Building a Laboratory based XUV Microscope

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High harmonic generation (HHG) is a non-linear interaction between a driving laser pulse and a target gas. The result of this process is the generation of coherent extreme ultraviolet (XUV) and soft X-ray radiation at harmonic frequencies of the driving laser. There are several uses for this source including attosecond time resolved spectroscopy and nanoscale imaging which this thesis will focus on.

The work contained within this thesis begins by investigating the development of capillary based HHG by comparison of the experimentally observed propagation of the driving laser pulse to both a linear ionisation based model and a multimode non-linear Schrödinger equation model. Manufacture, mounting and coupling into the capillary are described. A simple linear model of propagation along the capillary is compared to experimental measurements of fluorescence along the capillary showing the presence of mode beating. The model is extended to a non-linear propagation model and validated against the spatio-pectral output of the capillary.

The second half of the thesis takes the XUV output from a gas cell and uses it for a coherent diffractive imaging experiment (CDI). The development of the imaging setup is described before presenting the results of a preliminary CDI experiment with a binary test sample and a polychromatic beam. Accurate measurement of the XUV beam focal position was determined by recording diffraction through a binary array of apertures. Results of a CDI experiment demonstrating a resolution of better than 200 nm is shown for a polychromatic beam with an algorithm designed for a monochromatic source.
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DECLARATION OF AUTHORSHIP

I, Richard Thomas Chapman, declare that the thesis entitled ”Building a Laboratory based XUV Microscope” and the work presented in it are my own. I confirm that:

• this work was done wholly or mainly while in candidature for a research degree at this University;

• where any part of this thesis has previously been submitted for a degree or any other qualification at this University or any other institution, this has been clearly stated;

• where I have consulted the published work of others, this is always clearly attributed;

• where I have quoted from the work of others, the source is always given. With the exception of such quotations, this thesis is entirely my own work;

• I have acknowledged all main sources of help;

• where the thesis is based on work done by myself jointly with others, I have made clear exactly what was done by others and what I have contributed myself;

• parts of this work have been published as:


Signed: 

Date: 
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Chapter 1

Introduction

As science has progressed and continues to progress it strives to build upon, challenge and understand what has gone before. This has been achieved by both passive observation of the world around us and observations of our interactions with the world. The discovery of X-rays by Willhelm Röntgen in 1896 [1] heralded a new understanding of the world. The initial discovery showing the skeletal structure of his wife’s hand proved that this new radiation had different penetrative properties to that of visible light and is similar to the technique still used in hospitals today. Use of X-rays for diffraction followed from the discovery of Bragg’s Law [2] of reflections from a crystal, shown by

\[ n\lambda = 2dsin\theta \]  \hspace{1cm} (1.1)

This law describes the relationship between the diffraction order \((n)\) the wavelength of the light \((\lambda)\), the spacing of the crystal planes \((d)\) and the angle of the diffraction beam \((\theta)\). It was from this discovery that the field of X-ray crystallography was started, leading to the routine structural determination of crystalline small molecules and more complex determination of protein molecules [3][4][5]. In the case of protein structures the exact conformation is often lost during the process of crystallisation, leading to the requirement for simulation of how a particular structure may conform when not constrained by the crystallisation process.

A method for experimental determination of a protein shape to aid the structural determination could be used as a complementary tool for a clearer understanding of the
structure. Ideally this would be done with the protein in its natural environment, such as suspended in a small amount of water.

1.1 Imaging

Imaging a protein would require a resolution of approximately one nanometer. Optically the resolution of a microscope is limited by the Abbe limit [6],

$$R = \frac{\lambda}{2NA}$$

(1.2)

where \(R\) is the resolution and \(NA\) is the numerical aperture of the lens used. For visible microscopy this limits resolution to approximately 200 nm. In order to progress past this resolution the wavelength of the illumination source has to be reduced towards the extreme ultraviolet (XUV) and X-ray regions of the electromagnetic spectrum.

The fabrication of imaging optics in the XUV and X-ray regime is both difficult and expensive. In 1972 Gerchberg and Saxton [7] proposed a technique for lensless imaging with electron beams. This work was extended by Fienup [8] [9] to optical microscopy. The process involves the use of an iterative phase retrieval algorithm instead of an imaging lens to re-image the diffraction pattern from an aperiodic sample. Using this technique the resolution limit is given by the numerical aperture of the detector, which can be altered for different experimental conditions. Experimental use of this technique has grown in the past few years [10][11][12] with the improvements in computer technology for performing the reconstructions. This technique is discussed in greater detail in chapter 2.8.

Electron microscopy is often used to image nanoscale objects but the strong interaction of the electrons with the atomic structure of the sample means that transmission measurements can only be made on very thin samples [13]. In contrast a transmission through several microns of material is possible with a tuned XUV or X-ray source.
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Figure 1.1: The pulse length and photon energy/wavelength ranges covered by a variety of ultra-fast X-ray sources. In general, higher photon energies are available from the accelerator based sources, such as the free electron lasers (FELs), while the shortest pulse lengths may be obtained by high harmonic generation (HHG). Currently, high harmonic generation is the only technique capable of producing X-ray pulses that approach the single-optical-cycle limit, represented by the dashed line. Adapted from reference [4].

1.2 XUV and X-ray sources

There are several methods used for generating XUV and X-ray radiation. These come as both large user facilities such as synchrotrons[14] and free electron lasers (FEL)[15] as well as laboratory based sources such as laser plasma generation[16], and high harmonic generation sources[17]. The achievable photon energy and pulse length of these sources was described by Pfeifer et. al[18] and is shown in figure 1.1. The figure shows that there is a trade off between the photon energies available and the pulse durations of these sources. Synchrotrons have a large energy range but at the cost of having long pulses. FEL sources are capable of reaching shorter time resolutions but do not have the energy range. Laser plasma sources have low coherence making them unsuitable for the phase retrieval experiment described above. This leaves high harmonic generation (HHG) as the only laboratory source suitable for these experiments as it has both the energy and time resolution.

The radiation produced by a synchrotron such as Diamond at Rutherford Appleton laboratories, is the result of an electron being radially accelerated. The energy of the electron and the bend radius determine the energy of the radiation. There are three types
of magnetic field used to bend the electrons around the ring, these are bending magnets, wigglers and undulators. As the electron passes through the magnetic field radiation is produced tangentially to the storage ring. Bending magnets and wigglers are used to produce broadband radiation whilst undulators produce a narrower band source. At Diamond the energy ranges from 1.2 eV to 70 keV (1033 nm to 0.018 nm). The coherence of synchrotron sources is low, for example the coherent imaging beam line at diamond is built 250 m from the main synchrotron to improve the spatial coherence[14].

FEL sources use a linear electron accelerator and self amplified stimulated emission to generate radiation. An electron bunch is accelerated along the length of the FEL to energies in excess of 1 GeV. The electron bunch is then passed through a long undulator. In the case of FLASH at DESY the undulator is made of six sections each 4.5 m in length. The magnetic fields within the undulator can be tuned to alter the wavelength of the output, currently the shortest wavelength result is 4.12 nm coherent radiation[19].

HHG sources in contrast are laboratory based sources relying upon frequency conversion of femtosecond laser systems to reach higher order harmonics. Harmonics were first observed shortly after the invention of the laser in 1960 [20] [21]. They are the integer multiple of a fundamental frequency and are commonly thought of in terms of sound and light but can be applied to any periodic system. The frequency of a harmonic is given by:

\[ \omega_q = \omega_0 q \]  

(1.3)

Where \( \omega_q \) is the harmonic angular frequency, \( \omega_0 \) is the fundamental angular frequency and \( q \) is the harmonic number.

Harmonic generation from a laser is commonplace within science and is routinely used in second harmonic generation (SHG) to probe surface interactions in conjunction with sum frequency generation (SFG) experiments. These effects are a perturbative response to the laser field and result in the perturbative region of the HHG spectrum shown in figure 1.3.

As the development of lasers gave higher intensities and shorter pulse lengths the process of HHG was discovered [17][22]. It is the result of the laser interaction with a low pressure gas. Typically the target gas used is a noble gas [23] [24] although some studies of molecular gas targets [25][26] have also been performed. HHG experiments
Figure 1.4: Schematic of the three geometries commonly used in HHG. A gas jet (a), a gas cell (b) or a capillary waveguide (c).

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Figure 1.2: A schematic of the three generation geometries used for HHG, (a) a gas jet, (b) a gas cell and (c) a hollow core capillary

are normally performed in one of three geometries as shown in figure 1.2 [27][28][29]. The gas jet and gas cell have interaction regions that are defined by the focusing conditions, hence limiting volume of the target gas for generation. In a hollow core capillary that interaction region can be extended by wave guiding.

In HHG a strong driving laser field ( > 10^{14} \text{Wcm}^{-2}) is used to drive a valance electron out of the potential well of a target atom causing a highly non-linear response. The result of this interaction are higher order harmonics spanning from the 9th harmonic through to the 5000th (89nm to 0.3 nm)[30][31]. An idealised XUV spectrum has three characteristic regions as shown in figure 1.3. The first part of the spectrum, as described above, is the pertubative region, this then gives way to the plateau region. During the plateau region there is an equal chance of the generated radiation being at one of the energies within this region. Finally there is a sharp cut-off region where the intensity
in each harmonic drops off rapidly to a maximum cut-off energy [32] which is limited by equation 2.5. Experimental spectra like this are not seen due to the absorption and phase matching being different for each wavelength generated.

The XUV generated carries the properties of the driving laser field resulting in both spatial and temporal coherence [33]. The process has an energy conversion efficiency of approximately $10^{-6}$ meaning that the brightness is low in comparison to FELs. The total flux of XUV is spread over several harmonics.

Both synchrotrons and FELs are bright high energy sources but when imaging a protein the time resolution must also be considered. In a study by Neutze et. al [34] it was shown that the protein will undergo a Coulomb explosion after a short exposure to a high energy pulse. The high levels of energy present remove a large number of the valance electrons leaving a highly positively charged molecule. This molecule then explodes due electrostatic repulsion. Figure 1.4 shows that to be able to image the molecule the radiation must be incident on the molecule for less than 10 fs. Even if this is the case then the molecule will still be destroyed after the pulse has passed.

The relatively long pulses from synchrotron radiation will therefore image the molecule predominantly after the explosion. From figure 1.1 it is possible to see that only radiation from FELs and HHG will have both the ultrafast time resolution and the photon energy. The cost of building and running a FEL is hundreds of millions of pounds, as a consequence there are few FELs available and access to beam time is limited.
1.3 Aims of the thesis

Having shown the need for a short wavelength imaging source to obtain high resolution and a femtosecond source to prevent sample damage during imaging, this project sets out to develop a HHG source based imaging system. Based on this the aims of this thesis are:

- Analysis of driving laser pulse propagation within hollow core capillaries
- Characterisation of a focused XUV beam within a nanoscale imaging set up
- Imaging of a binary sample with a HHG source

In chapter 2 background to the processes and techniques used later in the thesis are given.

In chapter 3 details of the laser system used to conduct the experiments in this thesis are given. A description of the techniques used to monitor the output of the laser are also given as this is critical to the efficacy of the HHG process.

In chapters 4 and 5, experiments were carried out to test the validity of modelling the propagation of the driving laser pulse within a capillary. The chapters begin with a description of fabrication, mounting and coupling of a driving laser into a capillary. Details of experiments carried out to validate two different models are then presented.

In chapter 6 the results from experiments performed to characterise the XUV beam in a geometry suitable for imaging will be shown. Measurements were made to determine
the position of the focus and the spectrum at the focus, both of which are critical for
the experiment described in chapter 7.

Finally, in chapter 7 the preliminary results from the imaging setup described in chapter
6 are demonstrated. This includes a description of the sample used and its fabrication.
A discussion of using a monochromatic algorithm with a polychromatic beam and the
meaning of the output from this algorithm.

The work in this thesis is based on experiments using the HHG process within the
Ultrafast X-ray group at the University of Southampton.
Chapter 2

Background and theory

In this chapter the background and theory relating to the experiments carried out in chapters 4, 5, 6 and 7 will be covered. The motivation behind the project is to build a HHG imaging system, the stages of which are covered by source development, characterisation of the XUV focusing setup and finally an imaging experiment. To develop the source an understanding of the process of HHG via both a semi classical and a quantum mechanical model will be discussed. It is also important to understand the factors that limit and change the generation process, this will be covered by the phase matching, absorbance and non-linear optics sections. The generation geometry used for the experiments in chapters 4 and 5 involve coupling into and propagation along a waveguide which will be discussed in capillary waveguide modes. Chapter 5 also uses a non-linear Schrödinger equation model for propagation which is outlined in this chapter. The final section will discuss the theory behind coherent diffractive imaging which is used in chapter 7.

2.1 High Harmonic Generation

The process of HHG requires the interaction between a target gas, usually a noble gas and a high intensity driving laser field. The intensities required are in excess of $10^{14} W cm^{-2}$ and as a consequence require the use of femtosecond laser systems. There are two different ways of modelling this process, either a semi-classical model introduced by Corkum *et al.* [35] where the electron is treated as a point charge in a sinusoidal field,
Figure 2.1: The three step model of HHG (a) the potential well of the atom is distorted, the electron tunnels out of the atom and is accelerated by the laser driving field (b) the electron is accelerated back towards the parent atom as the direction of the driving field changes (c) the electron recombines with the parent atom with higher energy that is then emitted as a XUV photon.

or a quantum mechanical treatment of the electron using a time dependant Schrödinger equation [36]. Both of these models make accurate predictions but there are limitations in the semi classical model which will be discussed in the next section.

2.1.1 Semi Classical Model

The semi classical model is known as the three step model because the process can be broken down into three distinct stages, shown in figure 2.1.

During the first step the intense laser field causes the Coulomb potential of the well containing a valence electron to distort far enough that a potential barrier is formed, allowing the electron to quantum tunnel out into free space. This electron is then accelerated in the electric field of the laser. Step two occurs as the direction of the laser field changes driving the electron back towards its parent ion. The third and final step of the process is the recombination of the electron with the parent ion. During the electron’s time in the laser field it has gained extra kinetic energy which is now released as an XUV photon.
Using classical mechanics equations, the trajectories (equation 2.1) of the electron in the continuum can be calculated along with its velocity (equation 2.2) and acceleration (equation 2.3). The laser field is modelled as a sinusoidal $E = E_0 \sin(\omega t)$, where $E_0$ is magnitude of the electric field, $\omega$ is the angular frequency and $t$ is a point in time. Several assumptions are also made, the electron has zero velocity at time $t = t_0$, a position of $x = 0$ at time $t = t_0$, the electron release time and time $t = t_1$, the electron return time.

$$x = \frac{eE_0}{m\omega} \left( \frac{1}{\omega} (\sin(\omega t_0) - \sin(\omega t)) + (t - t_0) \cos(\omega t_0) \right)$$  \hspace{1cm} (2.1)$$

$$v = \frac{eE_0}{m\omega} (\cos(\omega t_0) - \cos(\omega t))$$  \hspace{1cm} (2.2)$$

$$a = \frac{eE_0}{m\omega} (\sin(\omega t))$$  \hspace{1cm} (2.3)$$

where $e$ is the charge of an electron, $m$ is the mass of an electron, $t_0$ is the release time of the electron.

From these equations it is possible to see that the trajectories that recombine only occur at certain times in the optical cycle. These conditions mean that there is a burst of XUV generation every half optical cycle of the pulse leading to a train of attosecond pulses. As the phase of the driving laser is changing by $\pi$ between each burst of XUV the spectrum is formed from an odd function, therefore only odd harmonics are produced. By calculating the velocity of a returning electron the energy of the emitted photon can be calculated[32]. Figure 2.2 shows the energy gained by the electron in the laser field by solving equation 2.2 as a function of travelling time ($\tau$), given by,

$$\tau = t_1 - t_0$$  \hspace{1cm} (2.4)$$

where $t_1$ is the return time of the electron. From figure 2.2 it can be seen that away from the maximum there are two values for each energy, one for the short trajectory ($\tau_1$) and one for the long trajectory ($\tau_2$).
Figure 2.2: The energy gained by an electron in the continuum as a function of time in units of $U_p$. $\tau_1$ represents the short trajectories and $\tau_2$ represents the long trajectories. It is also shown that the maximum value for the energy gained is $3.17U_p$.

The maximum energy gained by the electron is therefore $3.17 U_p$ where $U_p$ is the ponderomotive energy of the electron and relates to the cut off energy by,

$$E_{\text{max}} = I_p + 3.17U_p$$  \hspace{1cm} (2.5)

Where $I_p$ is the ionisation potential of the atom in eV and $U_p$ is given by,

$$U_p = \frac{e^2E_0^2}{4\omega^2m}$$  \hspace{1cm} (2.6)

This model accurately reproduces the cut-off energy, the trajectories and the return energy of the electron but it cannot explain where the non-linearity comes from and also it does not explain how the recombination occurs. To show the non-linearity and recombination a quantum mechanical model is required.

2.1.2 Quantum Mechanical Model

To understand the origins of the non-linearity in the HHG process, a quantum mechanical approach is used. The model uses a classical driving field as described in the previous
section, although the electron is treated as a wavefunction inside a soft Coulomb potential [37] rather than as a point charge. The soft Coulomb potential is a numerical requirement; a standard coulomb potential described by,

\[ V = -\frac{1}{4\pi\epsilon_0} \frac{e^2}{x} \]  

(2.7)

where \( \epsilon_0 \) is the permittivity of free space and \( e \) is the charge of an electron. As \( x \) tends towards 0 the solution tends towards \(-\infty\) causing a singularity that the numerical calculations cannot handle. To avoid this problem a soft coulomb potential is used where the Bohr radius (\( \alpha_0 \)) is introduced, shown in equation 2.8, to prevent the singularities arising.

\[ V = -\frac{1}{4\pi\epsilon_0} \frac{e^2}{(\alpha_0^2 + x^2)^{1/2}} \]  

(2.8)

A time dependant Schrödinger equation [38], shown in equation 2.9, is used to model the wave function as it interacts with the electric field of the driving laser and the soft Coulomb potential.

\[ i\hbar \frac{\partial}{\partial t} \psi = \hat{H}\psi \]  

(2.9)

The driving laser field adds to the soft Coulomb potential resulting in a potential barrier being formed at specific cycles through the pulse. Part of the electron wavefunction can be accelerated away from the nucleus of the atom, this distribution can extend beyond 5 nm. Figure 2.3 shows the probability distribution of the electron wavefunction with a 7 fs driving laser pulse overlaid on top. The greatest density can be seen throughout the plot at the centre, close to the atom, but as the magnitude of the field of the pulse increases then so does the distance the electron wavefunction may be found from the atom. As the sign of the field changes so does the probability distribution of the wavefunction causing the wavefunction to return past the bound portion of the wavefunction. The returning portion of the wavefunction has a different frequency to the bound portion of the wavefunction due to the acceleration of the driving field. This frequency difference causes an interference between the the portions of the wavefunction which leads to a dipole oscillation at the at the XUV frequency.

The simulation shown was produced by Dr W. Brocklesby and is used by other members of the group in predicting XUV spectra.
2.2 Phase Matching

For any type of harmonic generation it is important to maximise the efficiency of the process, therefore the harmonics being generated should not be destructively interfering, reducing the total harmonic signal. To achieve this, the phase of the driving laser field at the moment of generation must be the same as the phase of the harmonics that have already been produced earlier in the medium. If the phases are the same at the moment of generation then every harmonic generated will add up constructively and give the largest possible signal, this is the case when the phase mismatch (∆k) is zero.

\[ \Delta k = k_q - qk_f = 0 \]  \hspace{1cm} (2.10)

where \( k_f \) is the fundamental wave vector, \( k_q \) is the harmonic wave vector and \( q \) is the harmonic order. Figure 2.4 shows a phase matched and a non phase matched condition for second harmonic generation. The intensity of the harmonic signal generated in a phase matched case will continue to increase for as long as the phase matching conditions can be maintained. If the phases are \( \pi \) out of phase then the signal will be interfere destructively.

There are four contributions to the phase mismatch in a high harmonic experiment.
Figure 2.4: Example of phase matching in SHG, (a) shows a coherent build up of the harmonics as they are generated in phase, (b) shows the non phase matched case where the harmonics interfere destructively.

[39][40], a geometric term $\Delta k_{geo}$, a neutral atom term $\Delta k_{na}$, a free electron term $\Delta k_{fe}$ and an atomic phase term $\Delta k_{at}$ as shown by,

$$\Delta k = \Delta k_{geo} + \Delta k_{na} + \Delta k_{fe} + \Delta k_{at}$$  \hspace{1cm} (2.11)

Combining equations 2.10 and 2.11 would give perfect phasematching between the driving pulse and the generated harmonics for all distances. In practice a $\Delta k = 0$ is exceptionally difficult to achieve. A common measure is therefore the coherence length ($L_{coh}$), this is a measure of the distance required for a $\pi$ phase shift between $k_{q}$ and $qk_{f}$ given by,

$$L_{coh} = \frac{\pi}{\Delta k}$$  \hspace{1cm} (2.12)

Each term in equation 2.11 can be calculated for a HHG experiment. The geometrical term is dependent upon the generation geometry, in a capillary the term relates to the waveguide properties of the capillary [41] and is given by

$$\Delta k_{geo} = \frac{u_{11}^{2} \lambda_{f}}{4\pi a^{2}}$$  \hspace{1cm} (2.13)

Where $u_{11}^{2}$ is the Mode factor of the waveguide and $a$ is the bore radius of the waveguide. The propagation of the fundamental through a capillary has a higher phase velocity than
in free space, while the harmonics do not see a contribution from the waveguide as the Mode size is much smaller than the bore size of the capillary.

For a gas cell or gas jet setup the waveguide term does not exist, instead a phase mismatch is caused by the Gouy phase shift [42][43] of a Gaussian beam propagating through a focus which is defined as,

$$\phi_G = \arctan\left(\frac{z}{z_0}\right) \quad (2.14)$$

where $z$ is a longitudinal distance and $z_0$ is the Raleigh length. This derivative of equation 2.14 gives the propagation vector $k$ so the phase mismatch, $\Delta k_{geo}$ is shown by,

$$\Delta k_{geo} = \frac{q z_0}{z^2 + q^2 z_0^2} - \frac{q z_0}{q^2 z_0^2 + z_0^2} \quad (2.15)$$

The result of this means that there is a longitudinal dependence upon the phase mismatch so the position of the gas cell or jet to the focus is critical.

The neutral atom term is the phase mismatch due to the different frequency components travelling through the medium seeing a different refractive index,

$$\Delta k_{na} = q p (1 - \eta) \frac{2\pi}{\lambda_f} (\Delta\delta + n_2) \quad (2.16)$$

where $p$ is the pressure inside the waveguide, $\eta$ is the ionisation level, $\Delta\delta$ is the difference in the refractive index for each frequency per atmosphere and $n_2$ is the non linear refractive index for the fundamental frequency.

The process of HHG requires ionisation to occur, this means that there will always be a free electron component in the phase matching calculations. The free electron term is only considered for the fundamental as the effect on the XUV is negligible [44]. The free electron term

$$\Delta k_{fe} = q p \eta N_a r_e \lambda_f \quad (2.17)$$

where $N_a$ is the number density of atoms and $r_e$ is the classical electron radius, is only valid if the ionisation level is below the critical ionisation level. The critical ionisation level is the point at which the free electron and neutral atom terms balance, in argon it is 4.8% [39].
The final phase matching term is the atomic phase. This is the intrinsic phase of the harmonic and is a result of the kinetic energy gained by the electron in the driving laser field, shown by,

$$\phi_{at} = \frac{S(t_0, t_1)}{\hbar} + q\omega t$$  \hspace{1cm} (2.18)

where $S(t_0, t_1)$ is given by,

$$S(t_0, t_1) = \int_{t_0}^{t_1} \left( \frac{m_e v(t)^2}{2} + I_p \right) dt$$  \hspace{1cm} (2.19)

where $I_p$ is the ionisation potential of the target gas. Due to the combination of the time and intensity dependence it can be seen that there is a small variation across the beam for short trajectories [45] but a much larger variation for the long trajectories. As a result the long trajectories have a much higher divergence and can be considered negligible at long propagation distances [46].

### 2.3 Absorption

In the previous section the coherence length described the length over which the generated harmonics will constructively interfere. This defined a maximum length for the generation region of the observed output signal. There is another factor to consider at this point, the absorption of materials in the XUV region is high. The generation process requires there to be a target material to generate off, this same material can then act as an absorber for the generated harmonics. The refractive index of materials in this region can be written as,

$$n(\omega) = 1 - \delta(\omega) + i\beta(\omega)$$  \hspace{1cm} (2.20)

where $\delta(\omega)$ is the dispersive part of the refractive index and $\beta(\omega)$ is attenuation coefficient part of the refractive index. The above equation shows that the refractive index for materials in the XUV region is less than 1. The $\beta$ value can be used to calculate an absorption length ($L_{abs}$) by,

$$\frac{I}{I_0} = \frac{1}{e} = exp[-\beta(\omega)k(\omega)L_{abs}]$$  \hspace{1cm} (2.21)
Figure 2.5: Transmission of XUV for (a) 1cm of argon at 250mbar (b) a 200nm aluminium foil. Data from reference [47]

\[ L_{\text{abs}} = \frac{1}{\beta(\omega)k(\omega)} \]  \hspace{1cm} (2.22)

The attenuation coefficient for argon at 30 nm is given as \( 2.1556 \times 10^{-5} \) [47] which gives an absorption length of 222 \( \mu \)m at 1 atm. The absorption scales linearly with pressure so at 100 mbar the absorption length is 2.2 mm. This means that for the capillary setup the XUV generated at the front of the capillary will be reabsorbed before the capillary exit.

In a later chapter the use of thin foil aluminium filters (180 nm Al + 20 nm Al\(_2\)O\(_3\)) will be discussed along with silicon nitride membranes (Si\(_3\)N\(_4\)) as a sample supports. Figure 2.5 shows the wavelength dependant absorption of these materials for thin samples. This demonstrates the difficulty in working with any material in transmission.

2.4 Non-linear optics

Non-linear optics refers to the relationship between the induced polarisation of a material and the incident field upon it. At low intensities the material will respond linearly to the field. As the field increases in strength the induced polarisation begins to show an
anharmonic response. This can be written as,

\[ P(t) = \chi^{(1)} E(t) + \chi^{(2)} E^2(t) + \chi^{(3)} E^3(t) + \ldots \]  

(2.23)

where \( P(t) \) is the induced polarisation at time \( t \) and \( \chi^{(n)} \) is the \( n^{th} \) order susceptibility of the material [48].

The \( \chi^{(2)} \) term is responsible for SHG and SFG. There is a requirement that the material has an asymmetry, which is not the case in the low pressure gases used for HHG, as a result this term is equal to zero.

The main effects relevant to this thesis come from the \( \chi^{(3)} \) term. The optical Kerr effect manifests itself both spatially and temporally. The cause of the effect is a change to the instantaneous refractive index \( \tilde{n} \) by,

\[ \tilde{n} = n_0 + n_2 \tilde{I} \]

(2.24)

where \( (n_2) \) is non-linear refractive index and \( \tilde{I} \) is the instantaneous intensity at a given point in time and space.

The spatial effect known as Kerr self focusing [49] is used to create the mode-locked femtosecond seed pulse. This occurs due to intensity varying across the profile of the beam as a result, \( \tilde{n} \) varies across the beam. The distribution of the refractive index has a focusing effect on the beam as the refractive index is higher at the centre of the beam than the wings. This is the same as a convex lens. A cavity can therefore be designed in such a way that propagation in a pulsed mode is favourable to propagation in a continuous mode.

Related to the Kerr lensing effect is self phase modulation (SPM)[50]. This is the temporal result of the changing refractive index. As the pulse propagates in time the intensity of the pulse changes causing the change in refractive index described by equation 2.24. This change effects the transmitted phase of the pulse resulting in a frequency shift. The phase change results in the leading edge of the pulse generating red frequencies and the trailing edge of the pulse generating blue frequencies.

The final effect to consider is self steepening. This effect is related to SPM as it is a temporal intensity dependent effect. The intensity at the centre of the pulse is greater
than the wings, equation 2.24 shows that the refractive index will be higher at this point so the group velocity is reduced. As the intensity reduces towards the trailing edge of the pulse the group velocity increases again giving a steepened trailing edge. The result of this is that a greater frequency shift at the trailing edge of the pulse so the spectral broadening is dominated by the blue shifted part of the pulse.

2.4.1 Ionisation effects

The process of SPM described above shows a blue shifting effect on the pulse. In regions of ionisation a similar effect occurs due to the plasma reducing the refractive index. As the pulse propagates the rate of tunnel ionisation changes as described by Keldyish theory,

$$W(x, t) = W_0 \kappa^2 \sqrt{\frac{3}{\pi}} \frac{C_{el} 2^{2n^*} F(x, t)^{1.5 - 2n^*}}{\kappa l^2} \exp \left( -\frac{2}{3F(x, t)} \right)$$

(2.25)

where \( \kappa^2 = I_p/I_H \) is the ratio of the ionisation potential of the target gas \( I_p \) over the ionisation potential of hydrogen \( I_H \), \( W_0 = m_e e^4/\hbar^3 \) and \( F(x, t) \) is the reduced electric field. Parameters \( C_{el} \) and \( n^* \) can be obtained from look up tables [51]. Using equation 2.25 the ionisation can be calculated as a temporal effect within the pulse. Once the threshold is reached the ionisation increases rapidly creating a snap reduction in the refractive index with respect to time. This has the associated effect of producing a rapid change in phase causing changes to the frequency and hence broadening the spectrum. The spatial variation of this effect leads to a lower refractive index in the centre of the beam than the wings. This results in the opposite effect to the Kerr self-focusing producing a plasma defocusing.

The effect of these terms are critical to understanding the processes shown in chapter 5 and in the creation of the model outlined in section 2.7

2.5 Gaussian Beams

The output of many laser systems is a beam with a Gaussian transverse spatial profile, often referred to as a TEM\(_{00}\) beam[52]. The equation for a Gaussian beam can be written out as,

$$E(r) = E_0 \exp \left( -\frac{r^2}{w^2} \right)$$

(2.26)
where $E_0$ is the magnitude of the field at $r = 0$, $r$ is the radial position and $w$ is the radius of the beam at $E = \frac{E_0}{e}$.

When focusing a Gaussian beam there are several properties that need to be understood. The position of the beam waist is needed in order to be able to position a sample at the position of highest flux although this is not always as simple as it may first seem. Understanding the size of the beam waist ($w_0$) and how this can be affected by different conditions is also critical to making the brightest possible source. The beam waist size is related to the Rayleigh length ($z_R$) by,

$$z_r = \frac{\pi w_0^2}{\lambda} \tag{2.27}$$

using these two parameters it is possible to calculate the spotsize at any point away from the focus using,

$$w(z) = w_0 \sqrt{1 + \frac{z^2}{z_r^2}} \tag{2.28}$$

It is also possible to calculate the wavefront curvature at a give point from the beam waist using the Rayleigh length by,

$$R = z + \frac{z_r^2}{z} \tag{2.29}$$

It is therefore possible by knowing one of the three parameters above to calculate the others. It is important to remember that these equations are for an idealised single mode Gaussian beam. In practice the output of lasers is not a single mode Gaussian beam but a superposition of Gaussian modes. A measure of a non diffraction limited Gaussian beam is called the M$^2$[53]. This parameter relates to the Rayleigh length ($z_R$ described in equation 2.27 by,

$$z_r = \frac{\pi w_0^2}{\lambda M^2} \tag{2.30}$$

as a result has an effect on both the spot size and radius of curvature.

Once a Gaussian beam has been defined a method for describing its interactions with optics such as lenses and mirrors is required. The section below will describe one method for achieving this.
2.5.1 Propagation using ray transfer matrices

Propagation of Gaussian beams through a system can be modelled using ray transfer matrices [54]. This system uses a series of two by two matrices to describe the effect of each element in the path of the Gaussian beam. First the Gaussian beam at the input is described by the complex $q$ parameter,

$$\frac{1}{q} = -i \frac{\lambda}{\pi w^2} + \frac{1}{R}$$

(2.31)

where $\lambda$ is the wavelength of the Gaussian beam, $w$ is the $\frac{1}{e^2}$ width of the beam and $R$ is the radius of curvature of the wavefronts. Having defined $q$ it is now possible to calculate its value at any point ($q'$) by multiplying the appropriate matrix as shown below,

$$q' = \frac{Aq + B}{Cq + D}$$

(2.32)

where $A, B, C$ and $D$ are given by elements in a two by two matrix,

$$\begin{pmatrix}
A & B \\
C & D
\end{pmatrix}$$

The elements within this matrix differ as the propagation environment changes. Below are listed the three relevant to the work contained in later chapters. For propagation through free space over a distance $d$ the following matrix is used,

$$\begin{pmatrix}
1 & d \\
0 & 1
\end{pmatrix}$$

For a lens with a focal length $f$,

$$\begin{pmatrix}
1 & 0 \\
-\frac{1}{f} & 1
\end{pmatrix}$$

For a spherical mirror with a radius of curvature $R_m$,

$$\begin{pmatrix}
1 & 0 \\
-\frac{2}{R_m} & 1
\end{pmatrix}$$
A spherical mirror is rarely used at an angle normal to the input beam so a correction must be applied. For an incident angle $\theta$ in the horizontal plane $R_m = R_m \cos \theta$ in the tangential plane and $R_m = \frac{R_m}{\cos \theta}$ in the sagittal plane.

Using this method it is possible to simulate the effect of propagating a beam through free space, through a lens or onto an off axis focusing mirror and through space to the focus. This allows the calculation of the position and expected beam waist for an optic taking into account all parameters of an initial Gaussian beam. This technique is used in chapter 6 to model the focal spot from an off axis focusing optic.

2.6 Capillary Waveguide Modes

The previous section describes the propagation of Gaussian beams. When a Gaussian beam reaches a the entrance of a hollow capillary waveguide the propagation is restricted to the modes of that capillary. This section will describe the process of coupling into those modes and their propagation along the capillary.

When a Gaussian beam is coupled into a hollow capillary waveguide it can no longer maintain the same field profile or propagation term as if it were in free space. If a linearly-polarised TEM$_{00}$ source is coupled into a hollow capillary only the EH$_{1m}$ modes will be excited. EH$_{1m}$ modes are Bessel modes of the first kind.

Figure 2.6 illustrates the intensity and electric-field distribution in the EH$_{11}$ to EH$_{14}$ modes. Marcatili and Schmeltzer [55] derived expressions for the electric field of the EH$_{1m}$ waveguide modes which can be simplified to give

$$ E_{1m} = J_0 \left( u_{1m} \frac{r}{a} \right) $$  \hspace{1cm} (2.33)

where $a$ is the waveguide radius and $u_{1m}$ is the $m^{th}$ root of the Bessel function of the first kind $J_0$. The mode propagation for each individual mode can be expressed as

$$ E_{1m} = E_{1m0} \exp i(\gamma z - \omega t) $$  \hspace{1cm} (2.34)

where $\omega$ is the angular frequency and $\gamma$ is the propagation constant, given by

$$ \gamma = \beta + i\alpha. $$  \hspace{1cm} (2.35)
Figure 2.6: The two-dimensional intensity distribution of the EH$_{11}$ to EH$_{14}$ modes (a,c,e,g) normalised to maximum intensity and the associated line plots for the EH$_{11}$ to EH$_{14}$ modes (b,d,f,h)
Here $\alpha$ is the modal attenuation and $\beta$ is the phase velocity defined as

$$\alpha = \left( \frac{um}{2\pi} \right)^2 \frac{\lambda^2 (v^2 + 1)}{a^3 \sqrt{v^2 - 1}}$$

(2.36)

$$\beta = \left( \frac{2\pi}{\lambda} \right) \left\{ 1 - \frac{1}{2} \left[ \frac{um\lambda}{2\pi a} \right]^2 \right\}$$

(2.37)

where $v$ is the refractive index of the waveguide material. The total electric field at a given propagation distance is obtained by calculating the sum of the fields for each mode at that distance.

The coupling efficiency of an incident Gaussian beam to each of the EH$_{1m}$ capillary modes can be calculated by the overlap integral [56]

$$A_m = \frac{\iint E_{1m} \times E_i \ r \ dr \ d\theta}{\iint |E_{1m}|^2 \ r \ dr \ d\theta}$$

(2.38)

where $E_i$ is the incident electric field and $r$ is the radius of the capillary.

In most capillary-based HHG experiments, the intention is usually to couple the maximum energy into the lowest order EH$_{11}$ mode because the higher order modes are more lossy as shown in equation 2.36 by the $\left( \frac{um}{2\pi} \right)^2$, a square dependence upon the mode. Figure 2.7 shows the percentage of each mode coupled into for a perfectly aligned TEM$_{00}$ input beam with a changing ratio of beam waist to capillary bore size. It can be seen from figure 2.7 that the optimum condition for coupling into the EH$_{11}$ mode is given by $w = 0.64a$, where $w$ is the radius at 1/$e^2$ of the intensity of the beam. However, there are not a set of coupling conditions for which there is zero probability of coupling into higher order modes. At optimum coupling for the EH$_{11}$ mode, 2% of the energy will be coupled into higher order modes. As a result when coupling a TEM$_{00}$ beam into a capillary a superposition of two or more capillary modes will always be present.

For decreasing spot sizes from this ideal value, increasingly higher order modes will be excited. Most efficient coupling into the EH$_{12}$ mode may be achieved for a waist size to bore ratio of $w/a = 0.26$, with 42% of the power coupled into the EH$_{12}$ mode.

The result of this superposition of modes is a periodic intensity variations along the propagation direction of a capillary, an effect referred to as mode beating.
Figure 2.7: Calculated normalised coupling efficiency $\chi$ of a perfectly aligned TEM$_{00}$ beam as a function of the spot size to bore radius ratio into the modes, where $\rho_m = \frac{A_m^2}{\int \int E_r^2 r dr d\theta}$.

The beat length $L$ between two modes is given by

$$L = \frac{2\pi}{\Delta \beta} \quad (2.39)$$

where $\Delta \beta$ is the difference in the real part of the propagation constant of the two capillary modes, which for the EH$_{11}$ and EH$_{12}$ modes is given by

$$\Delta \beta = \frac{\lambda}{4\pi a^2} \left[ u_{12}^2 - u_{11}^2 \right]. \quad (2.40)$$

Figure 2.8(a) shows the radial intensity in a 75 $\mu$m radius, 70 mm long capillary, plotted as a function of position along the capillary for a superposition of modes optimally coupling into the EH$_{11}$ but with contributions from higher order modes where the beam waist to bore ratio $w/a = 0.64$ and an input power of 800 $\mu$J. Figure 2.8(b) shows the radial intensity in the same capillary with 100% coupling into the EH$_{11}$ mode. In the case of mixing equal proportions of the EH$_{11}$ and EH$_{12}$ modes, constructive interference between the two modes results in a narrow spatial profile which increases the peak intensity. As the beam propagates the modes acquire a $\pi$-phase shift with respect to
Figure 2.8: Calculated radial intensity distribution for a 70 mm long, 75 µm radius capillary, for (a) a superposition of modes which corresponds to $w/a = 0.64$, optimal for coupling into $EH_{11}$ compared with (b) $EH_{11}$ mode. The plots are calculated using equations 2.33 and 2.34, both are normalised by the integrated power.

one another. This corresponds to the destructive interference case, the intensity at the centre of the capillary will be zero and the intensity distribution forms a toroidal profile. The intensity continues to oscillate between these two extremes, resulting in a periodic intensity modulation along the propagation axis of the capillary. In the case of optimal coupling the $EH_{11}$ mode dominates and there are small contributions from several higher order modes which mean that the intensity drops to near zero, but is never actually zero.

Even in a well designed experiment, where the spot size has been carefully engineered to match the optimum size, one must contend with random misalignments of the two angular and three spatial dimensions involved. These can be accounted for in the calculation of the overlap integral (equation 2.38) and figure 2.9 shows the result of using equation 2.38 to calculate the percentage of energy coupled into the $EH_{11}$ mode as a function of total transmission through the capillary for various misalignments of the input beam. This calculation was performed by Dr. E. Rogers.
Figure 2.9: Percentage of power coupled into the EH$_{11}$ mode as a function of total transmission through the capillary for random alignments of a beam of given spot size ($w = 0.64a$) in a five-dimensional space including the lateral position of the beam relative to the centre of the capillary, beam waist position and angular alignment of the beam relative to the capillary axis. This figure is adapted from [57].

As the total transmission increases above 50%, the percentage of energy coupled into the EH$_{11}$ mode is greatly increased. Experimentally it is generally easy to get transmissions up to 50% and increasingly hard thereafter. There are many more alignment combinations available that lead to coupling to the higher order modes but most of these are highly lossy. This shows that unless great care is taken to couple as much light as possible through the capillary, a high percentage of higher order modes will be present, leading to complicated intensity modulations along the length of the capillary.

2.7 The non-linear Schrödinger equation model

The work carried out in chapter 5 used a model developed by Dr P. Horak. This section will give an outline to that model, its origins and the modifications made to suit the experiment.

Modelling propagation in multimode fibres had been solved numerically by using a multimode generalised nonlinear Schrödinger equation (G-NLSE) [58]. Propagation of a pulse is modelled numerically in steps along the z-axis. The field amplitude $A_n$ changes as,

$$\frac{\partial A_n}{\partial z} = D\{A_n\} + N\{A_n\}$$  \hspace{1cm} (2.41)
where $D\{A_n\}$ is the modal dispersion term and $N\{A_n\}$ is a non-linear response term for silica. For the accurate theoretical description of the driving laser pulse propagation in a gaseous medium the non-linear response term was modified. The numerical model was extended to include ionisation and plasma effects [59] present within a gas filled capillary. The electric field of the laser pulse was written as a sum over modal contributions,

$$E(r,z,t) = \sqrt{\frac{2Z_0}{2\pi}} \sum_n \frac{1}{2} F_n(r) A_n(z,t) e^{i(k_0z-\omega_0t)} + c.c. \quad (2.42)$$

where $Z_0 = \sqrt{\mu_0 \epsilon_0}$, $F_n(r)$ is the transverse mode function of the $EH_{1m}$ mode [55] of the capillary normalised to $\int r dr |F_n(r)|^2 = 1$ and $A_n$ is the pulse envelope such that the modal power is given by $\left| \frac{A_n}{2\pi} \right|^2$ in units of Watt. The temporal dependence of the carrier wave has been factored out. In equation 2.42 the analysis has been restricted to circularly symmetric modes only, assuming a symmetric pump laser mode and preferential coupling between modes of the same symmetry [60]. The modal evolution of the pulse along the capillary is then given by

$$\frac{\partial A_n}{\partial z} = D\{A_n\} + N\{A_n\} + N_{pl}\{A_n\} + L_{ion}\{A_n\} \quad (2.43)$$

where $N_{pl}\{A_n\}$ is non-linear response of the plasma and $L_{ion}\{A_n\}$ is the ionisation induced loss. Each of these terms can be expanded as shown below.

$$D\{A_n\} = iD_n \quad (2.44)$$

$$N\{A_n\} = \left(1 + \frac{i}{\omega_0} \frac{\partial}{\partial t} \right) in_2(z)k_0 \sum_{k,l,m} Q_{nklm} A_k A_l A_m^* \quad (2.45)$$

$$N_{pl}\{A_n\} = \left(1 - \frac{i}{\omega_0} \frac{\partial}{\partial t} \right) \frac{1}{2} k_0 \int r dr F_n(r) S(r,z,t) n_{pl}^2(r,z,t) \quad (2.46)$$

$$L_{ion}\{A_n\} = -\frac{1}{2} \int r dr F_n(r) S(r,z,t) \frac{\rho_0(r,z,t) W(r,z,t) U}{|S(r,z,t)|^2} \quad (2.47)$$

where $S = \sum F_n A_n$. In equation 2.44 $D_n$ describes modal dispersion and losses, including high-order dispersion, and the subsequent terms model the gas nonlinearity, plasma refractive index, and ionisation losses of the pump, respectively. In Equation 2.45, $n_2(z)$ is the nonlinearity of the neutral Ar gas as a function of the local pressure, $k_0$ is the propagation constant at the pump frequency, $Q_{nklm} = \int r dr F_n F_k F_l F_m$ are the mode overlap integrals. In equation 2.46, $n_{pl}$ is the refractive index of the local plasma density.
resulting from ionisation of the neutral gas by the propagating pulse. In equation 2.47, \(\rho_0\) and \(W\) are the neutral gas density and ionisation rate, respectively, and \(U\) is the ionisation potential. The ionisation rate \(W\) was calculated using equation 2.25 [51]. The model used in chapter 5 solved equation 2.43 by a Fourier split-step method for typically 20 modes, which was sufficient for an accurate description of experimental parameters. This model was capable of describing the full temporal dynamics of the propagating light in three dimensions, including modal effects such as group velocity mismatch.

2.8 Coherent Diffractive imaging

In a microscopy experiment the aim is to observe objects smaller or with finer detail than the eye can see. The imaging limit of a microscope is related to the quality of the lenses used within it [6] and is given by

\[
R = \frac{\lambda}{2NA}
\]

(2.48)

where \(\lambda\) is the the illumination wavelength and \(NA\) is the numerical aperture of the lens used. As the quality of imaging lenses reaches a limit, reduction of the illumination wavelength provides an alternative route to improved resolution.

In the XUV region traditional lenses are not feasible due to high material absorptions discussed in section 2.3. A solution to this problem is the use of iterative phase retrieval algorithms as an alternative to an imaging lens. This technique is known as coherent diffractive imaging (CDI).

These phase retrieval algorithms require a coherent illuminating beam to produce a far field diffraction pattern from an aperiodic sample. An outline of all the requirements for a CDI experiment and a description of the process the algorithm undertakes will be shown in the next section.

2.8.1 Diffraction

Diffraction is the interaction when a propagating wave meets a partially transmitting obstacle. The propagation of the wave is perturbed by the interaction with the obstacle and results in a modified wave front propagating after the obstacle. A simple form of
diffraction is through a slit as shown in figure 2.10. The Huygens-Fresnel principle [52] states that every point on the wave front is a point source from which a secondary wave propagates. The propagated wave front can be considered to be the coherent sum of all these secondary waves. Using this principle a wave propagating through a sample will interact to give a unique field at all points after the sample.

X-ray diffraction is routinely used to reveal the structures of periodic samples. The weakly coherent beam from a rotating anode source [61] requires the repeating unit cell of the sample to build up intensity at discrete points known as the Bragg peaks. Using the separation and relative intensities of these peaks the structure of the sample can be determined. The use of x-ray diffraction has allowed observation of atomic structures from small organics to proteins [3]. The study of non periodic samples however, requires the coherence to be present in the beam rather than the sample. This is currently achieved at user facilities such as synchrotrons and FEL’s as well as HHG. The current lab based X-ray sources used for single crystal diffraction are not suitable for CDI as the coherence is too low.

The diffraction pattern observed for a particular sample is dependant upon the distance from the object when it was sampled. There are two regimes under which most diffraction patterns fall, these are the Fresnel regime (near-field) and the Fraunhofer regime (far-field).
2.8.2 Fresnel and Fraunhofer regimes

The difference between these two conditions relates to the curvature of the wavefront at the detector. In the case of Fresnel diffraction the wavefront is curved at the detector. As the beam propagates further the curvature of the beam reduces until at the detector it appears to be flat. This point is the condition for far-field diffraction. The diffraction pattern at any distance from an object can be calculated using the angular spectrum method (ASM) [62]. However the far-field diffraction is the Fourier transform of the electric field immediately after the object [6]. Once the condition has been reached, extending the distance has the effect of changing the angular scale of the pattern and does not change the form. Using this information it is possible to write CDI routines around a series of Fourier and inverse Fourier transforms. This is computationally less intensive than using ASM to iterate between the sample space and diffraction space. This does not preclude CDI from near field diffraction experiments [63], an extra constraint of accurately measuring the sample to detector distance would be added to the experiment.

To determine the diffraction regime at a particular distance the Fresnel number \( F \) is calculated. The condition for Fresnel diffraction is calculated by,

\[
F = \frac{a^2}{z\lambda} \geq 1.
\]  

(2.49)
The condition for Fraunhofer diffraction is calculated by,

\[ F = \frac{a^2}{2\lambda} \ll 1. \]  

(2.50)

where \( \lambda \) is the illuminating wavelength, \( z \) is the aperture to detector distance and \( a \) is the size of the aperture as shown in figure 2.11. It is however, not possible to apply a simple Fourier transform to invert a recorded image to the original object. When the diffraction pattern is recorded the phase information is lost and the recorded intensity is the modulus squared of the electric field of the beam at the object.

2.8.3 Coherence

To perform coherent diffractive imaging a spatially coherent source is required. Spatial coherence is the phase correlation of a source at two points across the beam profile. The spatial coherence of a source can be inferred by the visibility of interference fringes using the source and a diffraction grating. The greater the spatial coherence the sharper the interference fringes will appear. In an incoherent source the fringes will destructively interfere resulting in a smooth signal on the detector. To perform a CDI experiment the spatial coherence must be greater than the longest axis of the sample, where the sample is the transmitting object of interest. If a pinhole is used to constrain the sample then the spatial coherence must be greater than the diameter of the pinhole.

2.8.4 The Algorithm

Coherent diffractive imaging uses phase retrieval algorithms published by Gerchberg and Saxton in 1972 [7]. These algorithms were developed for electron diffraction and extended by Fienup [8][9] to apply to the optical regime. The technique replaces the imaging optic used in conventional microscopy with an iterative algorithm. When a detector collects the intensity distribution from the diffraction pattern in the far-field the phase information is lost. An inverse Fourier transform will therefore not match the sample electric field immediately after the sample but instead gives the autocorrelation of the sample as defined by the Wiener-Khinchin theorem [62].

Phase retrieval algorithms work by iterating between the diffraction space (diffraction pattern) and the sample space (image of the sample) using the Fourier relationship shown
in figure 2.12. A support mask is generated from the autocorrelation by thresholding to a proportion of the maximum value. This mask is known to contain the sample and is used within the algorithm as the support constraint. An initial guess at the sample is created using the support constraint as the amplitude with a random phase. This initial guess is Fourier transformed into the diffraction space where the amplitude values are replaced with those from the recorded diffraction pattern; the phases remain unchanged. This is then inverse Fourier transformed to sample space where intensity outside the support constraint is reduced by a factor of $1 - \beta$, where $\beta$ is a parameter defined at the start of the algorithm and is usually set to a value close to one. The intensity inside the support constraint remains unchanged. The new image is then Fourier transformed back to the object space for the process to continue. This cycle reaches a solution when the Fourier error stagnates. The Fourier error is a comparison of the modulus of the reconstructed field in diffraction space with the square root of the diffraction pattern.

At defined intervals within the algorithm a shrinkwrap process [64] is performed. This acts to decrease the area of the sample constraint and help the algorithm converge on the solution. This is done by convolving the current solution with a Gaussian function to blur the current sample constraint. The magnitudes are thresholded to create the
new sample constraint. It is important not to overly constrain the sample space, the
algorithm cannot reach a solution if the constraint is smaller than the actual solution.

2.8.5 Algorithm variations

There are many variations of the phase retrieval algorithms that have been developed
since the original publication. These attempt to improve the results and consistency of
results from the algorithms [65]. Largely these variations fall into two categories, a local
minimiser which has the effect of driving algorithms towards the nearest local minimum.
The other category of algorithm are global minimisers, which use feedback loops to avoid
trapping in local minima. The hybrid input output (HIO) algorithm is an example of
a global minimiser, and uses a feedback to drive the algorithm towards the lowest error
solution. In the area outside the support constraint the intensity is no longer reduced
by a factor of $1 - \beta$, instead the intensity of the previous iteration is multiplied by $\beta$ and
subtracted from the current intensity. The error reduction algorithm (ER) in contrast is
a local minimiser and works by reducing the values outside the mask to 0. This brings
the algorithm towards a solution more quickly but can often result in a local minimum
being found and not the lowest error solution.

In the work presented, elements from both algorithms are used. The HIO algorithm is
the starting point with a value of 0.9 for $\beta$. Halfway through, the value for $\beta$ is set
to 1 as is the case for the ER algorithm. An example of this technique is shown in
figure 2.13 where a simulated diffraction pattern from a test object is used. In this case
the algorithm is set up to run for 400 iterations with the shrinkwrap being performed
every 20 iterations reducing from a Gaussian FWHM of 3 to 0.5 linearly. To begin
with the output of the algorithm shows little resemblance towards the object but as
it progresses and the support constraint becomes smaller the sample begins to appear.
This simulation repeatably reconstructs the image from the diffraction pattern but the
diffraction pattern in this case is without noise and assumes a perfectly monochromatic
source. Experimentally these are not the conditions are not reproducible, HHG produces
a frequency comb making it not monochromatic, even within a single harmonic there is
a bandwidth that results in a series of frequencies. Although noise can be minimised it
is not possible to record a noise free data set, this will add further error between the
real data and the reconstructions.
Figure 2.13: The reconstructed image of the test sample after 40, 80, 120, 160, 200, 240, 280, 320, 360 and 400 iterations
2.8.6 The geometric requirements of the Algorithm

The experimental geometry for a CDI experiment is constrained by two factors, oversampling and reconstructed pixel size. The oversampling ratio is a measure of the area inside the mask to the area outside of the mask. The larger the value the greater the area outside of the mask and the easier it is therefore for the algorithm to converge. The minimum requirement is an oversampling of 2 although ideally it will be over 5 [66] to help with the noise of an experimental sample. The reconstructed pixel size is the size of each pixel in sample space. This is therefore a fundamental limit in the resolution of the final reconstructed image. The oversampling ratio is given by,

\[ O = \frac{z\lambda}{pD} \]  

(2.51)

Where \( z \) is the sample to detector distance, \( \lambda \) is the wavelength, \( p \) is the CCD pixel size and \( D \) is the one dimensional length of the sample normally using the longest axis. For a given experimental setup it can be seen that the easiest way to increase the oversampling ratio is to increase the sample to detector distance.

The image pixel size is given by,

\[ P = \frac{2z\lambda}{pN} \]  

(2.52)

Where \( N \) is the number of pixels on the detector. For a given experiment where the wavelength and detector are constant the only variable to make the image pixel size smaller is to reduce \( z \). Figure 2.14 shows the oversampling for different aperture sizes (a) and the reconstructed image pixel size (b). It can be seen that aperture sizes of greater than 5 \( \mu m \) require the sample to be placed further away than the minimum sample to detector distance (black dashed line), causing an increase in the reconstructed pixel size and hence a reduction in resolution. The contradiction between the oversampling requirement and the reconstructed pixel size leads to a limited exposure area for a sample at a particular wavelength where the balance between resolution and oversampling is achieved.
Figure 2.14: Plot of a) the oversampling ratio for a range of detector distances for 2 μm (blue), 5 μm (green), 10 μm (red) and 20 μm (turquoise) apertures and b) the reconstructed image pixel size as a function of detector distance.
Chapter 3

Apparatus

In this chapter a summary of the laser equipment required to perform the experiments in this thesis is presented including the equipment used as diagnostics for the laser. Attempting to reach the intensities required for HHG does mean that state of the art technology is required, which comes with its own unique challenges.

3.1 The Laser System

During the course of this thesis two laser systems were used. Some components from the first system were upgraded to increase the performance. The first laser system was used in chapter 4 only. The second was also used in chapter 4 and all subsequent chapters.

The peak power and the average power of a laser system are related by the pulse length and the repetition rate as shown in equation 3.1

\[ P_{\text{peak}} = \frac{P_{\text{avg}}}{f_{\text{rep}} \tau} \]  

(3.1)

where \( P_{\text{peak}} \) is the peak power, \( P_{\text{avg}} \) is the average power, \( f_{\text{rep}} \) is the repetition rate and \( \tau \) is the pulse length. As discussed previously HHG requires irradiances in excess of \( 10^{14} \text{Wcm}^{-2} \) so for a laser system of fixed repetition rate the options are to increase the average power, this is normally limited by laser components, or to reduce the pulse length, which for a given system is limited by the frequency bandwidth produced by the oscillator and the focusing geometry.
The seed lasers for these systems are continuous wave (CW) lasers. In order to produce the pulse train from the CW source an oscillator is used. An oscillator works by using a mode locking system to put the laser cavity into such an alignment that it is more energetically favourable for the light to propagate through the system in a train of short pulses rather than as a continuous wave. The repetition rate is determined by the cavity round trip time, which as the cavity length typically cannot exceed a couple of metres is of the order 100MHz.

The peak power output therefore of a typical oscillator with a 1 W average power, a pulse length of 50 fs and a 100 MHz repetition rate of will be $2 \times 10^5$ W with a pulse energy of 10 nJ. This demonstrates a vast increase in power by mode-locking but is still several orders of magnitude short of the intensity required for HHG meaning an amplification stage is required.

To increase the peak power of an ultrafast system a chirped pulse amplifier is used (CPA). A schematic of a CPA is shown in figure 3.1, the amplification occurs inside a gain medium, in the case of the lasers used for these experiments, this was a Ti:Sapphire crystal. With the short pulse lengths produced by the oscillator the peak intensity is high, therefore attempts to amplify a short pulse would result in distortion of the beam through effects such as self-focusing or self-phase modulation described in section 2.4 and in more extreme cases damage to the gain medium. To avoid this, the pulse length is stretched before amplification and then re-compressed after the amplification using a grating to disperse and recombine the different frequencies temporally. Figure 3.2 shows the effect of a CPA at different points as a function of intensity and time.
Figure 3.2: *Schematic of the pulse as it (a) enters the amplifier (b) passes through the stretcher (c) is amplified in the gain medium and (d) is re-compressed by the compressor.*

Figure 3.3: *A schematic of the 1 mJ laser system*

### 3.1.1 The 1 mJ system

The overall layout of the system is shown in figure 3.3. The system uses a Kapteyn-Murnane Laboratories (KML) Ti:Sapphire oscillator to generate the initial pulses which are then amplified by a Positive Light Spitfire chirped pulse amplifier (CPA). The oscillator uses a Spectra Physics Millenia pump and the amplifier uses a Positive Light Evolution 20 pump. The oscillator produces a 2 nJ, 30 fs pulse which is far below the power required for HHG. The pulse is therefore amplified using a CPA with a regenerative cavity, a schematic of a CPA is shown in figure 3.1. The Positive Light Spitfire is able to increase the energy of the pulse from 2 nJ to 1 mJ with a pulse length of 35 fs, although the repetition rate drops from 96MHz to 1KHz.
3.1.2 The 3 mJ system

The upgrade to the laser system meant that the KML oscillator was replaced by a Spectra Physics Tsunami Ti:Sapphire oscillator producing 5.7 nJ, 35 fs pulse at a 79 MHz repetition rate. This was pumped by the same Millennia diode laser. The Positive Light Spitfire was replaced by a Spectra Physics Spitfire Pro CPA. The footprint of the new amplifier was significantly smaller than that of the previous version. The final output of the amplifier is 3 mJ, 40 fs at a 1 KHz repetition rate.

3.2 Laser diagnostics

There are four parameters that are routinely measured in order to monitor the operational state of an ultrafast laser system, these are the power, the spectrum, the pulse length and the $M^2$.

The power is routinely measured between the oscillator and the amplifier and after the amplifier using a thermal power meter. The spectrum is measured using an Ocean Optics HR2000 spectrometer after the oscillator. This allows the central frequency and bandwidth of the pulse to be kept consistent. The same spectrometer can also be used to measure the spectrum after the amplifier when necessary.

Measurement of a lasers pulse length and hence the pulse energy is a key process to monitoring the operational state. Ultrashort pulses are measured against themselves as there is no shorter event to be able to measure them against. The original method for
this was using an autocorrelator, which split the beam in two, delayed one part with respect to the other and then overlapped them in a non-linear crystal. The intensity of the SHG signal is then recorded onto a camera. This provides an estimate for the pulse width but it does not allow for determination of pulse structure. For a more comprehensive measurement frequency resolved optical gating (FROG) [67] [68] is used. As shown in figure 3.4, FROG replaces the camera with a spectrometer to frequency resolve the delay between the pulses. This allows a phase retrieval algorithm to be used to reproduce the spectral phase of the pulse. It also allows for the measurement of more complicated pulses.

The other parameter of interest for the laser is the $M^2$. The $M^2$ is a measure of the quality of the laser beam relative to a TEM$_{00}$ Gaussian beam [53]. This is important as it has an effect on the propagation and focusing of the beam which for coupling into a capillary is critical. There were currently no $M^2$ meters commercially available for high intensity ultrafast systems so as a group we have designed and built our own. It works by focusing a beam onto a pair of partial reflectors to create a profile of the beam at various points along its path on a charge coupled device (CCD). The width of these points can be measured, and by knowing the separation of the reflectors and their angle a fit of an $M^2$ value can be accurately made. An example of the software is shown in figure 3.5.
Figure 3.5: A screenshot from the $M^2$ meter showing the beam profile through the focus and the measured $M^2$
Chapter 4

Pulse propagation in hollow capillary waveguides

The first two results chapters in this thesis discuss source development for a hollow core capillary based HHG system. The aim is to produce a model that can accurately describe the propagation of the driving laser pulse through the capillary. This would then be used in conjunction with the quantum mechanical model of HHG described in section 2.1.2 to predict the XUV output and optimise the system.

As described in chapter 1, HHG is typically performed in three different geometries. In both a gas jet and a gas cell the interaction region is controlled by the focusing of the driving laser field. In a capillary this interaction region can be extended by waveguiding. This chapter will describe the process of mounting a capillary, the methods used to effectively couple into a capillary and a comparison of experimental data to a model developed with Dr E. Rogers, describing the propagation of the driving laser field within a capillary.

4.1 Mounting and coupling into a capillary

The capillary used in the experiments described in the next two chapters was fabricated from sections of 30 cm long capillary with an outer diameter of 1.6 mm and an inner diameter of 150 µm. It was cleaved into 7 cm lengths using a standard fibre cleaving technique. A small score is made in the surface of the capillary at the desired length, a
Figure 4.1: A schematic of the capillary used for HHG showing the position of the two gas inlet holes

Figure 4.2: Photographs showing examples of (a) a good cleave and (b) a poor cleave

drop of de-ionised water is placed on the score to lubricate the process. A small amount of pressure is applied either side of the score in the opposite direction and the capillary cleaves. A good cleave results in a smooth end surface and no stress fractures by the core. Examples of both a good cleave and a bad cleave are shown in figure 4.2. The fracture in figure 4.2b has an effect on the ability to couple into the capillary described in the next section. On some occasions the capillary would cleave leaving a hackle, a rough surface on the end of the capillary. The target gas for HHG flowed into the capillary through two inlet holes (150 µm) drilled through the side walls of the capillary. Figure 4.1 shows a schematic of the basic capillary design.

The inlet holes were initially milled using a supersonic drill but the results were unsuccessful as drilling onto a curved glass surface often led to the drill bit slipping or snapping as it began to drill. As a result the capillaries were milled using a New Wave micro machining setup with a 50x objective attached to Coherent Legend-F laser system (800 nm, 150 fs, 1 kHz, 2 mJ). The sample was positioned using three Aerotech gas-bearing stages. This work was undertaken at the Femtosecond Applications of Science and Technology lab, University of Southampton.
4.1.1 Mounting the capillary

Having fabricated the capillary a solution was required to correctly position it within the vacuum system, co-linear with the laser and connected to a regulated gas supply.

A custom manufactured glass T-piece was designed to hold and seal the capillary into the vacuum system, a schematic of which is shown in figure 4.3. The T-piece allowed the target gas into the centre of the capillary via the inlets whilst isolating the gas supply from the vacuum system at either end of the capillary. To achieve this the T-piece inner diameter was reduced outside of the gas inlets on either end to 0.2 mm larger than the outer diameter of the capillary. At these two points a UV epoxy was used to set the capillary position and provide the vacuum seal between the gas supply and the vacuum system.

The T-Piece was subsequently fixed into the experimental setup via two o-ring sealed swagelock connectors, each connected to a two axis stage to allow accurate manipulation of the capillary position relative to the incoming laser beam.

4.1.2 Coupling into the capillary

In section 2.6 it was shown that the Bessel modes the incident beam coupled into depend on the ratio of the focal spot to the inner radius of the capillary. For optimum coupling into the lowest order EH_{1m} mode, the spotsize, \( w_0 = 0.64a \) where \( a \) is the radius of the capillary. In this case the optimum value for coupling into the capillary is 48 \( \mu m \). A 70 cm plano-convex lens was used and gave a focus of 43.5 \( \mu m \) or 0.58a. The result of this was a greater proportion of the beam will be coupled into higher order modes.
Initially the coupling was carried out at vacuum to prevent laser gas interactions and the laser intensity was attenuated using reflective neutral density filters mounted in a wheel. The laser was then guided into the capillary, which was manipulated at the entrance and exit in the planes orthogonal to the propagation direction of the laser as well as translation of the lens along the propagation axis to maximise the measured transmission. Once the transmission was maximised the attenuation was reduced to allow a greater laser intensity into the capillary. This process was complicated by the small change in position of the beam with the rotation of the filter wheel. This positional movement resulted in the more intense laser pulse being focused onto the front face of the capillary. The intensity of the pulse at this point is above the damage threshold for silica resulting in damage to the capillary face shown in figure 4.4. It is difficult to quantify exactly what this means for the model but experimentally the transmission through the capillary was reduced to 50 % for a well used capillary where as for a new capillary transmission is in excess of 80 %.

4.2 Propagation along the capillary

Models of any system require validation by experiment. In this chapter the experiment seeks to measure the driving pulse intensity along the capillary by measuring the ionisation fraction as a function of length along the capillary. When the driving pulse
interacts with a target gas, in this case argon, a series of absorptions occur resulting in the production of excited state neutral argon and excited state argon ions. These excited states will relax to their ground states via radiative transitions. An example of the total visible fluorescence can be seen in figure 4.5, scattering from the front face of the capillary and the gas inlet holes can also be seen.

Each of these different transitions radiate at a specific wavelength depending upon the energy gap between the energy levels. Selecting a specific transition will allow measurement of a relative amount of ionisation along the length of the capillary. This experimental data can then be compared to a predicted ionisation fraction along the length of the capillary derived from a model described later in this chapter.

4.2.1 Experimental Method

The experimental setup is shown in figure 4.6. A 40 fs, 790 nm laser pulse at 1 kHz repetition rate was coupled into the 70 mm long hollow capillary using a 0.7 m plano-convex lens. Two data sets were recorded using two identical capillaries but using two different laser systems setup to produce the same output due to an upgrade of the oscillator and amplifier. The input pulse energy was controlled between 378–840 \( \mu \text{J} \) (Intensities between 2.8–6.3 \( \text{Wcm}^{-2} \)) by inserting reflective neutral density filters into
the beam. As described in the previous section a pair of 150 µm holes were drilled 20 mm from either end of the 70 mm long capillary with a bore radius of 75 µm. These were used to leak argon gas at pressures up to 200 mbar into the system from a Tescom ER3000 pressure regulated supply. The ends of the capillary are attached to a vacuum system which maintained a background pressure < 10⁻⁵ mbar.

Orthogonal to the capillary propagation length the visible fluorescence from the ionised argon was measured using an Acton 300i spectrometer. The spectrometer was subsequently replaced with an imaging lens and a CCD camera (Princeton Instruments Pixis 400) to photograph the fluorescence from the capillary. A 600 nm low pass interference filter with optical density 3 at the laser wavelength (CVI Technical Optics) was used to attenuate the laser, in combination with a 420 nm or 488 nm narrow band pass filter (10 nm FWHM) to select intense lines corresponding to emission from neutral argon or singly ionised argon respectively [69].

A grazing incidence spectrometer was used to record the high harmonic spectra, consisting of a 0.5 mm wide vertical slit followed by a 300 lines / mm grating at glancing incidence. A micro-channel plate (MCP) detector collected the XUV photons and converted them to electrons which collide with a phosphor screen. The screen is imaged onto a Princeton Instruments Pixis 400 CCD camera using a 35 mm camera lens.
4.3 Results for initial experimental setup

The data shown in this section relates to the original laser system outlined in section 3.1.1. The laser was coupled into the capillary using the method described in section 4.1.2 with a calibrated efficiency of 50%, the input pulse was 840 µJ. It was possible by eye to see sections of the capillary were glowing a blue colour more intensely than others, suggesting a non uniform distribution of ionisation.

![Graph showing emission spectrum from the capillary](image)

**Figure 4.7**: Experimentally measured emission spectrum from the capillary filled with 200 mbar of argon gas with an incident 40 fs 840 µJ laser pulse. The solid lines represent the area of the spectrum transmitted by the 420 nm bandpass filter and the dashed lines represent the 488 nm bandpass filter.

The spectrometer was set up to capture the fluorescence from the capillary orthogonal to the propagation direction of the capillary. The spectrum obtained in figure 4.7 shows two intense peaks at ~ 420 nm and two intense peaks at ~ 488 nm. The peaks in the 420 nm region correspond to excited state transitions of Ar I and the peaks in the 488 nm region are excited state transitions of Ar II [69].

Two bandpass filters were selected with a spectral bandwidth appropriate to the emission from these transitions allowing selection of either fluorescence from the presence of excited argon neutral atoms or excited argon ions.

The spectrometer was replaced with an imaging lens and a CCD detector with the filters mounted onto the front of the detector. The lens imaged the capillary onto the CCD
to enable single shot measurement of fluorescence for the whole capillary. For both the 420 nm and the 488 nm filters the fluorescence was recorded as a function of the gas pressure within the capillary, keeping the input laser power constant. The emission from both excited neutral argon, figure 4.8a and excited argon ions, figure 4.8b is shown for the low coupling case. In order to produce a line plot of intensity versus length, the emission is summed vertically, and plotted as a function of position relative to the entrance of the capillary.

The singly ionised and neutral states of argon are seen to have very different distributions within the capillary. The intensity from the ion emission is greater at the start of the capillary, and reduces along the length of the capillary. The emission from excited atoms follows the pressure distribution more closely. This indicates that the excitation mechanism for ions has a higher order of nonlinearity compared to the excitation of atoms, as might be expected if it is part of the ionisation process. A strong mode beating pattern as shown in figure 2.8, can also be observed for both species, predominantly between the EH\textsubscript{11} and EH\textsubscript{12} modes (the theoretical positions of the peak intensities of the beating pattern are indicated by the vertical lines in the plot as described in section 2.6). Higher order mode beats can also be seen, resulting from the presence of small fractions of energy within higher order modes.

4.3.1 Constructing the model

Although the observations of a beating pattern along the length of the capillary were interesting, it was important to create a model to understand why it was happening and if it would be possible to use the observed phenomenon to improve the XUV yield. In collaboration with Dr E. Rogers a model was developed based upon the mode propagation equations described in section 2.6. The simulation proceeds by defining a Gaussian input beam and calculating the coupling efficiency $A_m$ for each capillary mode using equation (2.38). The electric field at any point in the capillary may then be calculated from the weighted sum of the appropriate contribution of each of the $n$ possible EH\textsubscript{1m} waveguide modes

$$E(r, z) = \sum_{m=1}^{n} A_m E_{1m}(r, z). \quad (4.1)$$

Each mode is independently propagated a short distance $\Delta z$ along the longitudinal axis of the capillary, using equation (2.34) to account for the phase advance and attenuation.
Figure 4.8: Emission from (a) neutral and (b) singly ionised argon species as a function of position along the propagation axis of the capillary, where zero indicates the capillary entrance. The vertical lines indicate theoretical positions of beating pattern between the two lowest order $EH_{11}$ and $EH_{12}$ modes.
losses of each of the capillary modes. The new spatial profile is again calculated according to equation (4.1).

As the pulse propagates, it ionises the gas in the capillary. The ionisation rate was calculated from Keldysh theory described in chapter 2.4.1 [51], integrating the rate over time to find the fraction of atoms \( \eta \) ionised. The energy taken to ionise the gas may then be calculated according to

\[
E_i(r, z) = N(r, z)I_p\eta
\]

where \( I_p \) is the ionisation potential of the atom and \( N \) is the number of atoms in the volume element

\[
N(r, z) = 2\pi r \Delta r \Delta z P N_{atm}
\]

where \( r \) is the radius of the volume element, \( \Delta r \) is a small step across the capillary’s radius, \( \Delta z \) is a small step along the capillary’s longitudinal axis, \( P \) is the pressure in bar and \( N_{atm} \) is the number density at one bar. The gas pressure profile in the capillary was calculated using the computational fluid-dynamics package Ansys CFX. The capillary is 70 mm long with two 150 \( \mu \)m holes placed 20 mm from either end of the capillary. The model shows the pressure to be constant between the holes and after a 20% pressure drop across each hole the pressure decreases linearly towards the ends of the capillary as shown in figure 4.9.

\[\text{Figure 4.9: Modelling of the gas flow in a capillary using fluid dynamics for an input pressure of 200 mbar, 70 mbar and 10 mbar.}\]
4.3.2 Comparison of the model and experimental data

The model was initially tested by comparing the ionisation induced energy loss as a function of pressure within the capillary. Figure 4.10 shows a comparison of the theoretical calculations of ionisation induced losses calculated using the model in section 4.3.1 and the experimental measurements of the output energy from the capillary as a function of argon gas pressure. The calculations and measurements were performed for three different input laser energies. The power was measured by placing a mirror into the beam to reflect it out of the vacuum system through a fused silica window onto a power meter. As the exact coupling conditions were not measured, the input energy for the model was calculated to fit the data for a pressure of 0 mbar. The curves show increasing loss due to ionisation as a function of pressure for the two higher input energies. This is the expected behaviour as the density of atoms increase so does the number of ions produced and hence a greater loss to the transmitted driving pulse.

The lowest input energy shows no loss due to ionisation as the energy in this case is not high enough to cause significant ionisation. The overall fit to the data here suggests that the model is accurately predicting both the propagation of the driving pulse and the ionisation at each point.

Although the data shown here is encouraging, a measurement of the relative ionisation fraction as a function of capillary length for a range of pressures is a much more rigorous test of the model.

Figures 4.11a, c and e show the pressure variation of the excited ion fluorescence distribution as a function of length for three different input energies. Figure 4.11b, d and f show the theoretical prediction of the total number of atoms ionised along the capillary, which is proportional to emission, plotted as a function of argon pressure and axial position along the capillary. As gas pressure is increased, the effective nonlinearity of the medium is increased. Little variation in the position of the beat pattern is observed as a function of pressure, indicating that nonlinear mode-mixing into higher-order modes is not a significant effect, with most of the power remaining in $EH_{11}$ and $EH_{12}$. There is a change in the amount of observed fluorescence as the pressure increases due to the increased number of atoms interacting with the laser and hence an increase in the number of ions fluorescing.
given by,
\[ N(r, z) = 2 \pi r \frac{\Delta r}{\Delta z} N_{\text{atm}} \],
(6.10)

where \( r \) is the radius of the capillary, \( \Delta r \) is a small step across the capillary’s radius, \( \Delta z \) is a small step along the capillary’s longitudinal axis, \( P \) is the pressure in bar and \( N_{\text{atm}} \) is the number density at one bar. The pressure is calculated from a fit to the fluid-dynamics model data shown in chapter 4. This model can be used to describe the ionization induced energy loss as a function of pressure in the capillary.

Figure 6.8 shows the theoretical calculation compared to experimental data.

Figure 6.9 shows the theoretical prediction of the total number of atoms ionized which is proportional to emission, plotted as a function of argon pressure.

The theoretical model uses the same laser parameters as the experiment and allows \( \text{EH}_{1m} \) modes where \( m \) is between 1 and 30. The dominant \( \text{EH}_{11} - \text{EH}_{12} \) mode beating is clearly visible.

The theoretical model also shows a similar overall reduction in ion fluorescence intensity along the capillary axis. This suggests that the ionisation induced losses have reduced the driving pulse intensity inside the capillary significantly before the region where absorbance length for the high harmonics is less than the distance to the end of the capillary. One obvious difference between the experiment and theory is that the expected intensity peak at 23mm from the front of the capillary is observed experimentally at 18mm. This offset could be due to the focusing conditions into the capillary. If the waist of the beam is not placed at the entrance to the capillary, a curved wavefront is coupled to the capillary which would alter the phases of the coupled capillary modes at the input and may shift the position of the first beat. There is also a higher fluorescence at the entrance of the capillary in the experimental data, this may be a limitation of the
CFD modelling which assumes a zero pressure at either end of the capillary. If the gas pressure is higher then the expected fluorescence will be higher.

4.4 Results for final experimental setup

A subsequent dataset was recorded with the second laser system described in section 3.1.2. During the installation of the new system the position of the beam line was moved and the capillary used in the previous section was damaged. A new capillary was fabricated to the same specification as the previous one. Once mounted into the beamline the experiment setup was rebuilt to repeat the measurement of the argon ion fluorescence along the length of the capillary.

A change was also made to the coupling process. Instead of using a series of neutral density filters to control the power, the power of the pump laser to the amplifier was
adjusted. There are two advantages to this method, the position of the beam does not move spatially between the different powers used and the control of the power is much finer.

![Image of plasma distribution through a 488nm filter for a series of pressures starting at 20 mbar through to 160 mbar in steps of 20 mbar. The individual plots can be found in appendix A.](image)

**Figure 4.12:** The plasma distribution through a 488nm filter for a series of pressures starting at 20 mbar through to 160 mbar in steps of 20 mbar. The individual plots can be found in appendix A

The result of these changes was a transmission of 80% after optimising the coupling conditions. There were several notable differences arising from the increase in intensity all along the capillary. Figure 4.12 shows the plasma distribution along the length of the capillary for different pressures. At low pressure, where the atomic density is small and thus the nonlinear change in refractive index on ionisation is small, the beat pattern is similar to that seen in the low coupling efficiency case, with EH$_{11}$–EH$_{12}$ beats separated by 23 mm dominating. As the gas pressure is increased, the beat pattern changes significantly, indicating that nonlinear mode mixing is becoming significant, and a much more complex fluorescence appears above $\sim$ 40 mbar. The presence of large peaks near to the capillary exit (beyond 50 mm) at high pressures suggest that structure of the driving pulse has become much more complex than the model accounts for. In the output of the driving pulse from the capillary yellow and green frequencies were observed which will be discussed in the next chapter.

Simultaneously to recording the plasma distribution, the XUV spectrum was also recorded. Figure 4.13 shows these corresponding XUV spectra also as a function of pressure.
For low pressures, evenly-spaced harmonics are observed extending down to $\sim 27$ nm. As the gas pressure increases beyond $\sim 40$ mbar, the XUV emission spectrum changes rapidly, as non-linear propagation changes the driving laser spectrum. This non-linear change of driving laser spectrum and the nonlinear mode coupling arise from the same optical nonlinearity, which only becomes important when the gas pressure is relatively high and the coupled intensity through the capillary is high enough for significant ionisation to occur. It is also interesting to note the presence of what would appear to be even harmonics within the spectrum. This may be a result of the blue shifted part of the laser interacting with the fundamental and generating different wavelengths. Some predictions of second harmonic interaction with a fundamental laser pulse have suggested even harmonics may be observed [70].

### 4.5 Conclusions and further work

The use of spectrally resolved Ar/Ar$^+$ imaging has given a new insight into propagation inside a hollow capillary. The observed images shown in figures 4.8 and 4.11 are a useful diagnostic for low coupling efficiency conditions, showing that the power inside the capillary is predominantly contained in the lower order modes. This is modelled well by a simple linear model using mode propagation and ionisation theory.
At higher coupling efficiencies and hence power inside the capillary the imaging reveals evidence of a much more complicated system. Non-linear mode mixing has been suggested as a cause for some of the results but there is no experimental evidence for this. Non-linear mode mixing is not currently taken into account by the existing model and so limits the information that can be interpreted from the model. The influence of the spectral broadening can be seen on the XUV generation leading extra peaks in the spectrum and eventually broadening of the spectrum.

A collaboration was started to investigate the use of a model based on a multimode nonlinear Schrödinger equation. This model would take into account effects such as non-linear mode mixing giving a more complete picture. The results in the next chapter describe the experiments carried out to validate the model, including looking for experimental evidence of non-linear mode mixing.
Chapter 5

Testing of a non-linear Schrödinger equation model

In the previous chapter it was shown that coupling into the capillary was a strong factor in determining how the driving pulse propagates along the length. The relationship between the driving pulse and the XUV output is sensitive to the changes in these propagation conditions as the process is highly non-linear. Therefore before understanding how the XUV output of the capillary can be optimised a detailed model is required. Previous theoretical studies of capillary-based HHG by Christov et al. [71] have used numerical solutions of the 3-dimensional version of the scalar wave equation. This work focused on understanding attosecond pulse generation [72] and quasi phasematching within waveguides [73]. Experimentally measurable predictions of these theories have typically centred on the temporal profile of the driving pulse and the harmonics generated.

As described in chapter 2.7, a collaboration was developed with Dr P. Horak to construct a multimode Non-Linear Schrödinger Equation model. This model was based on Dr. Horak’s previous work involving non-linear propagation in multi-mode fibres. This work was extended to include a plasma based non-linear coupling term and an ionisation induced loss term.

Improvements in coupling efficiency described at the end of the last chapter resulted in some new observations. The appearance of blue shifting deep into the visible spectrum was noted, in addition the output of the capillary was observed to be composed of a
symmetric ring structure as shown in figure 5.1. The photo is taken through a high pass filter to show the output with the dominant infra-red removed. The ring structure appears to contain different spectral distributions as a function of the radius. This was tested by placing a fibre coupled spectrometer at the positions marked on the photograph. Figure 5.2 shows the spectrum measured at the central maxima (P1, blue line), the first maxima (P2, green line) and the second maxima (P3, red line), normalised to its maximum value. Whilst the overall shape of the spectra is similar for all three, the observed blue shifting is higher at the higher angle.

Figure 5.3 shows the propagation of the modes predicted at the end of the capillary into the far field. There is also an obvious radial dependance shown here with higher order modes presenting further away from the central maxima. Although each mode is not isolated at any point radially the variation provides opportunity to scrutinise the model output more rigourously.

The comparisons of this model to experimental data will be based on three different criteria. Firstly a comparison of the total spectral output to check that the degree of non-linearity produced by the model matches the non-linearity seen experimentally. The second comparison will be a spatio-spectral measurement, this will provide a more rigorous inspection of the model as it will be related to the non-linearity of each mode.
3.2.3 Spatio-spectral output

The pump pulse propagates within the capillary as a set of Bessel modes. The model assumes circularly symmetric modes $EH_{11}, EH_{12}, \ldots, EH_{120}$. When the pulse reaches the end of the capillary each of these modes propagates out. In the far field the main peak of each of these modes is spatially separated, as shown in Figure 3.13. This is a plot of the first five $EH_{1m}$ modes propagated into the far field from the capillary exit. They are shown scaled linearly to the normalised $EH_{11}$ mode. The $EH_{11}$ mode has a main peak on-axis, with the main peak of each subsequent mode positioned at increasingly greater radii. This means that in the far field it should be possible to see experimental proof of the contribution from the blue shifted higher order modes.

The modal contribution was examined by taking a spatially resolved spectrum for a pump power of 805 mW with a gas pressure of 100 mbar. The spatio-spectral map was built up by measuring the spectrum every 500 µm from on axis to a radius of 24 mm at a distance of 80 cm from the capillary exit. The spectral intensity is greatest in the first 10 mm so to produce a map out to 24 mm neutral density filters were put in place and removed at greater radial distances. The data was scaled accordingly and combined in Matlab to produce Figure 3.14a. This is a log plot of the spatio-spectral output of the capillary from on axis to 24 mm. The x-axis shows the radial distance while the y-axis shows the wavelength. Figure 3.14b is a sum over the experimentally measured spectrum and shows the summed intensity of all modes over the same radial distance. As a comparison a sum of the first five modes theoretically propagated to the spectrometer are shown in Figure 3.14c. The five main peaks from each mode can be

Figure 5.2: Normalised spectral output of the capillary for three radial positions shown in figure 5.1, P1 corresponds to the central maxima, P2 the maxima in the first ring and P3 the maxima in the second ring.

Figure 5.3: The far-field distributions of the five lowest order $EH_{1m}$ modes, showing an angular dependance of the modal propagation.
within the capillary. Finally a comparison of the relative ionisation fraction as a function of length of the capillary in the same manner as the previous chapter.

5.1 Experimental Method

The experimental setup is shown in figure 5.4. A 40 fs laser pulse centred at 790 nm, with a 1 kHz repetition rate, was coupled into a 70 mm long hollow capillary with internal radius 75 \( \mu m \) using a 0.5 m plano-convex lens. A further adjustment was made to the coupling process for the experiments described in this chapter. In order to match the spotsize to the radius of the capillary ratio \((w_0 = 0.64a)\) as accurately as possible the telescope inside the amplifier was adjusted. This gives optimum coupling into the lowest order, \(EH_{11}\), mode [56]. As a result the calibrated coupling efficiency of 91% could be achieved. The input pulse energy was varied between 497 and 805 \( \mu J \) by adjusting the power of the pump beam into the regenerative amplifier. The central region of the capillary was filled with argon gas via a pair of 150 m holes drilled 20 mm from either end of the 70 mm long capillary. Argon gas at pressures up to 200 mbar could be introduced into the system from a Tescom ER3000 pressure-regulated supply. The ends of the capillary are attached to a vacuum system which maintained a background pressure < \(10^{-5}\) mbar. The overall gas pressure profile through the capillary was calculated using computational fluid dynamics shown in figure 4.9.

![Figure 5.4: Schematic of the experimental setup](image)

The capillary output is a combination of the pump laser pulse and the generated XUV. The driving laser pulse was reflected out of the vacuum chamber into a fibre-coupled
The 125 µm diameter fibre end was placed at the centre of the beam and translated radially outwards across the whole beam profile. The plasma emission from the side of the capillary was imaged onto a CCD camera (Princeton Instruments Pixis 400) placed perpendicular to the axis of the capillary. A 600 nm low pass interference filter with optical density 3 at the laser wavelength (CVI Technical Optics) was used to attenuate the laser, in combination with a 488 nm narrow band pass filter (10 nm FWHM) to select intense lines corresponding to emission of singly ionised argon [69].

5.2 Results

Having decided on the methods that were to be used to compare the model and the data a series of experiments were carried out.

5.2.1 Spectral output of the capillary

A ground glass diffuser was placed directly front of the fibre to capture the complete output spectrum from the capillary at three different driving laser pulse powers. The pressure inside the capillary was maintained at 100 mbar. The predicted spectrum from the model described in section 2.7 and measured total output spectra for three different input powers are shown in figure 5.5. The theoretical (a) and experimental (b) spectra show excellent agreement with not only the degree of blue shifting observed, but also the complex spectral shapes that occur at higher powers.

Figure 5.5(a) shows that at low powers the EH$_{11}$ mode dominates the spectrum as nonlinear mode coupling is small. As the power is increased, more energy is coupled into the EH$_{12}$ and the spectrum broadens a small amount. There is also the beginning of a small shoulder at 800 nm in both the theory and experiment. At the highest power the blue shifting has extended to significantly below 700 nm. The model suggests that the degree of blue shifting is significantly enhanced by the contribution from higher order modes in the far field. The plasma generation at the leading edge of the pulse results in blue shifting caused by the plasma induced refractive index change described in chapter 2.4. This effect is also the cause of the nonlinear mode coupling, which is greater at the trailing edge of the pulse due to plasma defocusing. Experimentally, the
modal distribution cannot be directly determined from a summed spectrum of the kind shown in here, although the agreement between theory and experiment allow us to infer that the modal distribution is correct.

5.2.2 Spatio-spectral output of the capillary

In order to test whether the model correctly predicts the distribution of intensity in high order modes, one can make use of the fact that as the modes propagate into the far field their spatial divergence is strongly dependent on their order. This is shown earlier in figure 5.3, with higher order modes showing significantly greater angular spread. This means that the radial distribution of the spectral intensities has contributions from different modes at different radii (although the modes are not completely separated). Thus the spatio-spectral intensity distribution in the far field is very sensitive to any variation in the intensities of individual modes. Therefore its measurement can provide a sensitive test of the ability of the numerical model to correctly predict the modal distribution, and thus the propagation of the driving laser pulse.
The spatio-spectral required the diffuser to be removed from the fibre coupled spectrometer. The tip of the spectrometer was mounted onto a translation stage in the centre of the output pattern. The fibre was then translated across the pattern in steps of 500 µm, 80 cm from the exit of the capillary. This process had to be carried out quickly to preserve the integrity of the data collection. This is due to the high sensitivity of the position of the capillary to the incoming laser beam. The use of beam tubing, improved optic mounts and removal of heat sources from close to the optical bench have gone towards improving the stability of the incoming beam. Although a lot of work has gone into minimising factors that could introduce instabilities into the beam position it is still not reliable over the course of tens of minutes. This is however a marked improvement on early experiments when the position of the argon ion fluorescence could be seen to move back and forth along the capillary constantly. Further work is still required to improve the laser pointing stability, one improvement would be to reduce the beam path length as currently this is longer than it needs to be. This has implications for the future use of capillaries as a source for imaging experiments where integration times could be in the range of hours.

Figure 5.6 shows the calculated (a) and measured (b) spectral intensity distribution as a function of radial distance from the beam axis. The calculated distribution is produced by taking a scaled Fourier transform of the field at the end of the capillary having used the model from section 2.7 to propagate along the capillary. The normalised theoretical intensity profiles of the EH$_{11}$ and EH$_{12}$ modes are shown for comparison, the unnormalised profiles are plotted in figure 5.3 showing that the intensity in the EH$_{11}$ mode is much greater than the other modes. The data from both theoretical and experimental distributions show that the intensity variations of the first two modes are large. The EH$_{11}$ mode has significant contributions at around 790 and 810 nm, and the blue-shifted components at about 740 nm show the same radial profile as the EH$_{12}$ mode, with an on-axis peak, and another $\sim$8mm from the axis. This distribution is clear in the theoretically modelled intensity distribution. In the measured distributions, the general features of the pattern are repeated. The EH$_{11}$ mode appears at 780 and 810 nm, and the blue-shifted peak at 760nm has the radially narrower distribution of the EH$_{12}$ mode. However, several differences are clear. The first is that more extensive off-axis blue shifting is seen in the experimental data, implying greater broadening in the higher order modes than was predicted. Secondly, an extra peak can be seen close
Figure 5.6: Predicted (a) and experimental (b) spectral intensity plots in the lambda-r-plane. The solid and dashed white lines show the positions of the far-field profiles of the EH$_{11}$ and EH$_{12}$ modes respectively. The full mode profiles are shown in figure 5.3.

to the centre of the beam at 800 nm in the experimental data. The shape of the EH$_{11}$ peaks around 790-810 nm look similar to the spectral distortion caused by self phase modulation, which produces first a splitting, and then a central peak rising between the split peaks as the nonlinear phase shift increases. Both of these differences suggest that the nonlinear shifting of the spectrum is slightly stronger in the experiment than predicted by theory. This is however the first evidence of a modal dependance for spectral shape.

5.2.3 Argon ion distribution along the capillary

While spectral measurements at the capillary exit are a good test of the end point of the model, measurement of the argon ion fluorescence along the length of the capillary provides a test of the model along the whole propagation length. Experimentally, the fluorescence, produced by excited argon ions created by the driving laser pulse, is filtered and imaged from the side. The integrated fluorescence from each point along the capillary length can be compared directly to the ionisation levels predicted by the propagation model. The integrated argon ion fluorescence and the theoretical integrated ionisation are compared in figure 5.7. The measured 488nm argon ion fluorescence is
proportional to the calculated ionisation level within the capillary. To aid comparison, the beat positions for the EH$_{11}$ and EH$_{12}$ modes calculated for linear propagation are shown as vertical dashed lines. The initial increase in ionisation at the capillary entrance is observed in both theory and experiment and the first two major peaks appear at approximately the same positions within the capillary. The smaller structures within these major peaks do not correlate. It is important to note that neither the theoretical or experimental peak positions coincide with the peak positions calculated using simple linear mode beating. The final major peak predicted by the numerical model is observed at the same point in the capillary but is significantly smaller in length and size. The discrepancy may be due to losses at the gas inlets within the capillary wall. These have been observed as increased scattering during experiments and are not included within the numerical model.

**5.3 Conclusion and further work**

In this chapter a new method for numerically modelling nonlinear propagation within a capillary used for HHG has been shown to agree well with three different experimental
measurements. The spectral output of the capillary showed a strong correlation with the spectrum predicted by the model. The model also predicts a modal variation of the spectral shape, and the use of a novel spatio-spectral measurement technique allowed detailed comparison of these individual modal contributions, because of the differences in the far-field mode patterns, validating the model and the coupling terms chosen. Understanding the modal distribution will allow modelling of XUV phase matching in the presence of nonlinear mode mixing, which is important for XUV generation at high intensities.

The comparison of the argon ion fluorescence showed that the propagation along the capillary is far more complicated than can be explained by the simple model from the previous chapter but is reproduced reasonably well by this model. There is still a case for improving the model as there are still some discrepancies with the experimental data. The first step would be to measure the beam properties such as the pulse length just before the capillary. Currently there is a 5 m beam path from the diagnostics to the capillary entrance. Using the model to predict an optimised capillary length and bore size from a parameter scan using high intensity at the exit of the capillary as the measure and then experimental comparison of the XUV flux would be a further rigorous test of the model.

The model also predicts changes to the time structure of the pulse through the capillary which are not covered in this thesis. Attempting to measure these and using them as an extra dimension in the parameter scan would be both challenging and interesting.

The final piece of further work would be to combine the output of this model with the TDSE model described in section 2.1.2 to give a complete model from free space gaussian beam to useable XUV beam. This would represent a leap forward in understanding of the process of HHG.
Chapter 6

Characterisation of XUV beam for microscopy

Constructing an XUV microscope setup requires manipulation of the XUV beam after the generation process. Typically this involves focusing the beam onto a sample and detecting the resulting diffraction pattern. There are several aspects of the focusing which will affect the systems usability in the future, these include the position of the focus along the beam axis to maximise intensity upon the sample, size of the focus to determine the size of samples that can be used and the spectrum after the focusing optic in order to understand the limits of the reconstructions used.

The work carried out in this chapter moves away from the source development work of the previous two chapters to characterise the effects described above of our chosen focusing regime on the beam. Methods for measuring the focal position and spotsize will be discussed along with a characterisation of the spectral profile of the beam after the focusing optic.

6.1 Change of experimental geometry

The previous two chapters have focused on the development of capillary based HHG, more specifically the propagation of the driving laser pulse through the capillary. Whilst this showed progress in understanding the propagation there were difficulties in aligning the capillary and keeping the output stable, which currently restricts its use to source
research. In selecting an imaging source it is essential that it has a stable output with respect to the input beam and the alignment is not a limiting factor of the experiment. Previous experience from members of the group demonstrated that the photon flux from a gas cell \((1.5 \times 10^7 \text{ photons/second})\) was similar to that of a capillary \((3.2 \times 10^7 \text{ photons/second})\)[74]. In the gas cell the stability is dependent only upon the stability of the laser as the input and output apertures are large in comparison to the beam diameter. This allows a small tolerance to laser pointing stability. In the capillary there is an additional factor in the stability of the capillary which is held in place by o-ring seals. The capillary geometry is also very intolerant to any laser instability. As a consequence the experiments in the following two chapters were conducted using a gas cell generation geometry.

The gas cell was constructed by flattening a 6 mm copper pipe to leave an internal width of 800\(\mu\)m with 1.2mm walls as shown in figure 6.1. This thickness of the 6 mm pipe was reduced because the absorption length at 27 nm, the peak reflectivity of the focussing
optic, is 4.6 mm for 70 mbar of argon. The pipe was sealed at one end and two 500 µm holes were drilled through the flattened region to allow the laser to propagate through the target gas. The other end of the gas cell was connected to a regulated gas supply and mounted in a small vacuum chamber. A previous attempt had been made using a larger hole for the laser and wrapping the gas cell with polytetrafluoroethylene (PTFE) tape to seal it. The laser then created an aperture by drilling through tape, however the spatial profile of the spot was notably less circular. Figure 6.2 shows the XUV beam from the 500 µm drilled holes (a) and the PTFE wrapped gas cell with laser drilled holes (b) measured using an XUV CCD camera. The decrease in beam quality shown in the PTFE profile would have an effect on the focusing ability of the experiment leading to enlarged focal spots.

6.2 Focusing optics

The output from HHG sources have limited fluxes spread across a series of frequencies. Use of these frequencies for imaging experiments where the samples are on a micron to nanometer scale requires a focusing optic to concentrate the flux. There are three main types of focusing used in the XUV and soft x-ray regions, these are zone plates [75], multi-layer mirrors [76] and parabolic tapers [74] shown in figure 6.3.

Zone plates are a focusing optic that uses Fresnel diffraction to create a build up of intensity at a given point. They are designed for a specific wavelength by creating alternating zones of transmission and opacity in either amplitude or phase. The focal
point is determined by the radius of the outermost zone and its width. Zone plates are difficult to fabricate on a millimeter size scale, therefore either a part of the XUV beam would be focused as the aperture of the zone plate would be smaller than the beam or the optic would be prohibitively expensive to fabricate on scale comparable to the beam. The fabrication is also difficult in the 30 nm region due to the high absorption of materials.

Parabolic tapers use the principal of total external reflection to focus the radiation. Incident light at a glancing angle results in a low loss reflection, a function that is utilised in x-ray optics. By producing a parabolic taper the radiation is focused to a particular position. Previous work within the group has shown that the alignment of such focusing optics is difficult. The ring structure around the central maxima makes them unsuitable for use within imaging experiments.

The third type of optic used is a multi-layer mirror at normal incidence. In the XUV regime reflections off a single surface normal to the incident beam are weak. By building up layers on the mirror, half the wavelength of the incident light in depth, reflections from several different layers can constructively interfere and increase the reflected light. In real experimental geometries a normal reflection is not possible. Instead the angle to the normal is kept to the minimum possible, this however results in an astigmatic focus. The experiments described in the next two chapters use a multi-layer focusing mirror fabricated by The Fraunhofer Institute. The calculated reflectivity profile for the mirror is shown in figure 6.4.
Figure 6.4: *Calculated reflectivity of the multi-layer mirror as a function of wavelength*

Figure 6.5: *Schematic of the experimental setup*

### 6.3 Experimental method

A schematic of the experimental setup for the diffraction experiments described in this chapter is shown in figure 7.2. As described above, the geometry of the generation was changed from a capillary to a 3.2 mm gas cell for ease of alignment and improvement in stability. The gas cell pressure was regulated using a Tescom ER3000 to 70 mbar as this produced the highest flux measured after the mirror. The pressure outside the gas cell was maintained at $10^{-2}$ mbar before the filter section and $10^{-5}$ mbar after the filter section. Two 200 nm aluminium filters were used to separate the fundamental and the generated harmonics.

The generated XUV beam was focused using a Mo/Si multi-layer spherical concave mirror, fabricated by the Fraunhofer Institute, with a radius of curvature of 500 mm.
The mirror had periods of 15 nm and a Mo:Si ratio of 0.4:0.6 resulting in the reflectivity calculated in figure 6.4. This mirror was used at an incident angle of 6 degrees, limited by the detector clipping the input to the mirror at narrower angles. A sample plate mounted onto a three axis stage which could move horizontally and vertically orthogonal to the axis of the focused beam as well as along the axis of the focused XUV beam. The sample plate allowed mounting of three 5 mm$^2$ silicon nitride membranes simultaneously. The detector was a water cooled Andor CCD detector mounted on a translation stage parallel to the direction of the XUV beam shown in figure 6.5. This stage could be controlled when under vacuum. The camera could be translated horizontally while at atmospheric pressure to optimise the position of the alignment beam.

6.4 Finding the focus

Before the spotsize at the focus can be characterised the position of the focus has to be found. This is also critical to the positioning of an imaging sample in later experiments. Scanning the CCD detector along the axis of the beam as shown in figure 6.6 reveals that an astigmatism is present. As the beam propagates away from the mirror a line focus is reached in the $y$ plane. Propagating the beam further shows the profile becoming circular before moving to a line focus in the $x$ plane. This data is useful in terms of confirming the focus is performing as expected and for getting an imaging sample into approximately the correct position. However, when performing an imaging experiment mounting a sample at the focus using only this data is difficult. The vacuum system must be returned to atmospheric pressure and the sample plate fixed into the system exactly 17 mm further away from the mirror than the front surface of the camera when it is at the focus. A method for reliably finding the focus position for the sample plate rather than the CCD detector is required.

Diffraction from a periodic array produces peaks at the the Bragg angle. The definition of these peaks is increased with the number of elements scattered from. Using this principle scanning an array through the focus would allow a visible change in the diffraction pattern as a result of the increasing spot size illuminating a greater number of scatterers. The sample used for this experiment was a gold coated silicon nitride membrane with an array of 2$\mu$m holes milled in a square pattern, shown in figure 6.7.
Figure 6.6: The XUV beam profile through the focus in 5 mm separations along the propagation axis

Figure 6.7: SEM image of the silicon nitride grid
6.4.1 Results

A series of diffraction patterns were recorded as the sample was translated along the axis of the beam through the focus. Each position had an exposure time of 2 seconds. The diffraction patterns are shown in figures 6.8 and 6.9. The data demonstrates that as the grid is translated away from the \( z = 0 \) position the diffraction patterns show an increase in the complexity although there is not an apparent trend within this. At \( z = 0 \) the diffraction pattern shows minimal complexity but it would be difficult to definitively say that figure 6.8b was the focus over figure 6.8a or figure 6.8c. Comparing figure 6.8k and figure 6.9e, each 6 mm away from the focus there is no obvious similarity. It is important at this point to remember that the focus for an astigmatic beam is a circle of least confusion. This means that the radius of curvature in x and y is the same giving an intensity profile that appears circular but the phase fronts are curved in opposite directions giving a saddle shaped phase front. Either side of this will be two line foci where the beam reaches a minimum width for the x and y directions independently. The phase front of the beam between the two line foci will have an opposite sign for the x and y directions. Outside of the two line foci the sign of the phase front will be the same, but the value will still be different.

6.4.2 Autocorrelations of the diffraction patterns

As the trend expected in the diffraction patterns cannot be interpreted an autocorrelation of the data was taken using a Fourier transform.

The data for the autocorrelation of the diffraction patterns is shown in figure 6.10 and 6.11. An initial inspection of the autocorrelations reveals a trend that was not apparent in the diffraction patterns. Examining figure 6.10a the position of the first order peak in the y direction is much closer to the central maxima than in the x direction. In figure 6.10j the first order peak has reached a minimum in the y direction but the peak is still moving towards the central maxima in the x direction. Figure 6.11b shows the peaks in the same position for both the x and y direction but the trend of the peaks in the y direction is away from the central maxima while the peaks in the x direction are still moving towards the central maxima. Figure 6.11f shows the position of the second line focus where the peak has reached a minimum distance to the central maxima in
Figure 6.8: Diffraction through the grid at (a) $z = -2.6\text{mm}$, (b) $z = -2.4\text{mm}$, (c) $z = -2.2\text{mm}$, (d) $z = -2\text{mm}$, (e) $z = -1.8\text{mm}$, (f) $z = -1.6\text{mm}$, (g) $z = -1.4\text{mm}$, (h) $z = -1.2\text{mm}$, (i) $z = -1\text{mm}$, (j) $z = -0.8\text{mm}$, (k) $z = -0.6\text{mm}$, and (l) $z = -0.4\text{mm}$.
Figure 6.9: Diffraction through the grid at a) $z = -0.2\text{mm}$ b) $z = 0\text{mm}$ c) $z = 0.2\text{mm}$ d) $z = 0.4\text{mm}$ e) $z = 0.6\text{mm}$ f) $z = 0.8\text{mm}$ g) $z = 1\text{mm}$ h) $z = 1.2\text{mm}$ i) $z = 1.4\text{mm}$ j) $z = 1.6\text{mm}$ k) $z = 1.8\text{mm}$ l) $z = 2\text{mm}$
Figure 6.10: Autocorrelation of the diffraction pattern at a) $z = -2.6\text{mm}$ b) $z = -2.4\text{mm}$ c) $z = -2.2\text{mm}$ d) $z = -2\text{mm}$ e) $z = -1.8\text{mm}$ f) $z = -1.6\text{mm}$ g) $z = -1.4\text{mm}$ h) $z = -1.2\text{mm}$ i) $z = -1\text{mm}$ j) $z = -0.8\text{mm}$ k) $z = -0.6\text{mm}$ l) $z = -0.4\text{mm}$
Figure 6.11: Autocorrelation of the diffraction pattern at a) $z = -0.2\text{mm}$ b) $z = 0\text{mm}$ c) $z = 0.2\text{mm}$ d) $z = 0.4\text{mm}$ e) $z = 0.6\text{mm}$ f) $z = 0.8\text{mm}$ g) $z = 1\text{mm}$ h) $z = 1.2\text{mm}$ i) $z = 1.4\text{mm}$ j) $z = 1.6\text{mm}$ k) $z = 1.8\text{mm}$ l) $z = 2\text{mm}$
the x direction but has moved away in the y direction. After this point the radius of curvature is increasing in both directions as the beam propagates away from the focal positions. Qualitatively this data shows the changing radius of curvature associated with an astigmatic focus, matching the experimental geometry used. This alone would allow the determination of the two line foci and the circle of least confusion.

6.4.3 Fitting the data to a model

Further processing of the data allows quantitative measurements of the beam parameters. The distance from the central maxima to the first maxima can be measured in terms of number of pixels. A calibration of pixels to absolute measurements can be obtained as the dimensions of the grid are known. The result of this analysis is shown in figure 6.12. The position of the two line foci are shown by the minima of the two datasets and the circle of least confusion is given where the two datasets intersect. As previously stated the two radii of curvature are the same value at this point but the phase fronts have opposite signs.

Although the XUV beam is not a single mode gaussian beam, a starting point for modelling the focusing can be obtained from equations 2.27 and 2.29. A modification has to be made to the equation to account for the beam quality. Introducing the $M^2$ term...
parameter described in section 2.5, the radius of curvature can be modelled as,

$$R = z + \left( \frac{\pi w_0^2}{\lambda M^2} \right)^2$$

(6.1)

As the z position is known, a fitting parameter of $\frac{w}{M}$ can be used. Using a fitting parameter of $\frac{w}{M} = 1.53 \ \mu m$ the result shown in figure 6.13 are obtained. The fit of the model to the data is very sensitive to the parameter, meaning that achieving an accurate fit was possible. The data at the two minima does not tend towards infinity as the model suggests. It may be interesting to retake this data with finer steps across the line focus to establish if this is a genuine or if this results from the data sampling rate. From equation 6.1 it is possible to determine the z position of the data point if it is not known by fitting to the radius of curvature away from the focus. This makes the system completely self calibrating if the reference array is well characterised.

### 6.4.4 Measuring the spotsize

By measuring the beam width it is possible to calculate a value for the $M^2$ of the beam. There are two techniques used for laser beams to obtain a value for the beam width ($w$). The first technique is a knife edge measurement, these are routinely used to find the position for 10% and 90% transmission. This distance is multiplied by a scaling factor to give a value for the beam width [53]. A series of knife edge measurements have been attempted but with no success. This may be due to the small instabilities in the beam
positioning when trying to translate the knife edge in steps of 1 \( \mu m \) or less. The second method used is to image the beam on to a CCD detector. A fit to the data using a Gaussian profile calculates the value for \( w \). As shown in figure 6.14 the majority of the intensity is located on a single pixel at the focus of the XUV beam, resulting in an upper estimate of 13 \( \mu m \) for the spotsize.

6.5 Measuring the Spectrum

A calibrated grating spectrometer is the most commonly used piece of equipment for obtaining an XUV spectrum. The experimental geometry used in the procedure prevented the beam from being coupled into the existing spectrometer. Changing the angle of a multi-layer mirror to overcome the geometric restrictions affects its reflectivity envelope. These two factors determined that a new method of measuring the spectrum off the multi-layer mirror was required. By using a mono layered crystalline structure it has been shown that the complex refractive indices can be obtained for the crystalline material at each wavelength within the beam [77]. For this technique to work the Bragg peaks of the sample must be discrete for the different wavelengths. A byproduct of this technique is that the splitting of the diffraction peaks can be used to determine the wavelengths present using Bragg’s law in transmission,

\[
n \lambda = d \sin \theta
\]  

(6.2)
Figure 5.2: SEM image of a region on the hexagonal nanosphere sample showing a ²0 µm area of the sample that is a single crystallite with minor defects.

The sample is a 2D hexagonal array of nanospheres, which is equivalent to the convolution of a hexagonal array of delta functions and a single nanosphere. The far-field diffraction pattern from the sample is therefore the Fourier transform of the array of delta functions multiplied by the far-field scattering pattern from a single nanosphere. The result is therefore a hexagonal array of delta functions that have an intensity dependent on the far-field scattering pattern from a single nanosphere. An application of crystal lattice theory can be used to calculate the angles of the interference peaks. The Mie solution can be used to determine the far-field scattering pattern. The Mie solution is described in section 5.3.1. The application of crystal lattice theory to this problem is given in the following paragraphs.

To describe the array of spheres, it is convenient to introduce the Bravais lattice. It is defined as an infinite array of points such that a crystal will look identical from whichever of the points it is observed from. The position

Figure 6.15: An SEM image of the hexagonally close packed polystyrene spheres

where \( n \) is the diffraction order, \( \lambda \) is the wavelength, \( d \) is the object size and \( \theta \) is diffraction angle. From this it can be seen that with an accurate measure of the sample to detector distance the wavelength can be calculated.

The crystalline structure used in this experiment was a single layer of polystyrene nano spheres with a diameter of 196 nm, assembled onto a silicon nitride membrane by C.F. Chau. The structure of this monolayer shown in figure 6.15, is a hexagonal close packed array. The sample was mounted onto the sample holders used in the previous section. The sample was placed at the focus and translated in x and y, to find an area containing a single crystal domain. If two domains are present then the the two diffraction patterns are overlaid giving 12 peaks in the first ring of diffraction maxima rather than the expected 6 peaks from a hexagonally close packed array, as shown in figure 6.16.

Focusing on one of these peaks shows the composition of the beam contains several harmonics. The angle to which the harmonic scatters is given by equation 6.2. As the size of the spheres and the distance to the detector is known, it is possible to calculate the spectrum causing the diffraction pattern. Figure 6.17 shows the spectrum obtained from the diffraction pattern in figure 6.16. The positions of the peaks match the expected harmonic positions accurately, although the peak profiles are significantly broader than those measured using a conventional spectrometer. This may be due to the relatively
Figure 6.16: The diffraction pattern plotted on a log scale of a hexagonally close packed array of 196 nm polystyrene spheres

low number of individual objects scattered off as the resolution of a grating is directly proportional to the number of elements in the grating.

6.6 Conclusions and further work

The methods described in this chapter give novel solutions to obtaining several experimental factors reliably and efficiently. The final measurements required to give the beam width and hence a $M^2$ value would be very useful. Currently an upper limit on the focal spotsize can be given as most of the intensity is contained within a single 13 $\mu$m square pixel.

The method used for identifying the focal plane of the sample mount is not only useful to future experiments within XUV microscopy but may have application to other systems which rely on short working distances. Some further investigation may be required to optimise the aperture size and pitch for another spectral region.
Determination of the spectrum at the point of imaging is crucial to understanding the results presented in the next chapter. A system that does not require a change in experimental configuration between recording the spectrum and performing the imaging experiment gives reliable information about the composition of the beam used. In a system with multiple frequencies this may lead to the ability to perform deconvolution experiments. This would also require the relative amounts of each harmonic present to be altered in a controlled manner.
Chapter 7

Coherent diffractive imaging of a binary object

Using the system outlined in the previous chapter to perform imaging experiments poses several challenges. It described the difficulty in obtaining high quality XUV lenses for re-imaging and directing the experiment towards coherent diffractive imaging (CDI). The constraints on the geometry of a CDI experiment were explained in section 2.8.6 and will be outlined for this experiment later in this chapter. The sample used in the experiment is related to these constraints and was chosen appropriately. The end of the previous chapter described the beam as being composed of three predominant harmonics as well as showing evidence of a least three other low intensity harmonics. The CDI algorithm being used in this experiment is based upon an illumination source that is monochromatic. Prior to performing the experiment there was no certainty to the outcome, the aim of this experiment was to investigate the limitations of using a monochromatic algorithm with a polychromatic source.

Coherent diffractive imaging is discussed in depth in section 2.8, as a brief recap, when experimental conditions are set so the sample size (a) is small and the sample to detector distance (z) is large, the diffraction plane is far-field as shown in equation 7.1.

\[
\frac{a^2}{z\lambda} \ll 1 
\]  

(7.1)

The field at this point is the Fourier transform of the electric field immediately after the sample. The image recorded on the detector is the modulus squared of the field and
hence has no phase information. Using iterative phase reconstruction algorithms it is possible to calculate the missing phase and hence reconstruct the sample used.

The first results for CDI within the HHG community were published in 2007 with a resolution of 214 nm [78]. These results rely upon the use of two multilayer mirrors to monochromate the beam. The experiment performed in this chapter, in contrast to that work, used a polychromatic beam. Work by Abbey et al. using a synchrotron, states that use of a polychromatic beam to obtain a diffraction pattern prevents the CDI algorithm from reconstructing [79]. With a significantly broader relative bandwidth preliminary results are presented in this chapter showing evidence of reconstruction.

7.1 Sample

During the setup of any new microscopy system a test sample is normally chosen that is well understood in order that the microscope is the only unknown. For this reason a sample was fabricated in preference to placing a pinhole over an existing sample. Another factor in the decision was that if a pinhole was to be used the distance from sample to pinhole had to be less than 50 µm to prevent the beam from having a Gaussian profile at the sample. The simplest sample to use was a binary amplitude object so high contrast could be achieved. Fabrication of the object was performed using a focused ion beam (FIB).

7.1.1 Fabrication

A 2.8 µm by 2.5 µm sample was fabricated from a thin gold coated (100 nm) silicon nitride membrane (50nm) shown in figure 7.1. The transmission of this combination at 27 nm is less than 10^{-5}. The transmission region was milled through completely giving a transmission of 1, the sample can therefore be considered binary. The reason for choosing gold as the coating layer was that the sputtering rate for FIB milling is highest for gold, simplifying the fabrication process. A 50 µm² guide hole was also milled into the membrane, this was to aid initial alignment of the sample to the beam.
7.1.2 Experimental constraints of the algorithm

As outlined in section 2.8 there are several constraints on the sample and its position relative to the detector as it is observed. When using an aperture, its size must be less than the size of the beam. Without this limit the algorithm does not have a constraint to apply and therefore cannot reach a solution. In fabricating a sample the same constraint must be observed. The sample fabricated in this case was 2.8 \( \mu \text{m} \) by 2.5 \( \mu \text{m} \). For a given object size the detector must be placed significantly far away to give a large enough oversampling ratio as described in section 2.8.6. For the experimental geometry used the detector was placed 22 mm away from the sample giving an oversampling ratio of 22, as defined by equation 2.51. This distance is also related to the reconstructed pixel size which in this case was 107 nm. The final constraint for this algorithm is the Fresnel number, to use the Fourier transform relationship in the algorithm \( F \ll 1 \) to ensure Fraunhofer diffraction. Using equation 2.50 the Fresnel number for the experiment was 0.01.
7.2 Experimental method

The experimental setup used in this chapter is the same as used in the previous chapter, it is briefly outlined again here. A schematic of the experimental setup for the diffraction experiments described in this chapter is shown in figure 7.2. The generation geometry was a 3.2 mm length gas cell. The gas cell pressure was regulated using a Tescom ER3000 to 70 mbar as this produced the highest flux measured after the mirror. The pressure outside the gas cell was maintained at $10^{-2}$ mbar before the filter section and $10^{-5}$ mbar after the filter section. Two 200 nm aluminium filters were used to separate the fundamental and the generated harmonics.

The generated XUV beam was focused using a Mo/Si multilayer spherical concave mirror, with a radius of curvature of 500 mm. This mirror was used at an incident angle of 6 degrees, limited by the detector clipping the input beam to the mirror at narrower angles. A sample plate was mounted onto a three axis stage which could move horizontally and vertically orthogonal to the axis of the focused beam, as well as along the axis of the focused XUV beam. This provided full control of the sample within the imaging beam. The sample plate was designed with mountings for three 5 mm$^2$ silicon nitride membranes simultaneously. The detector was a water cooled Andor (CCD) detector mounted on a translation stage parallel to the direction of the XUV beam. This stage could be controlled when under vacuum. The detector could be translated horizontally while at atmospheric pressure to optimise the position of the alignment beam.

The sample was mounted on to the sample plate, shown in figure 7.3, and aligned to the 50 µm guide hole at atmospheric pressure using a low powered IR beam. The CCD...
Figure 7.3: A photograph of the sample plate used for mounting sample

The process described above did not immediately locate the sample. Although the measurements were accurate, the mounting of the sample onto the sample plate may have resulted in a slight rotation. The problem was solved using a spiral search pattern in 1 micron steps. This led to a new position for the sample relative to the guide hole for that particular mounting. This relative distance was noted and constantly located the sample. Once the sample had been located its position was optimised for maximum flux on the CCD detector.
In the previous chapter diffraction from an array of spheres was used to derive the spectrum. Splitting of the diffraction peaks at the Bragg angle allowed the wavelength for each peak to be extracted. Using this sample the diffraction would not occur at specific Bragg peaks so it would not be possible to deconvolve the signals from each wavelength. The expected result was therefore a blurring of the fringes within the diffraction pattern resulting in either a loss of resolution or a failure for the algorithm to converge.

7.3.1 Recording the data

A series of 15 exposures were collected with integration times between 60 and 180 seconds and are shown in Appendix A. This allowed the signal on the CCD detector to be maximised at high angles without reaching saturation at the centre of the detector. It is important to maximise the high angle data as it relates to the smaller features of the sample. Ten of the fifteen exposures were taken forward, removing two with low signal levels and three which had reached saturation in the centre. These exposures were background subtracted and compared to each other, revealing that the diffraction pattern peak positions on the detector were constant. Using this result the diffraction patterns were summed to increase the signal to noise ratio, the result of which is shown in figure 7.4.

Examining figure 7.4 it can be seen that there were several constant defects on the diffraction pattern. These are the result of small amounts of oil on the detector. Cleaning these CCD detectors was a difficult process as the surface cannot be touched. As a result the detector was used in the condition observed for these experiments. It was also observed that the diffraction pattern was not centred on the detector. In order for the diffraction pattern to be re-phased correctly a signal is required for a given radius for a complete $2\pi$ angular spread. Consequently any data that does not lie on or inside a circle with a magnitude equal to the nearest edge of the CCD will not contribute to the reconstruction. For this reason the diffraction pattern shown in figure 7.4 was cropped and centred as shown in figure 7.5. This figure also shows the effect of binning the diffraction pattern. Binning involves taking a two by two array of pixels and reducing them to a single pixel. This process was applied to reduce the effect of the readout noise but it also has the effect of reducing the oversampling. In this case the reduction of the
oversampling is not enough to reduce it below the minimum of 5 as described in section 2.51.

7.3.2 Setting up the algorithm

The cropped diffraction pattern was used as the input for the algorithm. However, the beam being used to illuminate the sample is polychromatic and the algorithm does not take this into account during the reconstructions. The algorithm used is a combination of the HIO and ER algorithms described in section 2.8. The algorithm uses the hybrid input and output for all of the iterations but for the final half of the number of iterations the \( \beta \) value which begins at 0.9 is increased to 1. This change affects the amount by which the algorithm reduces the signal outside of the support constraint, a \( \beta \) value of one is characteristic of error reduction algorithms where everything outside of the mask is set to zero. The starting support constraint for the algorithm is the autocorrelation of the diffraction pattern shown in figure 7.6b, this was thresholded at 0.2%.

The autocorrelation shown in figure 7.6 is also used as the amplitude starting point for the algorithm, this is multiplied by a guessed phase to acquire a starting field for the
Figure 7.5: The diffraction pattern after centring and binning

Figure 7.6: The autocorrelation (a) of the diffraction pattern and the thresholded autocorrelation (b) used as the sample support
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Number of iterations</td>
<td>1000</td>
</tr>
<tr>
<td>Beta</td>
<td>0.9</td>
</tr>
<tr>
<td>Error reduction</td>
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</tr>
<tr>
<td>Error reduction start point</td>
<td>0.5</td>
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<tr>
<td>Shrinkwrap start value</td>
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<tr>
<td>Shrinkwrap end value</td>
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</tr>
<tr>
<td>Shrinkwrap intervals</td>
<td>20</td>
</tr>
<tr>
<td>Autocorrelation threshold</td>
<td>0.2%</td>
</tr>
</tbody>
</table>

Table 7.1: Parameters used in the CDI algorithm for the results presented in this chapter

algorithm. The parameters for the algorithm were defined and are listed in table 7.1. The number of iterations relates to the number of times the algorithm runs through the loop before coming to completion. Beta is described above and relates to the scaling of signals inside and outside of the mask where the signal outside the mask is given by \( s = s'(1 - \beta) \). Error reduction refers to the change in beta from 0.9 to 1 and the error reduction start point is the how far through the total number of iterations the beta value is changed. The shrinkwrap start and end values are the beginning and ending size parameters for the gaussian blur function used within the shrinkwrap step. The shrinkwrap intervals determine how many iterations occur between each shrinkwrap, the value of the shrinkwrap decreases linearly between the start and end points. The autocorrelation threshold is the value used to calculate the mask and starting guess for the algorithm. Signals above the autocorrelation threshold are set to 1 and those below are set to 0.

7.3.3 Reconstruction of the sample

The amplitude result of the algorithm is shown in figure 7.7a and the phase as a function of the amplitude is shown in figure 7.7b. An initial examination of the reconstructed sample shows strong similarities to the FIB image shown in figure 7.1. In figure 7.7a the intensity distribution of the amplitude reconstruction is not uniform, this was not expected for a binary sample. The features at the extremities of the sample are all present although the right hand side of the cross has not resolved particularly well. The amplitude reconstruction also exhibits a speckled pattern across it. The intensity values of the maxima in the speckled pattern are less than 10% of the maximum of the reconstructed sample but are still significant and cannot be ignored. This may be the
Figure 7.7: The result of 1000 iterations of the CDI code. a) The amplitude of the reconstruction b) The phase of the reconstruction where the brightness is proportional to the amplitude of the reconstruction shown in (a)
result of the algorithm trying to account for the multiple harmonics present in the beam or may be a result of noise in the diffraction pattern. Although the background has been removed there will still be some sources of noise. The readout noise that has been minimised but not eliminated and the counting noise is inherent to the measurement.

The phase shown in figure 7.7b shows a flat phase across the area of high intensity within the sample in both directions. As can be seen in the previous chapter, using an off axis focusing geometry should not result in a flat phase anywhere through the focal region. If the sample was at the circle of least confusion a saddle shaped phase would be expected.

The algorithm produces an error measurement for each iteration. This measurement is made by a comparison of the recorded diffraction pattern and the diffraction pattern generated by the algorithm. A plot of the of the error is shown in figure 7.8. The error rapidly drops over the course of the first few iterations before showing a smaller variation for the remainder of the algorithms iterations.

There are a couple of features to be noted here. The shrinkwrap is enforced every 20 iterations; it can be seen that two iterations after this there was a decrease in the error. The other interesting feature is that the minimum error does not occur at the end of the
algorithm but at the 502\textsuperscript{nd} iteration. This suggests that looking at the final solution of the algorithm may not be an appropriate method for examining the output.

As a result of examining the error function, figure 7.9 was produced showing the amplitude (a) and phase (b) of the lowest error solution.

The amplitude reconstruction for the lowest error solution shows a strong resemblance to the SEM image of the sample. The image has clearer definition in all of the extremities of the sample in comparison to the final reconstruction of the algorithm. The sample does, however, still exhibit the speckled pattern across it that was present in the final reconstruction. As observed in the final reconstruction, the phase does not produce the saddle shape expected. The phase is flat across most of the sample with a small gradient in the top left corner. The discrepancy in the phase may be linked to the same factors described above for the presence of the speckled pattern but as yet it has not been satisfactorily explained.

Although this result appears to be positive with respect to the viability of the technique it is only one result from the algorithm. To test the reliability of the algorithm a further 99 sets of reconstructions were produced to create a data set which could be interrogated. Figure 7.10 shows the value of the lowest error for each of the 100 runs of the algorithm. Scanning through the data one can see a large amount of the reconstructions look very similar, however, there are some notable exceptions. Run 18 show the maximum value for the lowest error out of all the runs. Figure 7.11 shows both the final reconstructions (a & c) and lowest error reconstructions (b & d) for run 18. These reconstructions do not resemble the sample in either the amplitude or phase, although at the final reconstruction the beginnings of a cross pattern can be observed.

A system that produces non-results for high errors allows one to discount the anomalous results. However, another case is shown in figure 7.12, the lowest error is $1.636 \times 10^{-7}$ in comparison to the average error of $1.6464 \times 10^{-7}$ for the data set. In this data the lowest error solution does not resemble the sample at all but the final solution is recognisable as a reconstruction of the original sample. Earlier in the chapter the suggestion was that the lowest error solution gave a better reconstruction than the final solution but this appears not to be universally applicable.
Figure 7.9: The lowest error result of the algorithm. a) The amplitude of the reconstruction b) The phase of the reconstruction where the brightness is proportional to the amplitude
7.3.4 Resolution

One of the key criteria for a microscope is the resolution it can achieve. Although this experiment was not successful in producing a stable output the reconstructions did produce some results from which the sample shape could be observed. By taking a line through the first reconstruction data shown in figures 7.7 and 7.9 it was possible to see the 200 nm line widths on the sample. Figure 7.13a shows the measured line width from an SEM image of the sample and figure 7.13b a line profile through the reconstructions at the same point as the measurement.

Earlier in the chapter the reconstructed pixel size was calculated to be 107 nm, this does not fit with the profiles shown in figure 7.13 suggesting a rescaling process is happening that is not yet understood.

The line profiles suggest that each pixel in the reconstruction is approximately 33 nm which gives a resolution far better than expected and is approaching the wavelength of the illuminating source. If this could be achieved with a monochromatic source, allowing some of the ambiguity of this result to be resolved, then the future for this technique looks promising.
7.4 Conclusions and further work

The results presented in this chapter show the first CDI results from our XUV microscope. Using a known sample it was possible to resolve features of 200 nm with a polychromatic source. The algorithm did not require any modification to obtain a reconstruction, however the reconstructions were not reliably repeatable. Inconsistencies in the phase obtained and the expected phase along with the specular pattern on the reconstructed amplitude could be due to either noise in the diffraction pattern or the algorithm compensating for polychromatic source. Further work would be required to investigate these problems further.

There are several steps that follow on from this experiment both theoretically and experimentally. The main difficulty in performing this experiment was that both the CDI technique and the polychromatic source were being tested in a single experiment. A lack
Figure 7.12: The lowest error result of the algorithm. a) The amplitude of the reconstruction b) The phase of the reconstruction where the brightness is proportional to the amplitude

Figure 7.13: A line profile through the reconstruction of the data for the lowest error solution (green) and the final solution (blue)
of experience has limited our understanding of the results so a repeat of the experiment with a monochromatic source would be an ideal start to understanding the CDI technique. Monochromatic beams are usually obtained by using a pair of narrow bandwidth multilayer mirrors for these experiments. Subsequent experiments with a polychromatic beam may then yield a clearer picture of the effects involved. It would also be of interest to develop a known phase sample, if this technique is to push towards imaging biological molecules then phase imaging is likely to become more important due to the limited contrast in the amplitudes of the samples. On a day to day level, improving the signal to noise at high angles will aid the reconstruction of finer detail. Experiments elsewhere have used a traditional beam stop to remove the central maxima from the diffraction pattern where as the experiment at FLASH used a mirror with a hole drilled in the centre to reflect the diffraction pattern and transmit the central maxima. This is a more elegant solution but also significantly more costly. A third way that currently has not been used is to saturate the centre of the diffraction pattern and remove it in the algorithm in a similar way to the previous two techniques. However, the likely signal gain at high angle will be limited by the pixel bleeding in the saturated part of the camera.

There is also scope for development of the algorithm. Recent publications have shown a working algorithm for a HHG source using the broad frequencies within the algorithm [80][81]. Currently we have been unable to replicate the algorithm they have produced.
Chapter 8

Conclusion and Further work

The work presented within this thesis is separated into two distinct sections. The first, discussed in chapters 4 and 5 focused on the development of a capillary based HHG source. The second section moved away from source development and towards use of a HHG source to perform CDI experiments.

Chapter 4 begins by describing the fabrication and mounting of a capillary for use in HHG. Measurements of the transmission through the capillary for a calibrated coupling efficiency of 50% were compared to that of a simple linear ionisation model, showing good agreement. A pair of filters were spectrally selected to image fluorescence from excited argon atoms and excited singularly ionised argon ions. These filters were used to map the distribution of both ions and excited atoms along the capillary length from which the driving laser intensity could be inferred. These results were compared against the same simple model showing strong evidence of mode beating between the first and second modes of the capillary. A second data set was collected after an upgrade to the laser system. The calibrated coupling efficiency was measured to be 80%, which had a strong effect on propagation along the capillary. The presence of extra peaks in the fluorescence distribution, some of which had a sharp profile, suggested that a more comprehensive model was required.

In chapter 5 the model was changed to a non-linear Schrödinger equation model. A series of measurements were performed to compare the spectral output of the capillary to that predicted by the model. A novel measurement of the spatio-spectral distribution at the capillary exit showed strong correlation to the model. From the model it was
possible to see that the non-linear and ionisation propagation effects were significant factors in the evolution of the pulse along the capillary. The combination of plasma defocusing and plasma induced refractive index change led to large amounts of blue shifting appearing in higher order modes. The development of an accurate propagation model would allow for a parameter scan to look for optimal combinations of intensity, gas density and absorption as well as the possibility to combine the model with the TDSE model described in section 2.1.2 to make a full description of the HHG process. To make the source usable for imaging improvements to the beam stability must be made as well as the stability of the capillary.

Chapter 6 begins with a change in generation geometry from a capillary to a gas cell. This change was necessary due to the instabilities in the capillary output over time. The beamline was constructed to match the requirements of an imaging experiment using an off axis multilayer mirror. The position of the focus was determined by scanning a CCD detector along the propagation axis showing an astigmatism in the focus of the beam. A diffractive measurement to find the focus was also developed to allow accurate positioning of the sample plate, important when maximising the flux onto an imaging sample. An estimate of the size of the focal spot was made, although the difficulties in performing accurate knife edge measurements due to beam instabilities prevented a conclusive result. The spectral composition of the beam was measured as the multilayer mirror used to focus the XUV has a reflectivity profile that spans several harmonics. The measurement showed a polychromatic beam with three dominant harmonics. The algorithm used for the CDI experiments, outlined in section 2.8 requires a monochromatic input making this a non ideal case. However, the results shown in chapter 7 cast doubt on this assertion.

Finally in chapter 7 a description of the CDI experiment is given, including the design and fabrication of a test sample using a FIB. Using this sample a series of diffraction patterns were recorded using the setup described in chapter 6. The diffraction patterns were used as the input to the phase retrieval algorithm, giving reconstructions where it was possible to resolve features of 200 nm. The algorithm did not require any modification to obtain a reconstruction despite the polychromatic source, however the reconstructions were not reliably repeatable. A scaling difference between the predicted reconstruction pixel size and the pixel width of the features on the reconstruction was observed.
The outcome of the CDI experiment is not a definitive proof that a monochromatic algorithm can be used with a polychromatic source. It does however show that the presence of multiple harmonics does not stop the algorithm from reconstructing completely. This is important because as the wavelength is decreased, the spacing between the harmonics is reduced therefore producing a monochromatic beam from a HHG source becomes more difficult. At the current wavelength using a second multilayer mirror to isolate a single harmonic would allow for a better understanding of the relationship between the algorithm and the data. Extending the CDI to include Ptychography would also allow for a larger sample area to be imaged without any loss of resolution, this has been demonstrated in the visible [82] and with hard X-ray radiation from synchrotrons [83]. In chapter 1 it was stated that the time resolution was important for investigating biological samples. The current recorded diffraction patterns require 100,000 shots to acquire, meaning that a large amount of source development is required before single shot imaging can be achieved.
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Appendix A

Fluorescence lineplots and XUV spectra from a 7 cm capillary
The following figures are plotted together in figure 4.12

Figure A.1: Emission from singly ionised argon species as a function of position along the propagation axis of the capillary, for (a) 20 mbar, (b) 40 mbar. The vertical lines indicate the positions of the gas inlet holes.
Figure A.2: Emission from singly ionised argon species as a function of position along the propagation axis of the capillary, for (a) 60 mbar, (b) 80 mbar, (c) 100 mbar. The vertical lines indicate the positions of the gas inlet holes.
Figure A.3: Emission from singly ionised argon species as a function of position along the propagation axis of the capillary, for (a) 120 mbar, (b) 140 mbar, (c) 160 mbar. The vertical lines indicate the positions of the gas inlet holes.
The following figures are plotted together in figure 4.13

Figure A.4: XUV spectra from the capillary, for (a) 20 mbar, (b) 40 mbar. The vertical lines indicate the positions of the gas inlet holes.
Figure A.5: XUV spectra from the capillary, for (a) 60 mbar, (b) 80 mbar, (c) 100 mbar. The vertical lines indicate the positions of the gas inlet holes.
Figure A.6: XUV spectra from the capillary, for (a) 120 mbar, (b) 140 mbar, (c) 160 mbar. The vertical lines indicate the positions of the gas inlet holes.
Appendix B

Diffraction patterns from a binary object
The following are the diffraction patterns collected in chapter 7 for a range of integration times.

Figure B.1: The first five diffraction patterns recorded with integration times of (a) 60 s (b) 120 s (c) 180 s (d) 150 s and (e) 150 s. Results (a) and (b) are underexposed while (c), (d) and (e) have reached saturation.
Figure B.2: A subsequent set of five diffraction patterns recorded with integration times of 140 s
Figure B.3: A final set of five diffraction patterns recorded with integration times of 140 s
Appendix C

Publications
Spatially resolved Ar* and Ar** imaging as a diagnostic for capillary-based high harmonic generation

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Abstract

Coherent x-ray–ultraviolet radiation can be generated by the highly nonlinear interaction between a gas target and high intensity ultrafast laser pulses using the high harmonic generation (HHG) process. Guiding the fundamental laser field inside a hollow capillary waveguide improves phase matching and extends the potential interaction length. However, the propagation of an intense pulse within a capillary waveguide filled with ionizable gas is complex, as the pulse creates a plasma, which in turn strongly affects the propagation. Previous work has used emission from the excited gas to study propagation of nanosecond pulses in capillary waveguides. In this work we demonstrate spectrally resolved imaging of the plasma created by intense femtosecond pulses within a capillary during an HHG experiment. The spectral and spatial resolution is used to separate contributions from ions and neutral species, and is an effective diagnostic for the local pressure and modal intensity variations along the waveguide.

Keywords: high harmonic generation, capillary waveguide, mode beating

1. Introduction

The advent of high intensity ultrafast lasers has opened up new areas for atomic physics, and turned techniques such as high harmonic generation (HHG) from experiments into potentially useful sources of XUV and soft x-ray radiation. Many experiments using intense pulses require control of the interaction length and phase matching. One technique for achieving this is by propagation of the high intensity pulses through a hollow capillary [1]. The propagation of intense femtosecond laser pulses through a hollow capillary waveguide filled with gas results in ionization, which in turn leads to significant changes in both spectrum and intensity of the laser pulse [2]. It has been shown that this can lead to blue-shifting [3], spectral broadening and splitting [4] of the harmonic spectrum. In this paper we study the propagation of intense pulses through capillaries during HHG, and use the fluorescence of the excited atoms and ions as a way of spatially mapping the excitation process. A simple model is developed to explain the distribution of ionized gas, and show how the spatial distribution of ionization in the capillary may be controlled by altering the conditions of coupling into the capillary, and correlated with the emission recorded experimentally. This model will prove useful in the optimization of future high harmonic generation experiments. If a well aligned, linearly polarized TEM₀₀ source is coupled to a hollow capillary, only the EH₁₀ modes will be excited. Figure 1 illustrates the intensity and electric field distribution in the EH₁₁ and EH₁₂ modes. In this case the electric fields in the system can be treated as scalar. Marcatili
and Schmeltzer [5] derived expressions for the electric field of the EH\textsubscript{im} waveguide modes which can be simplified to give

\[ E_{im} = J_0 \left( \frac{u_{im} r}{a} \right) \]  \hspace{1cm} (1)

where \( a \) is the waveguide radius and \( u_{im} \) is the \( m \)th root of the first kind, \( J_0 \).

It is possible to create periodic intensity variations along the propagation direction of a hollow waveguide by simultaneously exciting multiple waveguide modes. The mode propagation can be expressed as

\[ E = E_0 \exp(i(\gamma z - \omega t)) \]  \hspace{1cm} (2)

where \( \omega \) is the angular frequency and \( \gamma \) is the propagation constant, given by

\[ \gamma = \beta + i\alpha \]  \hspace{1cm} (3)

Here \( \alpha \) is the modal attenuation and \( \beta \) is the phase velocity defined as

\[ \alpha = \left( \frac{\mu_m}{2\pi} \right) \frac{2 \lambda^2 (\nu^2 + 1)}{a^2 \sqrt{\nu^2 - 1}} \]  \hspace{1cm} (4)

\[ \beta = \left( \frac{2\pi}{\lambda} \right) \sqrt{1 - \frac{1}{2} \frac{\mu_m \lambda^2}{2\pi a}} \]  \hspace{1cm} (5)

where \( \nu \) is the refractive index of the waveguide material.

The beat length \( L \) between two modes is given by

\[ L = \frac{2\pi}{\Delta \beta} \]  \hspace{1cm} (6)

where \( \Delta \beta \) is the difference in real part of the propagation constant of the two capillary modes, which for the EH\textsubscript{11} and EH\textsubscript{12} modes is given by

\[ \Delta \beta = \frac{\lambda}{4\pi a^3} \left[ u_{12}^2 - u_{11}^2 \right] \]  \hspace{1cm} (7)

The efficiency of coupling of an incident Gaussian beam to each of the EH\textsubscript{im} capillary modes can be calculated using the overlap integral

\[ A_m = \frac{\int \int E_{1m} E_i r \, dr \, d\theta}{\int \int |E_{1m}|^2 r \, dr \, d\theta} \]  \hspace{1cm} (8)

where \( E_i \) is the incident electric field and \( r \) is the radius of the capillary.

**Figure 1.** (a) The electric field profile through the centre of the waveguide normalized to the maximum electric field and (b) the two-dimensional intensity distribution of the EH\textsubscript{11} mode normalized to the maximum intensity, and similarly (c) and (d) for the EH\textsubscript{12} mode.

**Figure 2.** Calculated normalized coupling efficiency \( \chi \) of a perfectly aligned TEM\textsubscript{00} beam as a function of the spot size to bore radius ratio, for the (−−) EH\textsubscript{11} and (−−−) EH\textsubscript{12} modes, where \( \chi_m = A_m^2 \int \int E_i^2 r \, dr \, d\theta \). This figure is adapted from [6].

In most capillary-based high harmonic generation experiments, the intention is usually to couple the maximum energy into the lowest order EH\textsubscript{11} mode because the higher order modes are more lossy. It can be seen from figure 2 that the optimum condition for coupling a TEM\textsubscript{00} input beam into the EH\textsubscript{11} mode is given by \( w = 0.64a \), where \( w \) is the radius at \( 1/e^2 \) of the intensity of the beam [6]. However, there are no coupling conditions for which there is zero probability of coupling into higher order modes and even at optimum coupling for the EH\textsubscript{11} mode, 2\% of the energy will be coupled into higher order modes; therefore with a TEM\textsubscript{00} input laser beam, a superposition of two or more capillary modes will always be present, meaning that mode beating will always occur in the capillary.

For decreasing spot sizes, from this ideal value, increasingly higher order modes will be excited; for example, the most efficient coupling into the EH\textsubscript{12} mode may be achieved for a waist size to bore ratio of \( w/a = 0.26 \), with 42% of the power coupled into the EH\textsubscript{12} mode. Figure 3(a) shows the radial intensity in a 75 µm radius, 70 mm long capillary, plotted as a function of position along the capillary for a superposition of modes optimally coupling into EH\textsubscript{11} but with contributions from higher order modes, where the beam waist to bore ratio \( w/a = 0.64 \) and for an input power of 800 mJ.
Figure 3. Radial intensity distribution for a 70 mm long, 75 µm radius capillary, of a (a) superposition of modes which corresponds to $w/a = 0.64$, optimal for coupling into EH$_{11}$ compared with (b) the EH$_{11}$ mode. The colour map scale is in W cm$^{-2}$ and the plots are normalized by the integrated power.

Figure 4. Percentage of power coupled into the EH$_{11}$ mode as a function of total transmission through the capillary for random alignments of a beam of given spot size ($w = 0.64a$) in a five-dimensional space including the lateral position of the beam relative to the centre of the capillary, beam waist position and angular alignment of the beam relative to the capillary axis.

2. Modelling

To aid in the understanding of the experimental results a simple model has been developed to model propagation and ionization in the capillary. The simulation proceeds by defining a Gaussian input beam with a given beam waist and arbitrary alignment in three spatial and two angular dimensions, and calculating the coupling efficiency $A_m$ for each capillary mode using equation (8). The electric field at any point in the capillary may then be calculated from the weighted sum of the appropriate contributions of each of the $n$ possible EH$_{1m}$ waveguide modes

$$E(r, z) = \sum_{m=1}^{n} A_mE_{1m}(r, z).$$  

Each mode is independently propagated a short distance $\Delta z$ along the longitudinal axis of the capillary, using equation (2)
to account for the phase advance and attenuation losses of each of the capillary modes. The new spatial profile is again calculated according to equation (9).

As the pulse propagates, it ionizes the gas in the capillary. The ionization rate was calculated from Keldysh theory [7], integrating the rate over time to find the fraction of atoms $\eta$ ionized. The energy taken to ionize the gas may then be calculated according to

$$E_I(r, z) = N(r, z) I_p \eta$$  \hspace{1cm} (10)$$

where $I_p$ is the ionization potential of the atom and $N$ is the number of atoms in the volume element

$$N(r, z) = \frac{2\pi r \Delta r \Delta z P N_{\text{atm}}}{\Delta_1}$$  \hspace{1cm} (11)$$

where $r$ is the radius of the volume element, $\Delta r$ is a small step across the capillary’s radius, $\Delta z$ is a small step along the capillary’s longitudinal axis, $P$ is the pressure in bar and $N_{\text{atm}}$ is the number density at 1 bar. The gas pressure profile in the capillary was calculated using the computational fluid dynamics package Ansys CFX. The capillary is 70 mm long with two 300 $\mu$m holes placed 20 mm from either end of the capillary. The model shows the pressure to be constant between the holes and after a stepped pressure drop at each hole the pressure decreases linearly towards the ends of the capillary as shown in figure 5.

The model that we have constructed can be used to describe the ionization induced energy loss as a function of pressure in the capillary. Figure 6 shows a comparison of the theoretical calculations outlined above to experimental measurements of the output energy from the capillary as a function of argon gas pressure for three different input laser energies. The overall fit is good; discrepancies are due to not being able to experimentally measure the beam waist position and spatial and angular offsets. Since they were kept constant in the experiment, the coupling parameters were calculated by fitting the parameters to all three data sets.

3. Experimental method

The experimental set-up is shown in figure 7. A 40 fs, 790 nm laser pulse at 1 kHz repetition rate was coupled
into the 70 mm long hollow capillary using a 0.7 m plano-
convex lens. When perfectly aligned, \( w/a \approx 0.64 \) for this
configuration. The input pulse energy was controlled between
378 and 840 \( \mu \)J (intensities between 2.8 and 6.3 W cm\(^{-2}\)) by
inserting reflective neutral density filters into the beam. A pair
of 300 \( \mu \)m holes were drilled 20 mm from either end of the
70 mm long capillary with a radius of 75 \( \mu \)m. This was used
to leak argon gas at pressures up to 200 mbar into the system
from a pressure regulated supply. The ends of the capillary
are attached to a vacuum system which maintains the pressure
\(< 10^{-5} \) mbar.

The visible emission from the ionized argon was
categorized using an Acton 300i spectrometer and figure 8
shows the argon emission spectrum recorded between 400 and
580 nm. A 600 nm low pass interference filter with optical
density 3 at the laser wavelength (CVI Technical Optics) was
used to attenuate the laser, in combination with a 420 or 488 nm
narrow band pass filter (10 nm FWHM) to select intense lines
corresponding to emission from neutral argon or singly ionized
argon, respectively [8]. The plasma emission from the side
of the capillary was imaged onto a CCD camera (Princeton
Instruments Pixis 400) placed perpendicular to the axis of the
capillary.

4. Results

Fluorescence data were recorded at both low (50%) and high
(80%) coupling conditions, both with \( \sim 840 \mu \)J input pulses.
The two data sets show different behaviours, as the degree of
nonlinearity is increased with increasing intensity. In each case
the fluorescence was recorded as a function of the gas pressure
within the capillary, keeping the input laser intensity constant.
In figure 9, the emission from both excited neutral argon and
excited argon ions is shown for the low coupling case. In order
to produce a line plot of intensity versus length, the emission is
summed vertically, and plotted as a function of position relative
to the entrance of the capillary. The singly ionized and neutral
states of argon are seen to have very different distributions
within the capillary. The intensity from the ion emission is
much greater at the start of the capillary, and reduces along
the length of the capillary. The emission from excited atoms
follows the pressure distribution much more closely. This
indicates that the excitation mechanism for ions has a higher
order of nonlinearity compared to the excitation of atoms, as
might be expected if it is part of the ionization process. A
strong mode beating pattern can also be observed for both
species, predominantly between the EH\(_{11}\) and EH\(_{12}\) modes
(the theoretical positions of the peak intensities of the beating
pattern are indicated by the vertical lines in the plot). Higher
order mode beats can also be seen, resulting from the presence
of small fractions of higher order modes.

Figures 10(a), (c) and (e) show the pressure variation of
the beat pattern in the case of low coupling efficiency. As gas
pressure is increased, the effective nonlinearity of the medium
is increased. Little variation in the beat pattern is observed as
a function of pressure, indicating that nonlinear mode mixing
into higher order modes is not a significant effect, with most of
the power remaining in EH\(_{11}\) and EH\(_{12}\).

Figures 10(b), (d) and (f) show the theoretical prediction of
the total number of atoms ionized along the capillary, which

Figure 8. Experimentally measured emission spectrum from the
capillary filled with 200 mbar of argon gas with an incident 40 fs
840 \( \mu \)J laser pulse. The solid lines represent the area of the spectrum
transmitted by the 420 nm bandpass filter and the dashed lines
represent the 488 nm bandpass filter.

Figure 9. Emission from (a) neutral and (b) singly ionized argon species as a function of position along the propagation axis of the capillary, where zero indicates the capillary entrance. The vertical lines indicate theoretical positions of the beating pattern between the two lowest
order EH\(_{11}\) and EH\(_{12}\) modes.
Figure 10. Experimentally measured beating pattern of singly ionized argon (488 nm line) for (a) 707 µJ, (c) 777 µJ, (e) 840 µJ and theoretically calculated beating pattern of singly ionized argon for a 40 fs laser pulse with input pulse energy of (b) 707 µJ, (d) 777 µJ and (f) 840 µJ. The vertical lines indicate the positions of the gas inlet holes.

Figure 11. The (a) plasma fluorescence through a 488 nm filter and corresponding (b) XUV spectrum, for a series of pressures starting at 20 mbar (bottom) through to 160 mbar (top) in steps of 20 mbar.

is proportional to the emission, plotted as a function of argon pressure and axial position along the capillary. The theoretical model uses the same laser parameters as the experiment and allows EH_{1m} modes where m is between 1 and 30. The dominant EH_{11}–EH_{12} mode beating is clearly visible.

The theoretical model also shows a similar overall reduction in ion emission intensity along the capillary axis as a result of ionization induced losses. One obvious difference between the experiment and theory is that the expected intensity peak at 23 mm from the front of the capillary is observed experimentally at 18 mm. This offset can be explained by variation of the focusing conditions into the capillary. If the waist of the beam is not placed at the entrance to the capillary, a curved wavefront is coupled to the capillary which alters the phase of the coupled capillary modes at the input, and shifts the position of the first beat in exactly the way seen from experiment.

A subsequent data set, recorded with a throughput of 80% after optimizing the coupling conditions, shows several notable differences arising from the increase in intensity all along the capillary. Figure 11(a) shows the plasma distribution along the length of the capillary for different pressures. At very low pressure, where the atomic density is small and thus the nonlinear change in refractive index on ionization
is small, the beat pattern is similar to that seen in the low coupling efficiency case, with $EH_{11} - EH_{12}$ beats separated by 22 mm dominating. As the gas pressure is increased, the beat pattern changes significantly, indicating that nonlinear mode mixing is becoming significant, and a much more complex beat pattern appears above 40 mbar. The large peaks near to the capillary exit (beyond 50 mm) at high pressures arise not from ion emission but from the scattered laser which has been blue-shifted into the window of the bandpass filters. Figure 11(b) shows the corresponding XUV spectrum recorded simultaneously with the visible fluorescence. At low pressures, evenly spaced harmonics are observed extending down to 27 nm. As the gas pressure increases beyond 40 mbar, the XUV emission spectrum changes rapidly, as nonlinear propagation changes the laser spectrum. This nonlinear change of harmonic spectrum and the nonlinear mode coupling arise from the same optical nonlinearity, which only becomes important when the gas pressure is relatively high and the coupled intensity through the capillary is high enough for significant ionization to occur.

5. Conclusions

Spectrally resolved $Ar^*/Ar^+*$ imaging has been observed during HHG in a capillary, with the observed variation at lower power matching well with the simple model described earlier. This provides a diagnostic for several aspects of ionization important for capillary-based XUV generation, including the distribution of the laser power between the capillary modes, the effect of loss on ionization along the capillary and the nonlinear absorption and propagation inside the capillary. Changes in the plasma distribution reflect changes in the output XUV spectrum.

At higher powers there is evidence of nonlinear mode mixing which cannot be taken into account by our present model. The model is at present being extended to take into account nonlinear coupling between modes induced by plasma related phase changes, using a formulation of the problem as a multimode nonlinear Schrödinger equation. The model also includes polarization effects and high order dispersion as well as wavelength-dependent mode coupling.

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Spatially resolved Ar* and Ar+* imaging as a diagnostic for capillary based high harmonic generation

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Abstract: Spectrally resolved imaging of Ar/Ar+ created by high harmonic generation is demonstrated, and used as a diagnostic of capillary geometry on XUV generation efficiency.

Introduction

Coherent XUV radiation can be generated by the highly non-linear interaction between a gas target and high intensity ultrafast laser pulses using the high harmonic generation (HHG) process. Guiding the fundamental laser field inside a hollow capillary waveguide [1] improves phase matching and extends the potential interaction length. However, propagation of an intense pulse within a capillary waveguide filled with ionizable gas is complex, as the pulse creates a plasma, which in turn strongly affects the propagation. Previous work [2] has used emission from the excited gas to study propagation of ns pulses in capillary guides. In this work we demonstrate spectrally-resolved imaging of the plasma created by intense fs pulses within a capillary during an HHG experiment. The spectral & spatial resolution is used to separate contribution from ions and neutral species, and is an effective diagnostic for the local pressure and modal intensity variations along the waveguide.

Experimental

The laser radiation at 800nm, from a 1 kHz Ti:sapphire chirped pulse amplifier system producing 1 mJ, 35 fs pulses is focused into a 150 µm hollow capillary waveguide. The capillary is 70 mm long with two radial 300 µm holes drilled 20 mm from each end. Gas flows through the two holes to define a central region of constant gas pressure, with almost linear pressure gradients from the holes to the ends of the capillary, which were maintained at a pressure of ~ 10⁻⁵ mbar. The resulting pressure variation along the capillary was calculated using computational fluid dynamics. The capillary was imaged at right angles to the laser propagation direction onto a CCD camera using a lens (fL. 5cm). A short pass filter was used to reject scattered laser light. In addition, 420nm and 488nm narrow band pass interference filters were used to image via transitions arising from excited neutral argon and argon ions respectively.
Results and Discussion

Figure 1 shows an example of a spectrum observed and cross-sections through the images of the capillary taken with the 420nm and 488nm band pass filters, showing the intensity of the neutral (b) and ionized (c) argon species respectively. Both these images show a series of intense maxima as a function of length along the capillary. These arise from mode beating between EH$_{11}$ and higher order modes (principally EH$_{12}$) launched into the capillary. The intensity of the emission from the argon ion species is significantly reduced as the beam propagates along the capillary.

Plasma generation in the capillary is modeled by calculating, firstly, the launched mode distribution via the overlap integrals with the incident laser focal spot. This intensity distribution at the capillary entrance is used to calculate the ionization via ADK theory, and the loss due to ionization calculated as a function of radius. The resulting intensity profile is then decomposed into a new set of capillary modes which are propagated a short distance, allowing for propagation loss differences between modes, and the process repeated. The results of this calculation for a range of pressures equivalent to those measured experimentally are shown in Figure 2a. These figures show the intensity distribution at ~488nm as a function of the pressure at the inlet. The model shows the calculated ionization level for sum over four capillary modes. In addition to this comparison the model can predict the laser transmission through the capillary as a function of pressure. Agreement between the calculated and measured transmission vs. pressure is good, demonstrating the validity of the model.
The peaks in emission at the beat positions predicted from mode propagation and loss are accurate. Further adjustment within the model of the laser wavefront shape at the capillary entrance has shown even better agreement, shifting the overall position of the beat pattern by several mm along the capillary. The significant decrease in emission intensity as the laser propagates along the capillary seen in the data is mirrored by a similar decrease in the calculated ionization level along the capillary. The highly nonlinear ionization process is very sensitive to small reductions in the intensity caused by loss due to ionization and capillary loss. In contrast, emission from excited neutral Ar, shown in Figure 1(b), does not show a similar decrease along the capillary. The excitation of neutral argon by processes such as multiphoton absorption is less strongly nonlinear than ADK ionization, and so will show less variation with intensity. Thus measurement of the total emission intensity is insensitive to the variations that affect the HHG process directly, i.e. the level of ionization.

The beat period does not vary significantly along the capillary length, indicating that nonlinear mode coupling out of the EH$_{11}$ and EH$_{12}$ modes is not strong even at the highest intensities and pressures. In the 20-30nm XUV region absorption by Ar along the capillary is significant, so the presence of most of the ionization (and thus XUV generation) at the capillary entrance implies that altering the capillary design could increase the amount of XUV at the output.

## Conclusions

Spectrally resolved Ar/Ar$^+$ imaging has been observed during HHG in a capillary. The observed images are a useful diagnostic for several aspects of ionization important for capillary-based XUV generation, including the distribution of the laser power between the capillary modes, the effect of losses on ionization along the capillary and the nonlinear absorption and propagation inside the capillary. Extension of the modelling using a multimode nonlinear Schroedinger equation, including polarization effects and high-order dispersion as well as wavelength-dependent mode coupling is in progress.

C.3 Optics Express paper based on the work from Chapter 5

Modal effects on pump-pulse propagation in an Ar-filled capillary

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Abstract: Accurate three-dimensional modelling of nonlinear pulse propagation within a gas-filled capillary is essential for understanding and improving the XUV yield in high harmonic generation. We introduce both a new model based on a multimode generalized nonlinear Schrödinger equation and a novel spatio-spectral measurement technique to which the model can be compared. The theory shows excellent agreement with the measured output spectrum and the spatio-spectral measurement reveals that the model correctly predicts higher order mode contributions to spectral broadening of the pulse. Fluorescence from the excited argon is used to verify the predicted ion distribution along the capillary.

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References and links


1. Introduction

High-power ultrashort laser pulses at near-infrared wavelengths propagating in gas-filled capillaries can form a compact source of extreme ultraviolet (XUV) and soft X-ray radiation by high-harmonic generation (HHG) [1]. Maximisation of the frequency conversion efficiency
requires a detailed understanding of the radiation-atom interaction mechanism as well as the propagation properties of both the near-infrared pump in the presence of a partially ionized gas, and of the generated XUV. Previous theoretical studies of capillary-based HHG by Christov et al. [2] have used numerical solutions of the 3-dimensional version of the scalar wave equation. This work focused on understanding attosecond pulse generation [3] and quasi phasematching within waveguides [4]. Experimentally measurable predictions of these theories have typically centred on the temporal profile of the pump and the harmonics generated. In this work, we focus on understanding the spectral and spatial distribution of the pump. We introduce a numerical model of pump pulse propagation based on a multimode generalized nonlinear Schrödinger equation. This model is verified firstly by simple measurement of the total output spectrum, a commonly used procedure [5, 6] which only tests the summation over all modes. More stringent testing of the model is then demonstrated by measuring the spatio-spectral distribution of the output power in the far field, and by measuring the evolution of the ionization along the capillary length. Both of these are much more sensitive to nonlinear mode coupling than the summed spectral output.

2. Experimental configuration

The experimental setup is shown in Fig. 1. A 40 fs laser pulse centred at 790 nm, with a 1 kHz repetition rate, was coupled into a 70 mm long hollow capillary with internal diameter 150 µm using a 0.5 m plano-convex lens. The ratio of beam spot size, \( w \), to capillary radius, \( a \), was 0.64, giving optimum coupling into the lowest order, \( EH_{11} \), mode [7]. The input pulse energy was varied between 378 and 840 µJ (corresponding to an intensity range of 2.8–6.3 \( \times 10^{14} \) W cm\(^{-2} \)) by inserting reflective neutral density filters into the beam.

The central region of the capillary was filled with argon gas via a pair of 300 µm holes drilled 20 mm from either end of the 70 mm long capillary. Argon gas at pressures up to 200 mbar could be introduced into the system from a pressure-regulated supply. The capillary was mounted within a vacuum chamber kept at ~ 10\(^{-4} \) mbar. The overall gas pressure profile was calculated using computational fluid dynamics and can be approximated by a central 30 mm flat region with linear pressure gradients in the 20 mm regions at each end of the capillary.

![Fig. 1. Schematic for measuring the spatio-spectral output and argon ion fluorescence from a capillary running in a HHG regime.](image)

The capillary output is a combination of the pump laser pulse and the generated XUV. The pump laser pulse was reflected out of the vacuum chamber into a fibre-coupled spectrometer mounted on a translation stage. The 125-µm diameter fibre end was placed at the centre of the beam and translated radially outwards across the whole beam profile. The argon ion
fluorescence was imaged using the setup shown in Fig. 1, consisting of a 35 mm focal length lens imaging the capillary from the side onto a CCD. A high pass and bandpass filter were used to single out the 488 nm emission line that is emitted by singly ionised argon [8].

3. Numerical model

For the accurate theoretical description of the pump pulse propagation we developed a numerical model based on a multimode generalized nonlinear Schrödinger equation [9] which we extended to include ionization and plasma effects [5]. To this end, the electric field of the laser pulse is written as a sum over modal contributions

\[ E(r, z, t) = \frac{\sqrt{2} Z_0}{2\pi} \sum_n \frac{1}{2} F_n(r) A_n(z, t)e^{i(k_0z-n\theta)} + c.c. \]  

(1)

where \( Z_0 = \sqrt{\mu_0/\varepsilon_0} \). \( F_n(r) \) is the transverse mode function of the \( EH_{in} \) mode of the capillary [10] normalized to \( \int r dr |F_n(r)|^2 = 1 \), and \( A_n \) is the pulse envelope such that the modal power is given by \( |A_n|^2 / (2\pi) \) in units of Watt, and the temporal dependence of the carrier wave has been factored out. Note that in writing Eq. (1) we have restricted the analysis to circularly symmetric modes only, assuming a symmetric pump laser mode and preferential coupling between modes of the same symmetry [9, 11].

The modal evolution of the pulse along the capillary is then given by

\[ \frac{\partial A_n}{\partial z} = i D_n + in_z(z) k_0 \sum_{k,l,m} Q_{nklm} A_k A_l A_m^* + \frac{i}{2} k_0 \int r dr F_n^\ast(r) S(r, z, t) n_{pl}^2(r, z, t) \]

\[ \frac{1}{2} \int r dr F_n^\ast(r) S(r, z, t) \rho_0(r, z, t) W(r, z, t) U \]

\[ \frac{1}{2} \int r dr F_n^\ast(r) S(r, z, t) n_{pl}^2(r, z, t) \]

(2)

Here, \( S = \sum F_n A_n^\ast \), \( D_n \) describes modal dispersion and losses, including high-order dispersion, and the subsequent terms model the gas nonlinearity, plasma refractive index, and ionization losses of the pump, respectively. In Eq. (2), \( n_z(z) \) is the nonlinearity of the neutral Ar gas as a function of the local pressure, \( k_0 \) is the propagation constant at the pump frequency, \( Q_{nklm} = \int r dr F_k F_l F_m \) are the mode overlap integrals, \( n_{pl} \) is the refractive index of the local plasma density resulting from ionization of the neutral gas by the propagating pulse, \( \rho_0 \) and \( W \) are the neutral gas density and ionization rate, respectively, and \( U \) is the ionization potential [5, 9].

The ionization rate \( W \) is calculated using the Keldysh theory [12]. For the results presented in this work, the set of equations (2) was solved by a Fourier split-step method for typically 20 modes, which we found sufficient for an accurate description of our experimental parameters. We emphasize that this model is capable of describing the full temporal dynamics of the propagating light in three dimensions, including modal effects such as group velocity mismatch. Therefore, as will be discussed in the following sections, the model allows us to investigate detailed spatial and spectral correlations, beyond what was possible with a simpler, earlier model [8].

4. Comparison of experimental results with numerical model

The predicted and measured total output spectra for three different input powers are shown in figure 2. The theoretical and experimental spectra show excellent agreement with not only the
degree of blue shifting observed, but also the complex spectral shapes that occur at higher powers. We believe this is the first evidence that the blue-shifted shoulder of the spectrum is primarily located in higher order modes.

Figure 2 shows that at lower powers the EH$_{11}$ mode dominates the spectrum as nonlinear mode coupling is small. The model also predicts a walk-off between the modes, this is calculated to be 8 fs between the EH$_{11}$ and the EH$_{12}$ modes along the 7 cm of the capillary. As the power is increased, more energy is coupled into the EH$_{12}$ and other higher modes. The theoretical model indicates that nonlinear mode coupling is greater at the trailing edge of the pulse due to plasma defocusing, which leads to the observed blue shifted spectra in the higher order modes. Experimentally, the modal distribution cannot be directly determined from a summed spectrum of the kind shown in figure 2, although the agreement between theory and experiment allows us to infer that the modal distribution is correct.

In order to test whether the model correctly predicts the distribution of intensity in high order modes, we make use of the fact that as the modes propagate into the far field their spatial divergence is strongly dependent on their order, with higher order modes showing significantly greater angular spread. This means that the radial distribution of the spectral intensities has contributions from different modes at different radii (although the modes are not completely separated). Thus the spatio-spectral intensity distribution in the far field is very sensitive to any variation in the intensities of individual modes, and its measurement can provide a sensitive test of the ability of the numerical model to correctly predict the modal distribution.

Figure 3 shows the calculated (a) and measured (b) spectral intensity distribution as a function of radial distance from the beam axis. The theoretical intensity profiles of the EH$_{11}$ and EH$_{12}$ modes are shown for comparison. It is clear from both theoretical and experimental distributions that the differences in the intensity distributions of the modes are large. The EH$_{11}$
mode has significant contributions at around 790 and 810 nm, and the blue-shifted components at about 740 nm show the same radial profile as the EH$_{12}$ mode, with an on-axis peak, and another ~8 mm from the axis. This distribution is clear in the theoretically modelled intensity distribution. In the measured distributions, the general features of the pattern are repeated. The EH$_{11}$ mode appears at 780 and 810 nm, and the blue-shifted peak at 760 nm has the radially narrower distribution of the EH$_{12}$ mode. However, several differences are clear. The first is that more extensive off-axis blue shifting is seen in the experimental data, implying greater broadening in the higher order modes than was predicted. Secondly, an extra peak can be seen close to the centre of the beam at ~800 nm in the experimental data. The shape of the EH$_{11}$ peaks around 790-810 nm look similar to the spectral distortion caused by SPM, which produces first a splitting, and then a central peak, rising between the split peaks as the nonlinear phase shift increases. Both of these differences suggest that the nonlinear shifting of the spectrum is slightly stronger in the experiment than predicted by theory.

![Fig. 3](image_url)

While spectral measurements at the capillary exit are a good test of the end point of the model, measurement of the Ar ion fluorescence along the length of the capillary provides a test of the model along the whole propagation length. Experimentally, the fluorescence, produced by excited Ar ions created by the pump pulse, is filtered and imaged from the side. The integrated fluorescence from each point along the capillary length can be compared directly to the ionisation levels predicted by the propagation model.

The integrated argon ion fluorescence and the theoretical integrated ionisation are compared in figure 4. The measured 488 nm argon ion fluorescence should be proportional to the calculated ionisation level within the capillary. The beat positions for the EH$_{11}$ and EH$_{12}$ modes calculated for linear propagation are shown as vertical dashed lines; as expected, both experimental and calculated beat positions are clearly shifted from these positions by the effects of nonlinear propagation. The initial increase in ionisation at the capillary entrance is observed in both theory and experiment and the first two major peaks appear at approximately the same positions within the capillary. The smaller structures within these major peaks do not correlate.
The final major peak predicted by the numerical model is observed at the same point in the capillary but is significantly smaller in length and size. The discrepancy may be due to losses at the gas inlets within the capillary wall. These have been observed as increased scattering during experiments and are not included within the numerical model.

**Conclusion**

In this paper we have introduced a new method for numerically modelling nonlinear propagation within a capillary used for HHG. We have compared this model to the spectral output of the capillary and observed strong correlation. The model predicts the modal variation of the pulse shape, and the spatio-spectral measurement technique allows detailed comparison of not just the integrated spectrum, but also the individual modal contributions, because of the differences in the far-field mode patterns, validating the model and the coupling terms chosen. Understanding the modal distribution will allow modelling of XUV phase matching in the presence of nonlinear mode mixing, which is important for XUV generation at high intensities. It will also allow better understanding of the spatial profiles of the compressed pump pulses predicted by the model, providing a route to their exploitation.

**Acknowledgments**

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C.4 Ultrafast XVI and CLEO Europe posters based on work from Chapters 4 and 7

The following pages show the scaled down posters that were presented at Ultrafast Phenomena, July 2008 and CLEO Europe, May 2011. The work is based on chapters 4 and 7 and covers the comparison of argon ion fluorescence to ionisation along the capillary and CDI of a binary object from a polychromatic source respectively.
Spatially resolved Ar* and Ar+* imaging as a diagnostic for capillary based high harmonic generation

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Abstract

Spectrally-resolved Ar* and Ar+* imaging has been observed during HHG in a hollow-glass capillary as an diagnostic tool. This diagnostic technique is valuable for understanding and optimising the XUV emission. The use of the Ar+* harmonic channel in the HHG process is essential for reducing the number of ionisation stages within the harmonic generation process. The dependence of the harmonic yield on gas distribution and pressure is investigated, with particular emphasis on the influence of capillary length. A good agreement between modelled and experimental ionisation profiles is found, suggesting that the model allows for the calculation of ionisation in hollow-glass capillaries.

Introduction

Non-collinear X-ray laser systems have been of great interest for over two decades. A non-collinear X-ray laser is one that produces a spatially-separated beam of X-rays. Non-collinear X-ray laser systems are capable of operating at low peak powers and can provide a wide range of X-ray wavelengths. The advantages of non-collinear X-ray lasers include reduced interference effects and increased tunability. The use of non-collinear X-ray lasers has led to a number of applications, including medical imaging, materials processing, and molecular imaging.

Experiments

The XUV spectra were recorded using a spectrometer with a resolution of 0.1 Å and a detection limit of 10^{-7} W/Å. The XUV spectra were recorded at different gas pressures and gas distributions. The gas pressure was varied from 1 to 10 Torr, and the gas distribution was varied from 1% to 10%.

Modeling

The model used in this study is a Monte Carlo simulation of the HHG process. The model takes into account the gas distribution, the gas pressure, and the capillary length. The model also takes into account the influence of plasma related phase changes.

Conclusions

The ionisation profiles observed in this study are in good agreement with the model predictions. The model allows for the calculation of ionisation in hollow-glass capillaries.

References


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Summary
High harmonic sources have excellent potential for coherent lab-based soft X-ray microscopy, with radiation consistently able to reach the soft X-ray “water window” (e.g. 90 nm). However, traditional imaging optics below 90 nm are hard to manufacture; instead, the far-field diffraction pattern is recorded and a computational technique called “Coherent Diffraction Imaging” (CDI) is applied to reconstruct the near-field spatial frequency.

A result of frequencies is generated, i.e., ideally, multiple harmonics would be used to make full use of the flux. In this case, an incoherent sum of wavelength dependent diffraction patterns is measured at the detector—a signal that cannot be interpreted correctly by traditional methods of CDI.

A standing result is presented for monochromatic CDI applied in a diffraction pattern obtained by illuminating a test sample with multiple harmonics. 90 nm resolution was obtained, measured by the value of the highest reconstructed spatial frequency, and the reconstructed pattern confirmed that the sample was at a line focus of the illuminating beam.

Coherent Diffraction Imaging

- Technique for lensless imaging developed for use in electron diffraction by Gerchberg and Saxton.
- Uses phase information in the far-field and the recorded far-field diffraction pattern to reconstruct the near-field phase distribution.

![Image of Coherent Diffraction Imaging](image-url)

Experiment Layout

- The recorded diffraction patterns are obtained by illuminating the sample through a partially coherent source.
- The sample is mounted on a pre-focus stage for alignment and the spatial coherence properties are measured.
- The sample is then illuminated through a partially coherent source and the far-field diffraction pattern is recorded at the detector.

![Image of Experiment Layout](image-url)

References


A Test Sample

- The test sample is an electron microscope sample, a microscopy sample, or a sample of unknown composition.
- The sample is mounted on a pre-focus stage for alignment and the spatial coherence properties are measured.
- The sample is then illuminated through a partially coherent source and the far-field diffraction pattern is recorded at the detector.

![Image of A Test Sample](image-url)

A Broad Bandwidth Reconstruction with 90 nm Resolution

- The recorded diffraction patterns are obtained by illuminating the sample through a partially coherent source.
- The sample is mounted on a pre-focus stage for alignment and the spatial coherence properties are measured.
- The sample is then illuminated through a partially coherent source and the far-field diffraction pattern is recorded at the detector.

![Image of A Broad Bandwidth Reconstruction](image-url)

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90 nm resolution reconstruction from a polychromatic signal using monochromatic phase retrieval technique...
C.5 Talks


Spatially resolved ar* and ar** imaging as a diagnostic for capillary-based high harmonic generation. - Talk presented at Photon 08. August 2008

Spatio-spectral measurements of a High Harmonic Driving Pulse - Talk presented to Southern Universities Spectroscopy Group. March 2010