. It was however modeled as an effective circular aperture with a radius of $d/l * r$, where $r$ is the distance to the source ($r = 170$ mm in the case of direct coupled system and 218 mm in the system including the focusing optic) with a maximum radius of 3 mm defined by the iris effect of the GCA holder. The transmission of the GCA is estimated to be ~30% for rays which fall within this effective aperture [Lucatorto et al., 1979]. The spectrometer was modeled in Shadow using a manual translational, $x$, of the grating given by $E4-1(b)$ and allowing Shadow to calculate the necessary rotational angle internally for the chosen center wavelength on the detector plane.

$$\varphi = \sin^{-1} \left( \frac{m\lambda}{2d \cos(D_r)} \right) \quad [E4-1(a)]$$
where $R$ (995 4 mm) is the focal length of the concave grating, $m$ is the spectral order, $\lambda$ is the central wavelength of interest, $d$ (1/1200 * $10^{-3}$ m) is the inter groove spacing of the grating and $D_\nu$ is the deviation angle of the spectrometer. No reflectivity of any optical component was considered explicitly in the ray tracing model. Rather, it was accounted for in throughput calculations by reducing the number of rays reflected from a mirror surface in accordance with a reflectivity. The blaze angle of the grating has no effect on modeling when reflectivity is not taken into account and so was ignored.

Shadow also allows for both the rotational and translational movements of the source and optical elements in the system. The optical element movements are defined by 6 variables, which describe the translations along the x-, y- and z-axis and rotations around the x-, y- and z-axis. Movement of an optical element also moves any succeeding screens, apertures and slits connected to the particular optical element. Therefore, in order to model a planar detector fixed in space, while allowing the grating to be moved, an optical element consisting of a reflective planar mirror was used which was tangential to the Rowland circle.

Shadow uses a Cartesian coordinate system and refers all of the ray components to this reference frame. The resulting output data files are in binary format and include columns containing x, y and z axis ray coordinates as well as $x'$, $y'$ and $z'$ coordinates in phase space. The coordinates defining rays on a particular mirror, lens or grating optical element may be plotted by use of the data in x and y columns. The coordinates for rays intercepting a particular continuation plane, slit, aperture, screen or source may be plotted by use of the data in the x and z columns. The phase space (sometimes referred to as momentum space) coordinates contain information on both the vertical and horizontal focusing conditions and also on the astigmatism and other aberrations present in the optical system. An ideal focusing system would result in a straight line in phase space, a deviation from this straight line directly indicates the existence of aberrations within the optical system. Phase space is composed of co-ordinates which describe the angle the ray makes with the
propagation direction, \( z' \) in the case of reflecting or refracting optical components and \( y' \) for all others [Shadow User's Guide, 1994]

![Diagram](image)

*Figure 4-3*  Shadow ray trace results with direct coupled system. Source size 1.45 mm spherical diameter, conical half angle divergence 0.004 radians, multilines 79.5, 79.25, 80, 80.5 and 80.75 nm and uniform angular radiance. Number of initial rays 25000 (upper left) Beam footprint on grating and (upper right) image at dispersion focal plane and histogram with 25 \( \mu \)m bins (lower left) vertical axis phase space and (lower right) horizontal axis phase space.

It can be seen from figure 4-3 (upper left) that the illuminated width of the grating with the direct coupled system using a slit of width 50 \( \mu \)m and height 3 mm is approximately 4 mm \( \times \) 10 mm and is the result of the small source size and position of GCA and spectrometer slit. This illuminated width did not change significantly with wavelength although the central position of the illuminated footprint moved horizontally on the grating. The latter result is the inherent consequence of the McPherson mount. The optic axis defined by the centre of the grating to the entrance slit is not fixed but varies slightly with wavelength. This
displacement of the optic axis can be calculated by simple geometry for the McPherson type mount and is given by the following equation.

\[
\theta = \sin^{-1}\left(\frac{x \sin(\theta)}{x^2 + (R \cos(\theta))^2 - 2xR \cos(\theta)^2}\right)
\]

[ E 4-2 ]

where \( R \) is the radius of curvature, 995.4 mm, and \( x \) is the displacement as given by equation 4-1 (b)
Figure 4-4  Graph showing the displacement of the optic axis from position at zero order as given by equation 4-2

The height of the spectral lines are dependent on the wavelength for which the grating is tuned and is ~ 2.6 mm @ 80 nm. The stigmatic property of the instrument is displayed in figure 4-3 (lower left) where the straight line centred at the origin indicates a linear relationship between the vertical position of a particular ray on the detector plane and its distance from the vertical focus. For a truly stigmatic instrument the slope of this data represents the reciprocal of the distance to the vertical focus. The optimum horizontal focus (i.e., that focus that gives sharpest spectral lines) is very close to the grating position as given by equation 4-1 (a) and (b). This is evident from the fact that the data points in figure 4-3 (lower right) are symmetric around the x-axis in horizontal phase space at the position of the detector plane. On the lower wavelength side of the chosen centre wavelength region the spectral lines are observed to be focused (horizontally) closer to the grating while on the higher wavelength side the spectral lines are focused (horizontally) further from the grating. It was found that the maximum deviation, as predicted by Shadow, from optimum horizontal focus across the entire detector plane (of width ~ 25 mm) was approximately ± 10 μm. In practice this small off-focus distance for wavelengths
at the edge of the detector does not interfere with the resolution capability of the Acton™ spectrometer when using the CEMA / PDA detector arrangement.

Figure 4-5 (solid line) Graph of spectral line height versus wavelength for direct-coupled system as predicted by ray trace results with 1.45 mm spherical monochromatic source, conical half angle divergence of 0.004 radians and uniform angular radiance. The maximum extent of the ray co-ordinates at the detector plane were used to quantify the height of the spectral line and (circular data point markers) experimental spectral line height for identified lines.

In figure 4-5 it is seen that the spectral line height as predicted by the ray tracing varies with regard to the wavelength of the particular line. Although the curvature of such lines is predicted by theoretical considerations, at near normal incidence and with the use of relatively large slit heights, the curvature is not apparent in ray traced or experimental results. The experimental spectral line height was recorded by the use of the direct illuminated back-thinned CCD array detector for the spectral range 30 to 160 nm with the upper wavelength limit set by lack of signal. The zero order spectral line is approximate 6 mm in height, measured as...
FWHM and being composed of predominantly visible light which is emitted from a larger source size deviates substantially from the trend predicted by the ray traced results. Identified spectral line heights versus wavelength were plotted in both first and second and third order. Apart from were the lines are very intense at ~ 50 nm the results show a gradual decrease in line height with increasing wavelength which follows that as predicted by the ray trace results indicating that a diameter of ~ 1.5 mm is a good estimate for the average source size in the VUV in the present experimental arrangement.

Once the direct coupled spectrometer arrangement was ray traced in Shadow and the focusing condition was confirmed (translational and rotational movements given by E 4-1 (a) and (b)) for the entire spectral range of the instrument the additional mirror was included and the GCA was repositioned 218 mm from the source. The spectrometer part of the model was unchanged from that used in the direct coupled system with the exception of the mirror orientation angle of the grating (changed from 0° to 180°) and the planar mirror acting as a detector (changed from 180° to 0°). A schematic of the modeled system showing axis conventions and labels of individual optic components as used by Shadow is displayed in figure 4-6.
Figure 4-6 Schematic of optical system showing co-ordinate system as used by Shadow in the modelling of the system with additional focusing element. The system is composed of three optical elements (mirr 01, mirr 02 and mirr 03 as labelled by shadow), an entrance slit attached to grating optical element and a circular aperture (serving to model the GCA). Other apertures were also included in order to observe the extent of the collected light at various different distances from the source.
Figure 4.7 Shadow ray trace results with the additional focusing optic. Source size 1.45 mm spherical diameter, conical half angle divergence 0.02 radians, multi-line 79.5, 79.25, 80, 80.5 and 80.75 nm and uniform angular radiance. Number of initial rays 25000. Working from left to right, top to bottom: (a) Footprint on spherical concave mirror, (b) footprint on slit showing only accepted rays, (d) vertical phase space at slit plane, (e) horizontal phase space at slit plane, (f) footprint on grating, (g) footprint on detector plane, (h) vertical phase space at detector plane and (i) horizontal phase space at detector plane.
From figure 4-7 (a) it can be seen that the illuminated width of the concave mirror is approximately 28 mm in diameter. The portion of the slit which is passing light is approximately 50 \( \mu \text{m} \times 19 \text{ mm} \). The width of the grating illuminated is approximately 27 mm in the vertical direction and 25 mm in the horizontal direction.

![Graph of spectral line height versus wavelength for system with additional focusing optic as predicted by Shadow](image)

**Figure 4-8** Graph of spectral line height versus wavelength for system with additional focusing optic as predicted by Shadow. The maximum extent of the ray coordinates at the detector plane were used to quantify the height of the spectral line.

From figure 4-8 it can be seen that the variation of spectral line height with wavelength displays a different shape than the direct-coupled system. In the direct-coupled system, the complete entrance slit of the spectrometer was filled. The instrument effectively images this slit onto the dispersion plane with unit magnification and the result is a more gradually varying function with respect to wavelength (figure 4-5). In the case of the system with the additional focusing element, the instrument effectively images a point source which achieves better filling of the grating and therefore the astigmatism and curvature due to coma of the instrument is more pronounced.

The throughput improvement can be estimated by considering the flux at each optical element and taking account of the surface reflectivity of both the mirror and
grating. The surface reflectivity of the concave mirror was estimated at ~ 20% for an iridium coating (from reflectivity data from Acton™). The GCA position is 170 mm from the source in the case of the direct-coupled system and 218 mm from the source in the system including the focusing optic.

<table>
<thead>
<tr>
<th></th>
<th>Direct coupled system.</th>
<th>Addition of the focusing optic.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial number of rays.</td>
<td>25000.</td>
<td>25000.</td>
</tr>
<tr>
<td>Rays which pass GCA.</td>
<td>17276.3.</td>
<td>11771.</td>
</tr>
<tr>
<td>Rays which reflected from concave mirror.</td>
<td>2354.2.</td>
<td></td>
</tr>
<tr>
<td>Rays which pass slit.</td>
<td>20.4.</td>
<td>110.8.</td>
</tr>
</tbody>
</table>

Table 4-9. Summary of Shadow throughput results. The source parameters remained the same for each system and a number of data-sets (10 in both cases) were averaged to achieve better statistics.

From a comparison of the ray traced results displayed in table 4-9, the throughput of flux with the system incorporating the spherical concave mirror is approximately 5 times that of the direct coupled system. It must be pointed out that this figure represents the maximum one can expect, as the system in practice will contain misalignments and possibly poor focusing. Also the efficiency of the additional mirror and most especially the grating will be a function of wavelength and therefore the increase in flux will also be a function of wavelength.

4.6 Preliminary construction and alignment of system with additional focusing optic.

Before assembling the complete vacuum optical beamline each main section was separately vacuum tested for leaks. When satisfied that no such leaks were present the mirror chamber section was fixed temporarily to the supporting table and both the table and the mirror chamber were leveled and adjusted in height to suit the spectrometer entrance arm flange (see figure 4-1). An alignment laser was positioned in the holder attached to the target chamber and was adjusted to make the beam central to 6 mm bore of the GCA holder (the actual GCA was not included in the system at this stage). It was important to ensure that the alignment laser was
normal to the GCA holder flange. This was achieved by minimizing the distance between back reflections of a half-silvered mirror and the rear window of the target chamber. The target chamber was moved in-order to center it with the optic axis of the mirror chamber and also to locate the source position to 1000 mm distant from the estimated mirror position. It was found that a half-silvered mirror when held against the entrance flange of the mirror chamber would allow further alignment to be made by the use of the back reflection thus ensuring that the alignment laser was normal to the front face of the flange.

Removal of the CEMA flange allowed an alignment laser to be positioned at the exit arm of the spectrometer by passing it back through the grating and entrance slit defining the optic axis of the spectrometer. Further use of the half-silvered mirror allowed the spectrometer alignment laser to be aligned normal to all spectrometer flanges as well as the exit arm flange of the mirror chamber by small adjustments to mirror chamber support table and spectrometer support table. At this stage a plane mirror was positioned at the location of the concave mirror and using the half-silvered mirror each alignment laser was verified to lie on the optic axis.

The plane mirror was replaced with the concave mirror and the focusing condition was verified by attaching a small light bulb to the target holder and using the filament as an object. Any necessary changes to the target chamber to mirror chamber distance and/or mirror chamber to spectrometer distance was followed by a repeat of the above alignment procedure with the plane mirror. As the mirror chamber consisted of long 40 mm diameter tubes with 40 KF flanges for convenience, the pump down time was increased greatly. It was found that after 3 to 4 days the mirror chamber reached an ultimate pressure of $3 \times 10^{-6}$ mBar as measured by a penning gauge positioned on a tee piece close to the spectrometer entrance slit.
Before spectrometer and target to mirror chamber flange opened

<table>
<thead>
<tr>
<th>Chamber Type</th>
<th>Before Pressure</th>
<th>After Pressure</th>
</tr>
</thead>
<tbody>
<tr>
<td>Target chamber</td>
<td>$1.6 \times 10^5$ Torr</td>
<td>$7.9 \times 10^5$ Torr</td>
</tr>
<tr>
<td>Mirror chamber</td>
<td>$3.5 \times 10^6$ Torr</td>
<td>$2.9 \times 10^6$ Torr</td>
</tr>
<tr>
<td>Spectrometer pressure</td>
<td>$1.3 \times 10^6$ Torr</td>
<td>$1.2 \times 10^6$ Torr</td>
</tr>
</tbody>
</table>

Table 4-12  Typical target, mirror and spectrometer chamber pressures before and after target chamber to mirror chamber and mirror chamber to spectrometer chamber in-line valves are opened

In the design and construction of an optical system some consideration must be placed on the possible sources of misalignment and sensitivity to such misalignments especially if the optical system is enclosed in a vacuum chamber where adjustment of the optical elements may be difficult to achieve. Although some form of initial alignment at atmospheric pressure is crucial if close to optimum conditions are to be reached, the process of pumping the system down may lead to movement of one or more of the aligned components. So long as this drift can be compensated for with relative ease, absolute fixture of the optical components may not be of primary importance when considering cost and size of support tables.

The initial testing consisted of positioning a laser plasma source on the optic axis of the spectrometer system and moving the grating to the zero order position. It was observed that the original alignment was somewhat disturbed upon pump down and therefore it was necessary to scan the resulting focused image of the source across the spectrometer’s entrance slit horizontally by rotation of the concave mirror. Once the zero order spectrometer image was observed the concave mirror was scanned in the vertical direction to optimize the detectors counts. Although the zero order signal displayed high intensity it was found to be predominantly of visible light. This was confirmed by use of a glass plate on the second target holder to attenuate the VUV portion ($\lambda < 200$ nm) of the zero order signal. Movement of the glass microscope slide to a position between the source and mirror resulted in little...
change to the zero order signal strength. The zero order signal is displayed in figure 4-13 with and without the glass microscope slide attenuating the source emission. The amount of VUV light contained in the un-attenuated emission is estimated to be 6%.

A mercury spectral lamp was also used and allowed for the testing of the experimental arrangement at the longer wavelength extreme ($\lambda > 270$ nm). The rear window of the target chamber was replaced by a quartz window and a quartz lens was used to couple the visible and UV light to the spectrometer slit. Although the spherical concave mirror should have a high reflectance in this spectral region no mercury emission lines were observed. This was due to the extremely poor efficiency of the grating in this region which is far from the blazed wavelength. Therefore, the spectral region were the grating is efficient ($\lambda < 100$ nm) and that spectral region were the spherical concave mirror is efficient ($\lambda > 160$ nm) do not overlap and so no spectral signal can be observed.

![Figure 4-13](image_url)

**Figure 4-13** Single shot zero order signal recorded by the system arrangement including the mirror chamber and using the emission from a polyethylene plasma with (a) and without (b) attenuation of the VUV emission by a glass microscope slide. Spectrometer slit 20 $\mu$m.
4.7 Concluding remarks on system modeling.

The focusing of collected VUV radiation onto the slit has been shown to increase system throughput even though the additional focusing optical element has relatively low reflectivity of approximately 20%. The increase has been estimated using both simple geometric arguments and also through the use of a ray tracing modeling software. The increase predicted by the simple preliminary calculations (~11) is found to be much higher than that predicted by the ray tracing experiments (~5). This can be attributed to the fact that a point source is an over simplification which does not correspond to the source used in the ray traced model.

In practice the reflectivity of the additional mirror will be a function of wavelength and will also have misalignment errors in position and so the increase might well be lower than the ray trace results suggest. The efficiency of the grating will vary more with wavelength than the spherical concave mirror, due to the fact that the grating is blazed for a single wavelength.

The height of the resulting spectral lines have been shown to follow quite different trends in the case with the source positioned a distance 382 mm away from the spectrometer entrance slit and that with the an additional concave mirror focusing the source object on to the spectrometer. In the former case the height of the spectral lines as predicted by Shadow decreases gradually with increasing wavelength, 2.6 mm @ 30 nm to 2.1 mm @ 325 nm. When the source object is focused onto the spectrometer entrance slit the spectral line height changes dramatically from 2.7 mm @ 30 nm to 4.5 mm @ 325 nm. By using an array detector the resulting intensity reduction at the detector plane can be compensated by binning the pixels in the vertical direction. Experiments using an array detector similar to the Andor™ back thinned CCD yielded spectral line heights for the wavelength range 30 to 160 nm similar to that predicted by the ray trace model. It was not possible to get accurate spectral line height measurements above 160 nm due to level of spectral signal. The similarity in spectral line height between that of the ray trace results and also from experimental measurements indicates that the source size used in the ray trace results (~1.45 mm) is indeed close to the emitting source size in the experiments.
The translational movement of the grating displaces the optic axis of the spectrometer instrument from its position at zero order given by equation 4.2. Although the displacement is relatively small and does not interfere with experiments using the direct-coupled arrangement, the displacement may in fact cause problems when the system is used with an additional focusing optic and precise alignment of the optical components is necessary.

The misalignment error associated with the additional focusing optic with respect to position relative to optic axis (x-y plane in figure 4-4) and displacement of mirror from ideal focusing position (z-axis in figure 4-4) have a dramatic effect on the flux throughput of the system. This is mainly due to the long lever arm effect of the beamline and therefore accurate angle movement of the mirror is critical.

Although an intense zero order signal was recorded it was mainly composed of visible light and so there was insufficient flux throughput in the entire spectral range covered by the instrument. In the short wavelength extreme of the spectral range of the instrument, \( \lambda < 100 \text{ nm} \) the grating is efficient due to operation close to the blazed wavelength. The quantum efficiency of the Galileo™ CEMA / EG&G™ PDA decreases rapidly at wavelengths longer than approximately 180 nm. The reflectivity of the spherical concave mirror available at the time of the experiments is now known to have negligible reflectivity in this region. Therefore no region exist where both the grating and spherical concave mirror is efficient.
Chapter 5. Concluding remarks

A multichannel normal incidence VUV spectrometer system has been described which enables both emission and absorption studies to be made on laser produced plasmas. Two detection configurations were used in the experiments, one a Channel Electron Multiplier Array / Photo Diode Array (CEMA / PDA) combination, the other a direct illuminated back thinned charge coupled device (CCD). Various emission studies were performed on line as well as continuum dominated plasmas in order to characterise the instrument's wavelength range, extent of order overlapping and wavelength resolution. The suitability of the spectrometer system for photoabsorption measurements was accessed by such experiments as absorption by gas as well as by a secondary plasma plume as in the DLP technique. The design, construction and preliminary testing of a normal incidence focusing optic was presented in chapter 4. The improvement in terms of light gathering ability has been illustrated by both simple geometric calculations as well as by ray tracing.

5.1 Conclusions and observations

Although the wavelength range possible with the present grating and mount extends from 30 to 325 nm the poor efficiency of the grating and problems with order sorting limits absorption studies to approximately 35 to 80 nm without the use of filters however emission studies in the range 35 to 325 nm are possible and the instrument is routinely used up to 220 nm for this purpose.

The Galileo™ CEMA intensifier and EG&G™ PDA detector arrangement provide reasonable wavelength resolution. The spectral lines from line dominated emission spectral are typically 6 to 7 pixels (FWHM) with a corresponding linear dispersion of ~ 0.021 nm pixel\(^{-1}\) in first order. Spectra obtained using the Andor™ back thinned CCD had spectral lines typically 3 to 4 pixels in width (FWHM) with a corresponding linear dispersion of ~ 0.022 nm pixel\(^{-1}\) in first order. The binning of the pixels of the CCD array in the vertical direction did not decrease the wavelength resolution achievable.

Generally speaking, the comparison of the detector sensitivity is rather crude because many of the factors involved, such as gain and efficiency of the detection
device, are strong functions of other parameters. Since the gains on both the MCP and of the phosphor screen stages of the CEMA are steep functions of applied voltage, and the CCD dark noise also depends on the operation temperature, an estimate of the sensitivity is more qualitative than quantitative. The CEMA / PDA detector arrangement with a CEMA bias of 2500 / 800 V (V_gap / V_plate) showed a similar sensitivity range as that of the Andor™ back thinned CCD over the entire range of the spectrometer. Both detectors show good single shot sensitivity at wavelengths below about ~140 nm. The back thinned CCD with its inherently low noise and 16 bit ADC has a higher dynamic range than the CEMA / PDA based detector.

Photoabsorption measurements on both gas and secondary plasma plumes are possible in the wavelength range approximately from 30 to 80 nm with the upper limit determined by the falloff in efficiency of grating. Both samarium and tungsten plasma prove to be good backlight sources for reproducible continuum emission. The samarium plasma emission in the normal incidence VUV region is essentially 'line free' with some spectral lines arising from lower ion stages and also from oxygen present as a surface contaminant. The reproducibility of these low ion stage features is excellent and it is the oxygen line emission which is responsible for some of the peaks in the ratio plots.

The average flux of the emission (centred at ~50 nm) from a samarium plasma is reproducible to ~2 % when the Spectron™ SL803 laser is used in contrast to ~16 % with use of the JK Lasers™ Ruby laser. Although the tungsten emission is characterised by numerous lines, the reproducibility of these lines is excellent. Therefore, a tungsten plasma is also ideal for use as a backlight as the collected emission is stable over hundreds of shots without rotation to a fresh surface. The reproducibility of the continuum emission is improved by incorporating a secondary slit of suitable width between the spectrometer slit and the source. This reduces the effect which the variation of source size has on the collected flux. The use of a secondary slit is also important in DLP photoabsorption studies for limiting the region through which the collected continuum backlight passes.

The inclusion of such a secondary slit has numerous advantages. These include, (1) limiting the amount of front plasma emission collected by the
spectrometer, (2) limiting the range of species present in photoabsorption measurements and (3) reducing the amount of continuum signal collected which has not passed through the absorption plume and also reducing the scattered light present in general.

Also it is important to note that saturation in photoabsorption measurements reduces the wavelength resolution achievable as well as increasing the effect the underlying noise has on the relative photoabsorption cross sections. The system parameters which determine the saturation level include (1) detector noise, (2) stray light, (3) higher order light present and (4) strength and reproducibility of backlight emission, $I_0$.

Higher order contributions have been observed to dominate at wavelengths above ~100 nm. By use of a gas such as helium or argon this upper limit may be extended to ~140 nm. Solid state filters are limited by their high absorption of VUV light above about ~100 nm.

In chapter 4 the design, construction and preliminary testing of the spectrometer system with an additional focusing element is described as well as the benefits of such a system. Although both the simple geometric calculations as well as ray trace results suggested that the throughput would be increased with the introduction of the focusing optic, no spectral data could be obtained in experiments to verify this. The main difficulty encountered was the poor reflectivity of the aluminium mirror with MgF$_2$ coating. It was evident that the wavelength range were the grating was efficient (30 to approximately 120 nm) did not overlap that range for which the additional concave mirror (aluminium with MgF$_2$) had an adequate reflectivity. In view of this an iridium coated mirror is presently on order from Acton™.

The focusing of collected VUV radiation onto the spectrometer slit has been shown to increase system throughput even though the additional focusing optical element has relatively low reflectivity of approximately 20%. The increase has been estimated using both simple geometric arguments and also through the use of a ray tracing modelling software. The increase predicted by the simple preliminary calculations (~11) is found to be much higher than that predicted by the ray tracing
experiments (~5). This can be attributed to the fact that a point source is an oversimplification which does not correspond to the source used in the ray traced model.

Spatial resolution can be important for photoabsorption studies as the dominant ion stage can be selected somewhat by imaging a specific portion of the plume onto the entrance slit of a spectrometer. The use of a slit allows for the collection of light from a vertical chord of the plasma plume. Usually slits reduce the amount of light coupled into an optical system reduces considerably and therefore the signal-to-noise ratio of the recorded spectrum suffers. Hence it is important that the system is as close to aperture matched as possible.

In practice, the reflectivity of the additional mirror will be a function of wavelength and will also have misalignment errors in position and so the increase might well be lower than the ray trace results suggest. The efficiency of the grating will vary more with wavelength than the spherical concave mirror due to the fact that the grating is blazed for a single wavelength.

Although an intense zero order signal was recorded it was mainly composed of visible light and so there was insufficient flux throughput in the entire spectral range covered by the instrument. In the short wavelength extreme of the spectral range of the instrument, \( \lambda < 100 \text{ nm} \), the grating is efficient due to operation close to the blazed wavelength. The quantum efficiency of the Galileo™ CEMA / EG&G™ PDA decreases rapidly at wavelengths longer than approximately 180 nm. The reflectivity of the spherical concave mirror available at the time of the experiments is now known to have negligible reflectivity in this region. Therefore no region exist where both the grating and spherical concave mirror is efficient.
5.2 Future developments

Some of the planned applications of this normal incidence VUV spectrometer instrument are briefly described below.

5.2.1 Laser driven ionization of a laser generated barium plasma

When a DYE laser with a pulse energy of greater than ~ 200 mJ and pulse duration of typically 500 ns is tuned to a resonance or intercombination transition of an atomic species, it will produce almost complete ionization of a metal vapour within typically a microsecond of the initial irradiation [Lucatorto and McIlrath, 1980 and Kent et al., 1987]. The ability to form a controlled ionized sample of a metal vapour with all of the ions residing in the ground state makes RLDI ideally suited to absorption studies.

The RLDI technique was first performed by both Lucatorto and McIlrath, [1976] to study the absorption spectra of various metal vapours including lithium, sodium, manganese and barium. These results permitted temporal measurements on the state of ionization of the sample and also provided data on the mechanisms involved in the ionization.

Some of the necessary pumping transitions are shown below all of which are readily obtainable with a DYE laser [Kallenbach et al., 1986].

\[
\begin{align*}
\text{Ba I} & \quad 6s^2 \text{S}_0 \rightarrow 6s6p \text{P}_1 \quad \lambda = 553.5 \text{ nm} \\
\text{Ba II} & \quad 6s^2 \text{S}_{1/2} \rightarrow 6p^2 \text{P}_{1/2} \quad \lambda = 493.5 \text{ nm}
\end{align*}
\]

5.2.2 Gating of channel electron multiplier array

XUV and VUV imaging of laser produced plasmas have often used channel electron multiplier array (CEMA) devices because of their high spatial resolution (typically 40 lp mm\(^{-1}\) for 10 μm pore diameters) and high gain (typically \(10^5\) for a single stage device) [Galileo™, 1986]. Normally a steady state dc voltage is used to bias the intensifier thus producing time-integrated images of the plasma. In photoabsorption studies, it is possible to use a second plasma as a continuum source.
This allows for the 'natural' gating of the photoabsorption of the front sample plasma with a temporal resolution of the order of the duration of the continuum back-light. In the XUV region, the continuum emission duration follows that of the excitation laser pulse duration. At longer wavelengths, the continuum emission is significantly longer. In the case of temporally resolved emission studies some form of intensifier or detector gating is required.

Time resolution in the nanosecond or sub nanosecond range may be achieved using uni-dimensional streak cameras. Quite often the plasma displays axial symmetry and so the spatial resolved image of a single dimension is acceptable. A detector which exhibits bi-dimensional spatial resolution and very short time resolution would be advantageous in viewing the complete plasma in both spatial dimensions [Luppi et al., 1984].

One way such a bi-dimensional detector could be realised would be to gate the bias voltage of a CEMA based intensifier. Producing a transient high potential pulse with sharp transition times is difficult because component and stray capacitance must be kept to a minimum and also devices must be used that can both handle the high voltage and high peak power of the pulse. Circuits for such gating usually consist of power MOSFETS or avalanche bipolar transistors [Fulkerson and Booth, 1994].

On avalanching, a bipolar transistor can be made to produce extremely fast, high peak current pulses. Most transistors are capable of producing an avalanche response but only a few are able to produce a reproducible and sustained operation. Recently a number of transistors which are characterised specifically for avalanche pulse operation have become available, namely the ZTX415 from Zetex Semiconductors™. The avalanche transistor depends for its operation on the negative resistance characteristic at the collector. When the collector voltage exceeds a certain level, somewhere between $V_{CEO}$ and $V_{CEO}$, depending on the circuit configuration, the voltage gradient in the collector region exceeds the sustained field strength, and electron-hole pairs are liberated. These are accelerated by the field, liberating others in their turn. Consequently, the current increases rapidly, although the voltage across the device is actually falling [Zetex Semiconductors™, 1996 and Hickman, 1996].
5.2.3 Application to FEL characterisation.

It has been decided that the Acton™ VM 521 normal incidence VUV spectrometer will be used in the characterisation of the free electron laser (FEL) experiment located at DESY synchrotron facility Hamburg in Germany. In a conventional laser, the emission of photons results from the transitions of atom-bound electrons from a higher to lower quantum state. In the free electron laser, the electrons that produce the photons are not bound to any atom but are free; they are generated in an electron accelerator and emit coherent radiation as they undergo periodic oscillations within an insertion device such as an undulator. Although the emission from insertion devices presently in use is coherent it is not monochromatic and the radiation and therefore the power is distributed across a broad wavelength range (IR to VUV).

By increasing the length of the undulator it becomes possible to amplify a desired wavelength which can be tuned by varying the period between the magnets in the undulator or varying the energy of the electrons entering the undulator. The operation of the proposed FEL will be based on self amplification of spontaneous emission (SASE) and therefore will not require a seed laser to initiate lasing action. As the proposed design uses a single pass of the undulator to provide adequate amplification it removes the necessity of obtaining efficient mirrors suitable for operation at short wavelengths.

The grating in the Acton™ spectrometer will be replaced with one which has a higher groove density (either 2400 or 3600 lines mm\(^{-1}\)) thereby increasing the wavelength resolution of the instrument. It is proposed to incorporate a wedge behind the new grating to extend this coverage to 130 nm. Another possibility would be to get Acton™ to modify the grating mount to achieve the necessary coverage. A Photonics™ back thinned direct illuminated CCD detector similar to the Andor™ model described in previous chapters will be used to record the diffracted spectrum. Initial testing and alignment has shown that with the 1200 lines mm\(^{-1}\) grating a FWHM line width of just 2 pixels (approximately 0.2 nm pixel\(^{-1}\)) is readily possible with this detector. Calibration of the spectrometer instrument will be made through the use of a calibrated ECR source. This will provide both wavelength as well as
absolute efficiency across the entire range of the instrument. The output from the FEL will be coupled to the spectrometer with one of two arrangements, either a plane mirror or a spherical concave iridium mirror.
# References

<table>
<thead>
<tr>
<th>Reference</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
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</tr>
<tr>
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<td>Celliers, P, Mrowka, S, Matthews, D 1995 Optimization of X-ray sources from a high-average power Hd Glass laser produced plasma for proximity lithography <em>UCRL-LR-105821-95-3</em></td>
</tr>
</tbody>
</table>


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**Appendix I  Acton™ VM-521 1 metre normal incidence spectrometer specifications**

<table>
<thead>
<tr>
<th>Specification</th>
<th>Details</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Model</strong></td>
<td>VM - 521</td>
</tr>
<tr>
<td><strong>Focal length</strong></td>
<td>0.9954 m</td>
</tr>
<tr>
<td><strong>Optical system</strong></td>
<td>Near normal incidence 7.5° / 7.5°</td>
</tr>
<tr>
<td><strong>Horizontal aperture ratio</strong></td>
<td>F / 10.4 with 96 mm wide grating</td>
</tr>
<tr>
<td><strong>Grating</strong></td>
<td></td>
</tr>
<tr>
<td>Manufacturer</td>
<td>Bausch &amp; Lomb</td>
</tr>
<tr>
<td>Type</td>
<td>Concave reflective</td>
</tr>
<tr>
<td>Grooves mm⁻¹</td>
<td>1200</td>
</tr>
<tr>
<td>Concave radius</td>
<td>9.954 mm</td>
</tr>
<tr>
<td>Coating</td>
<td>Indium</td>
</tr>
<tr>
<td>Blaze angle</td>
<td>2.75°</td>
</tr>
<tr>
<td>Blaze wavelength</td>
<td>80 nm (First order)</td>
</tr>
<tr>
<td>Ruled width</td>
<td>96 mm</td>
</tr>
<tr>
<td>Ruled height</td>
<td>56 mm</td>
</tr>
<tr>
<td>Wavelength range</td>
<td>30 - 325 nm (Max mechanical movement)</td>
</tr>
<tr>
<td>Resolution capability</td>
<td>0.014 nm (Entrance and exit slit width 10 µm)</td>
</tr>
<tr>
<td>Reciprocal linear dispersion</td>
<td>0.83 nm mm⁻¹</td>
</tr>
<tr>
<td>Nominal wavelength coverage</td>
<td>~ 21 nm for both CEMA / PDA and back thinned CCD based detectors</td>
</tr>
<tr>
<td>Wavelength accuracy</td>
<td>± 0.05 nm (mechanical mount movement)</td>
</tr>
<tr>
<td>Wavelength reproducibility</td>
<td>± 0.005 nm (mechanical mount movement)</td>
</tr>
<tr>
<td>Scanning mechanism</td>
<td>Sine drive mechanism with precision lead screw external to the vacuum system</td>
</tr>
<tr>
<td>Focusing</td>
<td>Cam controlled translation motion with Micrometer adjustments external to vacuum system for both short and long wavelength focus adjustments Long wavelength adjustment micrometer closest to spectral drive Micrometer settings for Galileo™ CEMA and Andor™ back thinned CCD detector using present flanges Short wavelength 2.21 Long wavelength 1.85</td>
</tr>
</tbody>
</table>
Figure AI-1  Schematic of Acton™ VM-521 normal incidence spectrometer showing approximate external dimensions in inches
### Appendix II. EG&G™ photo diode array (PDA) detector specifications

<table>
<thead>
<tr>
<th>Specification</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>PDA model</td>
<td>1453A</td>
</tr>
<tr>
<td>Interface model</td>
<td>1471A</td>
</tr>
<tr>
<td>Detector type</td>
<td>Silicon photodiode array</td>
</tr>
<tr>
<td></td>
<td>Silicon behind fused silica face plate</td>
</tr>
<tr>
<td>Wavelength range</td>
<td>180 nm to 1100 nm</td>
</tr>
<tr>
<td>Quantum efficiency</td>
<td>70 % @ 650 nm</td>
</tr>
<tr>
<td>Gain</td>
<td>1350 photo electrons / count</td>
</tr>
<tr>
<td>Sensitivity</td>
<td>1960 photons / count @ 550 nm</td>
</tr>
<tr>
<td>Maximum deviation from linear response</td>
<td>1 %</td>
</tr>
<tr>
<td>Dynamic range</td>
<td>32768 @ 15-bit</td>
</tr>
<tr>
<td>Uniformity of sensitivity</td>
<td>± 5 %</td>
</tr>
<tr>
<td>Maximum geometric distortion</td>
<td>1 channel</td>
</tr>
<tr>
<td>Line width</td>
<td>2 channels FWHM</td>
</tr>
<tr>
<td>Array length</td>
<td>25 mm</td>
</tr>
<tr>
<td>Pixel number and array size</td>
<td>1024 pixels, 25.6 mm * 2.5 mm</td>
</tr>
<tr>
<td>Centre to centre spacing</td>
<td>25 μm</td>
</tr>
<tr>
<td>Dark current in counts</td>
<td>1024 (800 ms @ 5 °C)</td>
</tr>
<tr>
<td></td>
<td>120 (1 s @ -20 °C)</td>
</tr>
<tr>
<td>Maximum system noise</td>
<td>1.5 (counts / scan)</td>
</tr>
<tr>
<td>Temperature range</td>
<td>-40 °C to 18 °C</td>
</tr>
<tr>
<td>Gas requirements</td>
<td>Must be flushed with dry nitrogen for temperatures lower than room temperature during operation and after operation until detector system has reached room temperature</td>
</tr>
</tbody>
</table>

† Specifications taken from "Models 1452, 1453, and 1454 silicon photodiode detectors" Operating and service manual EG & G Princeton Applied Research
Appendix III Andor Technology™ back thinned CCD specifications.

CCD head model: DH 425 - V
Serial number: CCD - 359
Array number: 1024 * 256
Pixel size: 26 µm²
Thermal electric cooling: 3 - stage
Controller card: CC100 (1 MHz)
A/D resolution: 16 - bit
Readout speed: 1, 2, 16, 32 us per pixel

Readout noise

<table>
<thead>
<tr>
<th>A/D rate</th>
<th>R/S</th>
<th>CCD gain</th>
<th>Single pixel</th>
<th>Full vertical binning</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 MHz</td>
<td>1 µs</td>
<td>10</td>
<td>40</td>
<td>40 6</td>
</tr>
<tr>
<td>500 kHz</td>
<td>2 µs</td>
<td>10</td>
<td>38</td>
<td>37 9</td>
</tr>
<tr>
<td>62 kHz</td>
<td>16 µs</td>
<td>10</td>
<td>8 4</td>
<td>9 1</td>
</tr>
<tr>
<td>31 kHz</td>
<td>32 µs</td>
<td>5</td>
<td>6 4</td>
<td>7 7</td>
</tr>
</tbody>
</table>

Base mean level

<table>
<thead>
<tr>
<th>A/D rate</th>
<th>R/S</th>
<th>Counts</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 MHz</td>
<td>1 µs</td>
<td>555</td>
</tr>
<tr>
<td>500 kHz</td>
<td>2 µs</td>
<td>460</td>
</tr>
<tr>
<td>62 kHz</td>
<td>16 µs</td>
<td>210</td>
</tr>
<tr>
<td>31 kHz</td>
<td>32 µs</td>
<td>234</td>
</tr>
</tbody>
</table>

CCD dark current

Minimum dark current achievable: 0.37 @ -54.3 °C with 12 °C water cooling
CCD dark current uniformity: Better than 0.07 electrons / pixel / sec
Linearity: Better than 1 % over 16 bits
Response uniformity: Better than 3.3 %
Dark current defects: No hot spots

Response defects (Dark pixels)
<table>
<thead>
<tr>
<th>Power supply unit</th>
<th>PS 150</th>
</tr>
</thead>
<tbody>
<tr>
<td>Multi I/O box</td>
<td>IO 159</td>
</tr>
<tr>
<td>CCD chip</td>
<td>EEV CCD30 - 11</td>
</tr>
<tr>
<td>CCD chip serial number</td>
<td>A4593 - 13 - 11</td>
</tr>
<tr>
<td>Format</td>
<td>Back illuminated</td>
</tr>
<tr>
<td>Coating</td>
<td>None</td>
</tr>
</tbody>
</table>

**Figure A III-1**  (a) Andor Technology™ back thinned CCD and (b) drawing showing main dimensions of front flange
### Appendix IV Galileo™ Channel Electron Multiplier Array (CEMA) specifications

<table>
<thead>
<tr>
<th>Specification</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>MCP model</td>
<td>Galileo 6025 - FM MCP</td>
</tr>
<tr>
<td>Plate outside diameter</td>
<td>32.74 mm</td>
</tr>
<tr>
<td>Plate thickness</td>
<td>0.43 mm</td>
</tr>
<tr>
<td>Active area</td>
<td>4.9 cm²</td>
</tr>
<tr>
<td>Electrode material</td>
<td>Inconal or nichrome</td>
</tr>
<tr>
<td>Pore size</td>
<td>10 μm</td>
</tr>
<tr>
<td>Centre to centre spacing</td>
<td>12 μm</td>
</tr>
<tr>
<td>Open area ratio</td>
<td>55%</td>
</tr>
<tr>
<td>Bias angle</td>
<td>12°</td>
</tr>
<tr>
<td>Solid glass border</td>
<td>Yes</td>
</tr>
<tr>
<td>Minimum electron gain</td>
<td>$3.3 \times 10^4$</td>
</tr>
<tr>
<td>MCP / phosphor gap distance</td>
<td>1.2 mm</td>
</tr>
<tr>
<td>Bias current</td>
<td>$4 \times 10^{16} \mu$Amps per plate</td>
</tr>
<tr>
<td>Maximum linear output density</td>
<td>10% of bias current density</td>
</tr>
<tr>
<td>Fibre optic phosphor</td>
<td>P20</td>
</tr>
<tr>
<td>Coating</td>
<td>MgF₂ Magnesium fluoride</td>
</tr>
<tr>
<td>Maximum bakeable temperature</td>
<td>350 °C</td>
</tr>
<tr>
<td>VUV wavelength range</td>
<td>(detection efficiency &gt; 10%) 20 - 180 nm</td>
</tr>
</tbody>
</table>

![Diagram of Galileo™ CEMA and vacuum assembly](image)

**Figure A IV-1 (a) Galileo™ CEMA and (b) vacuum assembly showing position of feedthroughs with vacuum side showing**
## Appendix V. Galileo™ glass capillary array (GCA) specifications.

<table>
<thead>
<tr>
<th>Individual pore size</th>
<th>50 μm diameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>Percentage open area</td>
<td>60 %</td>
</tr>
<tr>
<td>Thickness</td>
<td>3 mm</td>
</tr>
<tr>
<td>Diameter</td>
<td>25 mm</td>
</tr>
<tr>
<td>Glass material</td>
<td>Soda lime glass</td>
</tr>
</tbody>
</table>
Appendix VI. Photographs of system.

Figure AIX-1  Photograph showing system configuration used in helium gas photoabsorption and barium photoabsorption
Figure AIX-3
CEMA / PDA detector arrangement

Figure AIX-4
(a) Mirror chamber showing 40KF flanges at 15° and (b) mirror holder and positioner for both focusing and angle alignment