Controlling Laser High-Order Harmonic Generation Using Weak Counter-Propagating Light

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CONTROLLING LASER HIGH-ORDER HARMONIC GENERATION
USING WEAK COUNTER-PROPAGATING LIGHT

by

Sergei L. Voronov

A dissertation submitted to the faculty of
Brigham Young University
in partial fulfillment of the requirements for the degree of

Doctor of Philosophy

Department of Physics and Astronomy
Brigham Young University
April 2003
This dissertation has been read by each member of the following graduate committee and by majority vote has been found to be satisfactory.

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ABSTRACT

CONTROLLING LASER HIGH-ORDER HARMONIC GENERATION USING WEAK COUNTER-PROPAGATING LIGHT

Sergei L. Voronov

Department of Physics and Astronomy

Doctor of Philosophy

Laser high-order harmonic generation in the presence of relatively weak interfering light is investigated. The interfering pulses intersect the primary harmonic-generating laser pulse at the laser focus. The interfering light creates a standing intensity and phase modulation on the field, which disrupts microscopic phase matching and shuts down local high harmonic production. Suppression of the 23\textsuperscript{rd} harmonic (by two orders of magnitude) is observed when a counter-propagating interfering pulse of light is introduced. A sequence of counter-propagating pulses can be used to shut down harmonic production in out-of-phase zones of the generating volume to achieve quasi phase matching. Harmonic emission is enhanced in this case.

A new high-power laser system with higher pulse energy has been constructed to further investigate quasi phase matching of high-order harmonics generated in difficult-to-ionize atomic gases (e.g., neon as opposed to argon). The new system can also be used
to study harmonic generation in ions. A new counter-propagating beam produces a train of 5 pulses with regulated timing. In preliminary tests, the new system has produced high harmonics up to the 65th order in neon. This should increase with additional adjustments to the laser system. The high-order harmonics have also demonstrated to be useful for polarized reflectometry measurements of optical surfaces in the extreme ultraviolet (EUV) wavelength range.
I would thank my advisor Dr. Justin B. Peatross for his support, encouragement, and teaching. Thanks to those in our research group for their support and help. Thanks to the members of my graduation committee for discussing my dissertation. I wish to thank my parents for teaching me the principles that finally helped me reach the goals. I want to thank my beautiful wife, Julia, for being loving, patient, and supportive during all the years I devoted to the study. I would like to thank my children Anton, Victoria, and Sofia for their love and understanding.
PUBLICATIONS


CURRICULUM VITAE

The author was born in Ukraine on March 11, 1966. He attended Tomsk State University, Russia, from 1983 to 1991, and graduated with a Master of Science degree in Optics and Photonics. He came to Brigham Young University in fall of 1996 and began graduate studies in the Department of Physics and Astronomy. His research work was performed under the direction of Professor Justin B. Peatross.
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CHAPTER 1

PRINCIPLES OF HIGH HARMONIC GENERATION

1.1 INTRODUCTION TO HIGH-ORDER HARMONICS

Lasers capable of generating terawatt peak powers with pulse durations on the order of few tens of femtoseconds, or even shorter, are now routinely available in small-scale laboratories. Such lasers are capable of generating light intensities corresponding to electric fields that easily exceed the fields binding valence electrons to atoms [1]. Electrons that experience these extreme conditions can radiate high-order harmonics of the stimulating laser field. Over the past decade, there has been much interest in the process of high-order harmonic generation because of many potential applications in science and technology [2,3].

P.A. Franken et al. at the University of Michigan [4] reported the first observation of harmonic generation in 1961. This was soon after the invention of the laser itself (1960) and represents the first experiment in non-linear optics. They used a pulsed ruby laser that produced electric field strength $10^5$ V/cm (wavelength 694.2 nm, pulse energy 3-joule, millisecond duration). When the laser light was focused into a non-isotropic material such as quartz, the second harmonic was emitted at 347.2 nm. The efficiency of converting the laser energy into the second harmonic was only a small fraction of a percent.

Since that time, non-linear frequency-conversion techniques have typically relied on crystalline solids as nonlinear media. For efficient conversion, the electromagnetic waves corresponding to the driving and the generated signal must be phase-matched.
That is, the fundamental and the harmonic must have the same phase velocity as they travel through the generating medium. In this case, the nonlinear polarization (described in section 1.2) adds coherently as the waves co-propagate, resulting in a quadratic increase in the signal intensity. Phase matching has generally been achieved in birefringent materials with carefully chosen field polarization orientations, so that the light of the fundamental and the harmonic frequencies travel with the same speed in the material. Otherwise, the harmonic light produced up- and down-stream do not join in phase.

The reliance on solid materials limits the application of nonlinear optical techniques to wavelengths around 200 nm or longer [5]. In order to study high-order harmonics, which have much shorter wavelengths, very high laser intensities must be used. Solid materials are not suitable for this not only because they are opaque in the extreme ultraviolet (EUV) range, but they also are unable to withstand the relatively high intensities required for high-order processes. In contrast, many gases are transparent well into EUV spectrum. Light at these shorter wavelengths propagates with only moderate absorption in low-pressure gases. In addition, gases are “self-healing” which means that extreme intensities can be used. However, because gases are isotropic, established phase-matching techniques cannot be used, and thus, phase-matched frequency conversion techniques are not well developed for the EUV and soft x-ray regions of the spectrum.

For many years, only second through fifth harmonics were studied. However, in 1987 McPherson et al. [6] from the University of Illinois at Chicago observed harmonics as high as the seventeenth order produced in neon gas. Their ultraviolet (248-nm) laser had the maximum pulse energy of 20mJ and pulse duration of 350 femtoseconds, creating
10^{15-10^{16}} \text{W/cm}^2 \text{ focused intensity. Since then, many other researchers have studied High-Order Harmonic Generation (HHG) in an effort to achieve the highest possible conversion efficiency and maximum available order, using a variety of wavelengths, pulse durations, focused intensities and different gases to generate high harmonics. Anne L’Hullier et al. [7] studied high harmonic generation using a 1-picosecond Nd:Glass laser with wavelength 1.054 microns. They observed the 29th harmonic in Xe, the 57th in Ar, and at least up to the 135th harmonic in Ne. They also investigated the influence of gas density and focal geometry on the conversion efficiency. Macklin et al. observed harmonics up to the 109th order using 125 fs, 806 nm pulses in neon [8]. Preston et al. obtained the 37th order of a 380 fs, 248 nm KrF laser from helium ions [9].

The highly nonlinear nature of the ionization process makes it possible to generate and detect harmonics up to several hundred orders, which correspond to wavelengths below 3 nm. Z. Chang et al. [10] reported such harmonics using 26 fs, 20 mJ, 800 nm laser pulses with 10^{15} \text{W/cm}^2 intensity at the focus. The pulse duration of the harmonics they produced is expected to be less than 3 fs, since the high-harmonic emission is believed to occur on the rising edge of the 26 fs pulses. At very short pulse durations, the combination of enhanced single-atom harmonic response and the tendency for reduced phase mismatches can lead to relatively efficient production of sub-femtosecond (attosecond) x-ray pulses, the output being 100 times brighter for 5 fs laser pulses than for 10 fs laser pulses [11]. Nevertheless, the efficiency of converting laser energy into potentially useful high harmonic light is typically poor (< 10^{-7} conversion efficiency of laser light into one high harmonic order). Despite the low efficiency of this process, the
photon flux generated is sufficient to be useful for some applications in time-resolved ultrafast x-ray spectroscopy.

An attractive feature of HHG is that the harmonics emerge from the generating region as coherent beams with polarization similar to that of the generating beam. The harmonics are embedded in the residual laser beam but with a typically narrower beam divergence [12-14]. Fig. 1.1 represents a typical setup to produce the high-order harmonics in a jet of gas inside a vacuum chamber.

![Typical setup for producing laser high-order harmonics in gas. The gas is injected into the focus of the laser.](image)

The low divergence of the high harmonic light may be useful for applications such as the probing of dense plasmas, EUV lithography, ultrafast time-resolved spectroscopy, EUV reflectometry or ellipsometry, or basic atomic physics research [1-3,15]. Nevertheless, the harmonics generated in atomic gases (in particular the harmonics of highest orders) have so little energy per pulse that their prospects of usefulness in science and technology are limited. Geometrical, free-electron, and intensity-dependent
phase mismatches (between the generated high harmonic light and the laser field) play a dominant and quite undesirable role in limiting the harmonic emission.

In my dissertation, I present a model and experimental verification that phase mismatches in high harmonic generation can be significantly counteracted with weak interfering light. The remainder of this chapter provides additional background information about the high-harmonic generation process and review methods for enhancing the efficiency of converting laser light into high-order harmonics. Chapter 2 describes how weak counter-propagating laser pulses can be used to disrupt harmonic generation in out-of-phase regions in the laser focus. Chapter 3 describes our first experiments in laser high-order harmonic disruption and enhancement using interfering weak light. Chapter 4 outlines the development of a compact polarized EUV reflectometer utilizing high-order harmonics. This EUV reflectometer illustrates a possible application of high harmonics. Chapter 5 describes a new laser system we constructed for the purpose of extending the phase matching technique to laser pulses with higher energy and, hence, to higher harmonic orders. The new system enables the simultaneous disruption of harmonic generation in several out-of-phase emission zones for an overall harmonic emission enhancement (quasi phase match). Chapter 6 outlines the new experimental setup for high harmonic production with multiple counter-propagating pulses. This setup will be used for future experiments that extend my dissertation work.
1.2 HARMONIC GENERATION AT THE ATOMIC LEVEL

When intense laser light interacts with an atom, the bound electrons undergo violent and non-linear motions that can radiate high harmonics of the applied frequency. The atomic potential is an even function because the medium is isotropic and centrosymmetric [16]. As a result, the harmonics generated in gases are all odd order. In the high-harmonic generation process, intense laser light with angular frequency $\omega$ passes through a non-linear medium and generates light with angular frequency $N\omega$, where $N$ is the harmonic number. This is in contrast with the common situation for low-intensity light where the polarization of the medium depends linearly on the electric field:

$$P = \varepsilon_0 \chi E, \quad (1.1)$$

where $\chi$ is the linear susceptibility. Hence, if light of angular frequency $\omega$ passes through the medium, a polarization also oscillating at $\omega$ is produced. This is only an approximation, which breaks down at high laser intensities when non-linear terms in the electric field polarization become important. The polarization is often represented by an expansion in terms of electric field:

$$P = \varepsilon_0 \chi^{(1)} E + \varepsilon_0 \chi^{(3)} E^3 + \varepsilon_0 \chi^{(5)} E^5 + \ldots \quad (1.2)$$

In this case, the polarization is composed of non-linear terms such as $\varepsilon_0 \chi^{(3)} E^3$, $\varepsilon_0 \chi^{(5)} E^5$, etc., as well as the linear term $\varepsilon_0 \chi^{(1)} E$. These elements in the polarization may oscillate at frequency $3\omega$, $5\omega$, etc., giving rise to harmonics. Because the higher order terms in the susceptibility are small compared with the first term, non-linear optical
effects were not observed [4] until the development of the laser. In order to achieve
harmonic generation, the generating laser light must be linearly polarized (or close to it).

Until the early work of McPherson et al. [6], HHG had been described within a
framework of perturbation theory [17-19]. Perturbation theory suggests that each
harmonic order depends on the order immediately below it and that each harmonic order
should be progressively much dimmer than the last. Nevertheless, high harmonics are
observed with a long “plateau” in the conversion efficiency as a function harmonic order,
where many orders have roughly the same strength. The plateau ends abruptly with a
sharp cutoff past which higher harmonics are not produced. A schematic representation
of this phenomenon is shown in Fig. 1.2 for krypton, argon and helium noble gases [7].
The occurrence of the long plateau is a departure from a major characteristic of
perturbation theory, which suggests that the strength of each harmonic order N should
scale as the input field to the Nth power. It has been shown experimentally that for
harmonics with photon energies above the ionization potential of the atom, the non-linear
polarization scales instead as the input field to an effectively lower power (e.g., ~5th) [20].

The plateau in the emission strength of different harmonic orders occurs because
each harmonic order not only depends on the previous order but also on the order after it
[21,22]. As an example, the 15th harmonic depends on the 13th and the 17th order which
are both coupled all the way back to the fundamental. The plateau of harmonic orders
decreases in intensity by typically only a factor of 2 or less between each order where in
the perturbation regime the decrease is typically several hundred times between each
successive order. This dependence on both higher and lower harmonics has a tendency to
flatten out the harmonic production up to a maximum harmonic order where a sharp cut-
off occurs. As it is seen on Fig. 1.2, the plateau region is longer for He, than for Ar or Kr, though the relative conversion efficiency is lower. The interpretation of these results is not an easy task because it involves both the single-atom response to the laser interaction and the collective coordination of many atoms, the capability of the medium to ensure proper phase matching between the incident field (the driving field) and the generated harmonics.

Fig. 1.2 A schematic representation of high harmonic emission cut-off seen in krypton, argon, and helium noble gases (taken from Ref.[7]).
Traditionally, harmonic generation is described in terms of the non-linear polarization induced in the medium by the high strength input electric field [16,23]. For low-order harmonics, this polarization, described in Eq.(1.2), comes about from an effective potential well wherein the electron resides, which varies from the parabolic shape of a purely “harmonic” system. However, very strong external electric fields change the dynamics of the interaction, especially when the field becomes comparable to the Coulomb field (on the order of $E=5*10^{11} \text{ N/C}$ in hydrogen), which binds the electron to the atom. As atomic electrons are driven strongly in a field, the potential that the
electron sees is extremely anharmonic – to the point where the electron may leave the atom and become ionized.

Fig. 1.3 depicts the potential of a commonly used model for strong field-atom interactions. The figure shows a hydrogenic atomic potential well (solid line) and the potential of an applied strong oscillating electric field \( E=E_0 \cos(\omega t) \) (dashed line) at an instant in time. The combined potentials become

\[
V(x) = -\frac{e^2}{4\pi|\lambda|\epsilon_0} - exE,
\]

(1.3)

where \( e \) is the charge of the electron, and \( \epsilon_0 \) is the permittivity constant.
We can estimate the intensity at which the laser field becomes comparable to the atomic field, in which case the electron may freely escape according to the Barrier Suppression Ionization (BSI) model [24,25], which has proven to be remarkably accurate. This provides a benchmark for the maximum intensity an atom can experience before ionizing. If the field is strong enough, the coulomb barrier falls below the initial binding energy of the electron. Fig. 1.4 shows the combined potentials of the atom and a field sufficiently strong to allow the electron can escape over the barrier.

Of course, there can be quantum tunneling before the barrier is suppressed completely to the level of the binding energy [26]. There can also be quantum wave packet reflections from the barrier even when it is suppressed below the binding potential. Nevertheless, we ignore these effects in this estimate.

To find the electric field $E$ needed to allow the electron to escape, we find the extremum of $V(x)$ in Eq.(1.3) and solve for $x$, which yields

$$x = \sqrt{\frac{e}{4\pi\varepsilon_0 E}}.$$  \hspace{1cm} (1.4)

This is the position of the highest point on the barrier. We substitute this position back into Eq. (1.3) and set the potential equal to the binding energy $-V_0$:

$$V\left[x = \sqrt{\frac{e}{4\pi\varepsilon_0 E}}\right] = -\frac{e^{-3/2}}{\sqrt{\pi\varepsilon_0}}\sqrt{E} = -V_0.$$ \hspace{1cm} (1.5)

From this, we see that the electric field needed to ionize the atom is

$$E = \frac{\pi\varepsilon_0 V_0^2}{e^3}.$$ \hspace{1cm} (1.6)

The corresponding intensity for linear polarized laser light is found to be

$$I = \frac{\varepsilon_0 c E^2}{2} = \frac{\pi^2 \varepsilon_0^3 c}{2 e^6} V_0^3.$$ \hspace{1cm} (1.7)
where \( c \) is the speed of light in vacuum. Table 1.1 shows the ionization potential in eV for neutral atoms and ion states of the most commonly used noble gases [24,27].

<table>
<thead>
<tr>
<th>Gas Species</th>
<th>Neutral</th>
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<th>II</th>
<th>III</th>
<th>IV</th>
<th>V</th>
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<tr>
<td>Helium</td>
<td>24.6</td>
<td>54.4</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Neon</td>
<td>21.6</td>
<td>41.0</td>
<td>63.4</td>
<td>97.1</td>
<td>126.2</td>
<td>157.9</td>
</tr>
<tr>
<td>Argon</td>
<td>15.8</td>
<td>27.6</td>
<td>40.7</td>
<td>59.8</td>
<td>75.0</td>
<td>91.0</td>
</tr>
<tr>
<td>Krypton</td>
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<td>37.0</td>
<td>52.5</td>
<td>64.7</td>
<td>78.5</td>
</tr>
<tr>
<td>Xenon</td>
<td>12.0</td>
<td>21.2</td>
<td>32.1</td>
<td>47.6</td>
<td>59.0</td>
<td>72.0</td>
</tr>
</tbody>
</table>

Table 1.1 Ionization potentials in eV for neutrals and ions for the noble gases.

Table 1.2 shows the ionization intensities calculated with Eq. (1.7). In order to achieve the intensities shown in the Table 1.2, a millijoule-class laser pulse with the duration of a few tens of femtoseconds need only be focused to a diameter tens of micrometers wide. The peak intensity of a laser pulse may be estimated as follows:

\[
\text{Intensity} = \frac{\text{Power}}{\text{Area}} = \frac{\text{Energy}}{\text{duration} \times \text{Area}}.
\]

(1.8)

For example, our first ultrafast laser setup at Brigham Young University designed to produce high-order harmonics utilized 0.5-millijoule, 30-fs laser pulses focused onto a spot of the radius of 25 microns (Gaussian profile). Under these conditions, the peak laser intensity of the laser in focus was approximately \( I = 2 \times 10^{15} \text{ W/cm}^2 \). According to Eq.(1.7), the electric field of the laser was on the order of \( 10^{11} \text{ N/C} \). This electric field strength is
comparable to the Coulomb field binding the electron in a hydrogen atom. Modern lasers can routinely produce electric field strengths as high as $10^3$ Volts per Angstrom (laser intensity $\sim 10^{19}$ W/cm$^2$).

<table>
<thead>
<tr>
<th>Gas Species</th>
<th>Ionization intensity, W/cm$^2$</th>
<th>Ion state I ionization intensity, W/cm$^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Helium</td>
<td>$1.5 \times 10^{15}$</td>
<td>$3.5 \times 10^{16}$</td>
</tr>
<tr>
<td>Neon</td>
<td>$8.7 \times 10^{14}$</td>
<td>$1.1 \times 10^{16}$</td>
</tr>
<tr>
<td>Argon</td>
<td>$2.5 \times 10^{14}$</td>
<td>$2.3 \times 10^{15}$</td>
</tr>
<tr>
<td>Krypton</td>
<td>$1.5 \times 10^{14}$</td>
<td>$1.4 \times 10^{15}$</td>
</tr>
<tr>
<td>Xenon</td>
<td>$8.3 \times 10^{13}$</td>
<td>$8.1 \times 10^{14}$</td>
</tr>
</tbody>
</table>

Table 1.2 Ionization intensities for some neutral noble gases and their ionization state I.

A typical high-intensity laser produces 800 nm wavelength radiation (Ti:Sapphire active medium, photon energy 1.55 eV). The ionization suggested by the above model is often referred to as multiphoton ionization. For example, $\sim 11$ photons are needed to ionize a neutral atom of argon gas.
1.3 CLASSICAL AND QUANTUM MODELS FOR HIGH HARMONIC GENERATION

A. Classical model

We now consider high harmonic emission by electrons undergoing the ionization process described in the previous section. We can apply classical kinematics to estimate the maximum energy available to an atomic electron in the applied oscillating electric field, which might be emitted as a high-harmonic photon.

Kenneth Kulander et al. [28] and Paul Corkum [29] first suggested theoretically that the maximum photon energy in harmonic generation follows the universal law:

\[ K_{\text{max}} = V_0 + 3.17U_p, \]  

where \( V_0 \) is the energy needed to ionize the atom, and \( U_p \) is the mean kinetic energy of a free electron oscillating in the laser field (no drift velocity), called the ponderomotive potential. According to the model, the electron acquires a maximum kinetic energy of \( 3.17U_p \) as it is pulled from the atom by the laser field (leaving the atom momentarily ionized) and subsequently slammed back into the atom when the polarity of the laser field reverses.

Applying classical kinematics, the equation of motion of a free (not influenced by the atomic potential) electron under the applied laser field is

\[ F = m\ddot{x} = -eE_0 \cos(\omega t + \phi), \]  

where \( m \) is the mass of the electron, \( \omega \) is the angular frequency of the laser radiation, \( E_0 \) is the electric field amplitude of the laser radiation, and \( \phi \) determines the phase of the
field at the moment the electron becomes free of the atom. The coordinate $x$ indicates the distance from the atomic center to the electron oscillating in the field.

Upon integration of Eq.(1.10), we get

$$\dot{x} = -\frac{eE_0}{\omega m} \left[ \sin(\omega t + \phi) - \sin(\phi) \right],$$

(1.11)

assuming that the particle is at rest when $t=0$. If $\phi$ is not zero, then we see from Eq.(1.11) that there will be a net drift to the motion of the electron.

The mean kinetic energy of the electron in the case of no net drift, or the ponderomotive potential, is

$$U_p = \langle K \rangle = \frac{1}{2} m \langle \dot{X}^2 \rangle = \frac{1}{2} \frac{e^2 E_0^2}{m \omega^2} \langle \sin^2(\omega t) \rangle = \frac{1}{4} \frac{e^2 E_0^2}{m \omega^2},$$

(1.12)

which is a useful energy benchmark. In terms of the intensity of an applied linearly polarized laser field, $I_0 = \varepsilon_0 c E_0^2 / 2$, the ponderomotive potential may be written as

$$U_p = \frac{e^2 I_0}{2 m \omega^2 \varepsilon_0 c},$$

(1.13)

![Fig. 1.5 Escape of an electron from the atom.](image-url)
Fig. 1.5 shows a case when $\phi$ in the Eq.(1.11) is some positive value. As the electron oscillates in the electric field, it never returns again to the nucleus because of a net drift, owing to the particular initial condition. On the other hand, if $\phi$ is negative (even slightly), the drift is such that the electron returns to the nucleus with considerable velocity. Integrating Eq.(1.11) we obtain

$$x = \frac{eE_0}{mw^2}[\cos(\omega t + \phi) - \cos(\phi)] + \frac{eE_0}{mw}t\sin(\phi),$$

(1.14)

assuming that the particle is at the origin when $t=0$. The time when the particle returns to the nucleus (i.e., to the position $x=0$) can be found by setting Eq.(1.14) equal to zero and solving the following transcendental equation for $t>0$:

$$[\cos(\omega t + \phi) - \cos(\phi)] = -\omega t\sin(\phi)$$

(1.15)

The kinetic energy of the electron upon return is

$$E_{\text{max}} = \frac{1}{2}m\dot{x} = \frac{e^2E_0^2}{2m\omega^2}[\sin(\omega t + \phi) - \sin(\phi)]^2 = 2U_{\text{p}}[\sin(\omega t + \phi) - \sin(\phi)]^2,$$

(1.16)

where the value for $\omega t$ is found from Eq.(1.15).

Only those electrons that return to the nucleus and get reabsorbed can emit high harmonics. Fig. 1.6 depicts the case when the electron returns to nucleus, which occurs if the phase $\phi$ is negative.

![Fig. 1.6 Return of an electron to the atomic center, positioned at $x=0$.](image-url)
We solve Eq.(1.15) and Eq.(1.16) simultaneously to find maximum kinetic energy of the electron upon return, Eq.(1.9). In order to find the result $E_{max} = 3.17 \ U_p$, we should plot Eq. (1.16) as a function of the phase parameter $\phi$. Fig. 1.7 shows the kinetic energy that the electron possesses upon return to the nucleus as a function of the phase of release $\phi$. As is apparent, the maximum kinetic energy $3.17 \ U_p$ occurs for $\phi = -17^\circ$.

Fig. 1.7 Maximum electron kinetic energy of $3.17U_p$ occurs for $\phi = -17^\circ$. 
Upon return to the atom, the electron can recombine, giving up not only all of its kinetic energy, but also the ionization potential of the atom in the form of a single photon. Thus, the highest possible photon energy that the electron can generate is given by Eq.(1.9). Of course, if the electron does not recombine it can re-scatter giving up part of its energy as radiation, i.e. it may produce a high harmonic photon, but not the highest possible photon energy.

If the intensity $V_0$ in Eq.(1.7) (i.e., the BSI model) is used in evaluating the $U_p$ (Eq.(1.13)), the highest harmonic photon energy can be calculated for the intensity at which the atom ionizes. Again, this is a slight over-estimate of the intensity since the electron can tunnel through the barrier at slightly lower laser intensities [24,25]. However, the results do not vary substantially from the results of more sophisticated models based on the tunneling ionization rates [27] for noble gases developed by Ammosov, Krainov, and Delone (ADK model) [30].

<table>
<thead>
<tr>
<th>Gas Species</th>
<th>Maximum harmonic order</th>
<th>Ion state I maximum harmonic order</th>
</tr>
</thead>
<tbody>
<tr>
<td>Helium</td>
<td>199</td>
<td>4295</td>
</tr>
<tr>
<td>Neon</td>
<td>119</td>
<td>1365</td>
</tr>
<tr>
<td>Argon</td>
<td>41</td>
<td>297</td>
</tr>
<tr>
<td>Krypton</td>
<td>27</td>
<td>187</td>
</tr>
<tr>
<td>Xenon</td>
<td>17</td>
<td>113</td>
</tr>
</tbody>
</table>

Table 1.3 Maximum available harmonic orders for some neutral and singly ionized noble gases.
According to Eq. (1.9), the maximum harmonic order that can be produced is

\[ N_{\text{max}} = \frac{V_0 + 3.17U_p}{\hbar \omega}, \]  

(1.17)

where \( \hbar \omega \) is the photon energy of the generating light. From Eq. (1.13) and Eq. (1.17), we can estimate the maximum harmonic order possible for 800 nm laser field. We employ the data from table 1.2 to construct table 1.3.

An increase of laser intensity increases \( U_p \) and, hence, the maximum photon energy. Thus, the gases that require more intensity to ionize are able to produce higher harmonic orders. The ionization process sets an upper limit on the intensity to which an atom can be exposed. The ionization intensity can be increased somewhat by making the laser pulses shorter, because the atom can survive to higher intensities [31].

Compared to the heavier noble gases, the electronic structure of neon and helium are more tightly bound. The heavier noble gases therefore produce fewer harmonic orders. However, they have significantly higher conversion efficiencies [7,32]. This happens because a more tightly bound initial electron wave packet (e.g., He or Ne) undergoes a larger spread before returning to recombine with the atom (see Fig. 1.6). This lessens the probability of recombination to the ground state of the atom [18].

J. Zhou et al. [32] measured the number of harmonic orders generated as a function of laser pulse duration. They found that for short pulses, 25 fs, the maximum harmonic order is much higher than that predicted using the BSI assumption because the field grows appreciably during the excursion time of the electron. For example, orders up to the 61\textsuperscript{th} were produced from neutral argon atoms (compare with order 41 in table 1.3.) The possibility of generating high-order harmonics from neutrals with much higher
maximum harmonic orders using extremely short laser pulses is the subject of theoretical and experimental investigation [11, 33-40].

Eq.(1.17) together with the BSI model gives a reasonable estimate for the maximum harmonic order. However, collective effects (phase mismatches in the harmonic emission from an ensemble of atoms) can alter this rule [31]. It is interesting to mention that the maximum harmonic order for some gas species in the Table 1.3 is on the order of hundreds, or even thousands. In practice, however, the overall emission of the highest harmonics can be quite small. Moreover, it can be difficult to reliably detect the faint signal of the highest harmonics for which mismatches can be severe. Nevertheless, this short-wavelength source has important implications technologically if it can be made sufficiently bright [1-3,15]. The search for methods to counteract phase-mismatches of high-order harmonic emission in atomic gases, especially for higher harmonic orders, is a main priority.

Using ions to generate high harmonics is a possibility since they typically have higher ionization potentials, but the problem with ions is that they necessitate the presence of free electrons, which make phase matching of high harmonic generation difficult.

B. Quantum model

To obtain a more complete understanding of high-harmonic generation, an investigation of the atomic wave function obeying the Schrödinger equation can be very instructive. Lewenstein et al. [41] developed a quantum mechanical model of high-harmonic generation in an oscillating field with photon energy much less than the atomic
binding potential. They used the adiabatic assumption, meaning that the laser intensity varies slowly with respect to an optical period. Their theoretical work corresponds well to experimental evidence [31] that the cut-off harmonic energy tends to be slightly lower than that predicted by Kulander and Corkum.

If the electric field of the laser changes direction while part of the electron wave packet escapes to the continuum states, that part of the electron wave packet can approach the nucleus and recombine to the ground state, producing a photon corresponding to the kinetic energy acquired by the electron wave packet plus the ionization potential. In the approach of Lewenstein et al., the contribution of all excited bound states is neglected, as well as the depletion of the ground state. The other basic assumptions in this model are:

1) only a single bound state and the continuum states are needed to describe the wave packet;
2) in the continuum, the electron can be treated as a free particle moving in the electric field with no effect of the atomic potential $V_{atomic}$;
3) the acceleration of the wave function position $\langle \dot{\vec{r}} \rangle$ is Fourier transformed to find the spectrum of high harmonic emission.

Lewenstein et al. showed that the classical model gives essential agreement with the quantum-mechanical approach. They verified that the maximum photon energy in harmonic generation agrees reasonably well with Eq.(1.7), but tends to exhibit a slightly lower harmonic order. In order to make a realistic comparison between the Lewenstein et al. model and experimental data [31], one has to take into account propagation effects.
1.4 PHASE MISMATCHES IN HIGH HARMONIC GENERATION

As mentioned in section 1.1, in order to have strong harmonic emission, the generating laser light and the created harmonics must travel in phase for a coherent buildup of energy. To understand phase matching, consider a wave front of the laser, which propagates in an atomic medium. The atoms along the path of the laser emit harmonics. These join with previously emitted harmonics with some kind of definite phase relationship. As the harmonics propagate with the laser field, they are joined by additional harmonics emitted by atoms “down stream”, their phases being associated with the laser wave front that creates them. Beneficial phase matching occurs for an individual harmonic order when the laser wave fronts and the harmonic wave fronts propagate in phase. However, this does not happen typically in a real process because the speeds of the incident and harmonic waves can vary.

Typically, a particular generated harmonic and the laser field (fundamental) are matched in phase over distances that are many times the wavelength of the incident light. However, the length over which the harmonic phases are matched (called “the coherence length”) can be much shorter than the focal depth (typically, 1 mm) owing to different rates of diffraction for different wavelengths. Fig. 1.8 shows a particular harmonic produced at different positions in the focus, propagating over many wavelengths. After a coherence length, new harmonic production is out of phase with previously generated harmonic light. Fig. 1.9 indicates how the production of a particular harmonic can go in and out of phase many times in the laser focus.
Fig. 1.8 A particular harmonic produced from atoms at different locations in the laser focus. Harmonic light generated at the first position is out of phase with the harmonic light generated at the second position, but in phase with the light generated at the third position.

Phase mismatches can cause severe destructive interference for harmonic light emerging from different locations in the laser focus. The efficiency of converting the
fundamental light into high-order harmonics is limited by several phase mismatch phenomena:

1) geometrical phase mismatches arise from discrepancies between the diffraction rates for the laser and for individual harmonics;

2) the refractive index for the fundamental laser light differs from the refractive index for the shorter-wavelength harmonics (this occurs mainly due to free electrons);

3) the intrinsic phase of harmonic emission can vary longitudinally and radially throughout the focus because of varied atomic response to the local laser intensity.

Obviously, to produce the brightest possible high harmonic light, we must mitigate phase-mismatches in the generation volume. This is a separate issue from the possibility of re-absorption of harmonic light within the relatively dense generating medium. We should keep in mind that phase issues do not alter the single-atom response to the laser field, which is the primary driver of the harmonic generation process.

A. Plasma phase mismatches

Since the strongest harmonic emission occurs as atoms undergo ionization by the laser field, we are often interested in harmonic generation in the presence of plasma consisting of free electrons and their parent ions together with the remaining intact neutral atoms. Under typical high-harmonic generation conditions [42], if the gas becomes fully ionized, the appearance of free electrons associated with ionization causes the length over which the harmonic phase slips by \( \pi \) relative to the phase of new harmonic production to be as little as few tens of microns.

The index of refraction for plasma is given by
\[ n_{\text{plasma}}(\omega) = \sqrt{1 - \frac{\omega_p^2}{\omega^2}} , \quad (1.18) \]

where \( \omega \) is the laser frequency and \( \omega_p \) is the plasma frequency. The plasma frequency is given by

\[ \omega_p^2 = \frac{N e^2}{\varepsilon_o m} , \quad (1.19) \]

where \( N \) is the density of free electrons. The index of refraction in plasma is less than unity for all light traveling through the plasma and depends on the frequency of the traveling light. Eq. (1.19) can be approximated for the case of high harmonic generation as

\[ n_{\text{plasma}}(\omega) \equiv 1 - \frac{\omega_p^2}{2\omega^2} , \quad (1.20) \]

since the laser frequency \( \omega \) is typically far above the plasma frequency \( \omega_p \).

The coherence length \( L_c \) for generating the \( q^{th} \) harmonic in a medium with frequency-dependent refractive index is found from

\[ \left[qk - k_q\right]L_c = \frac{2\pi q}{\lambda_o} \left[n_{\text{plasma}}(\omega) - n_{\text{plasma}}(q\omega)\right]L_c = \pi . \quad (1.21) \]

When the refractive index for plasma, Eq. (1.20), is used in this formula, we find

\[ L_c \equiv \frac{\lambda_o \omega^3}{q\omega_p^2} \equiv \frac{4\pi^2 m}{q\lambda_o N e^2 \mu_o} , \quad (1.22) \]

where we have assumed that \( q > > 1 \). As an example, if one electron per atom is ionized in 1 torr of gas (initially at room temperature, i.e., \( N \equiv 3 \times 10^{22} \ \text{m}^{-3} \)), the coherence length is found to be \( L_c \ (\text{cm}) \equiv 4/q \). Thus, the coherence length for the 39\textsuperscript{th} harmonic in this case is 100\textmu m, and is ten times shorter if the pressure is increased to 10torr. Plasma phase-
mismatches become of greater importance (i.e., the coherence length gets shorter) with increased harmonic order. This means that a higher order harmonic goes in and out of phase with the fundamental field more often.

Eq.(1.22) shows that the coherence length can decrease as gas density increases. This means that the harmonic signal does not necessarily improve merely because the density of harmonic emitters increases [43]. The conversion efficiency can also be limited by the defocusing of the laser light if the on-axis density of free electrons becomes too large. The density of free electrons depends on the level of ionization. For lower order harmonics (e.g., orders in the teens or less), which can be generated with little ionization, the phase mismatch due to plasma is of relatively minor importance. For very short pump pulses (tens of femtoseconds and less), ionization is delayed until after significant amounts of harmonic radiation are emitted. Therefore, if the pressure is not too high, the coherence length in this case can approach that of a neutral gas.

B. Geometrical phase mismatches

Geometrical phase mismatches arise primarily from the Gouy shift \( \tan^{-1} \frac{z}{z_o} \), which gives the phase deviation between a laser wave front that undergoes focusing compared to the phase front of a plane wave (see Ref.[18], p. 486). Here, \( z_o \) is the laser Rayleigh range, \( z_o = \frac{\pi \omega_0^2}{\lambda} \), and \( z \) is the distance from the laser focus along the axis of laser beam propagation, and \( \omega_0 \) is the beam waist radius. The phase for the \( q^{th} \) harmonic order at each point where it is produced along \( z \) is \( \phi = q \tan^{-1} \frac{z}{z_o} \). The Gouy shift for the harmonics themselves as they propagate is negligible compared to this shift, which is \( q \) times the laser Gouy shift [29,44]. We can estimate the distance over which the phase \( \phi \)
changes by $\pi$. For this purpose, we take the derivative of this phase term:
\[
d\phi/dz = q\left(z_0 + z^2/z_0\right).
\]
This implies a phase change of $\Delta\phi = \pi$ over an approximate coherence length of
\[
\Delta z = \frac{\pi (z_0 + z^2/z_0)}{z_0},
\]
or simply $\Delta z = \pi z_0/q$ near the center of the laser focus.

For a typical f/50 geometry with an 800nm wavelength, the Rayleigh range is about $z_0=2.5\text{mm}$. In this case, the coherence length for the harmonic number $q=39$ turns out to be 200 $\mu\text{m}$ near the center of the focus. According to Eq.(1.23), the coherence length becomes somewhat longer if evaluated away from the focus. To reduce geometrical phase mismatches, the depth of focus can be increased relative to the gas jet width to confine the interaction region to a single geometrical coherence length, thus avoiding destructive phase cancellations [44,45]. However, loose focusing requires greater laser pulse energy to achieve the same intensity. Unfortunately, the high-intensity regions in the beam before and after the single coherence length cannot be utilized in this case.

C. Intensity-dependent phase mismatches

The intensity-dependent phase of single-atom emission of a particular harmonic can have macroscopic effects since different atoms throughout the focus experience different intensities. The laser intensity can significantly vary along its axis within the gas medium. The intensity-dependent phases can therefore vary axially as well. This can lead
to phase mismatches and deterioration of harmonic emission. This has been demonstrated experimentally [46,47] and is in agreement with theoretical expectations [48]. It is possible to find situations in which the intrinsic phases can be played against geometrical phase mismatches for a modest improvement to overall phase matching [46,47]. Our group has also made attempts to learn more about the nature of intrinsic phase mismatches [49].

1.5 PHASE-MATCHING IN HIGH HARMONIC GENERATION

Since high harmonics were discovered in 1987 [6], a basic understanding of the phase matching for high harmonics generated in gases has emerged. Wahlstrom et al. [40] and Altucci et al. [50] investigated harmonic generation as a function of atomic density. They found well-defined limitations to the gas density depending on harmonic order and atomic species beyond which index-related phase mismatches overcame the benefit of having an increased number of harmonic emitters.

L’Huillier et al. [44,45] investigated extensively the phase matching limitations associated with focal geometry. Because of these issues, it is typical to confine the interaction region to a single geometrical coherence length to avoid destructive phase cancellations (geometrical only). This is usually done by lengthening the laser focus relative to the width of the gas jet or by working outside of the focus. As mentioned above, this method does not utilize the high-intensity focal volume outside of the coherence length.

A remarkable approach for improving the conversion efficiency of high-order harmonic generation has been demonstrated by the group headed by Margaret Murnane.
and Henry Kapteyn. By generating the high harmonics inside of a hollow core fiber, they reached harmonic conversion efficiencies beyond $10^{-5}$, owing to a greatly extended coherence length in the wave-guide [51,52].

Hergott et al. [53] demonstrated relatively high conversion efficiency for harmonic orders 23-31 using a long focal length lens (~2 meters). They increased the energy of high harmonics by at least one order of magnitude (achieving microjoule harmonic energies). In the experiments, Hergott et al. controlled the diameter of the laser aperture placed before the focusing lens, interaction length, pressure, and position of the gas medium relative to the laser focus [44,45].

Braun et al. proposed a phase-matching technique [54] that relies on a self-channeling effect of the laser in gas near atmospheric pressure with laser intensities near $10^{14}$ W/cm$^2$. A single filament (a self-focused beam) can form with pulse energy as high as 30 mJ. A limitation of this self-channeling technique is its interaction length. When the laser intensity exceeds $10^{14}$ W/cm$^2$, which is typically required to produce high-order harmonics, self-focusing is no longer in balance with plasma defocusing. Nevertheless, the confocal parameter can elongate to 10 times that in vacuum [55], leading to improved phase matching for some of the lower high harmonic orders [55,56].

More recently, Tamaki et al. [57] demonstrated phase matching of high harmonics via self-focusing of the laser in a low-density gas. They observed elongated focal lengths up to 7 mm. The high-order harmonics appeared to be emitted from all the gas volume within this length. The coherence length for harmonic emission was reported experimentally to be similar in length (few mm). This improved the conversion efficiency of the 49th harmonic of 800 nm light by 40 times. The scheme enhances the conversion
efficiencies in the cut-off region to as much as $\sim 10^{-6}$ (i.e., up to two orders of magnitude enhancement for the harmonic orders in upper forties). Tamaki and coworkers used a gas cell with two 300-micrometer diameter pinholes, isolating the gas from the ambient vacuum chamber. Pinholes were used to transmit the focusing laser beam into and out of the cell. The cell was pressurized with neon atomic gas at 50 torr and the ambient vacuum chamber pressure was held to below a millitorr. Their approach is now in wide use due to its greatly reduced geometrical phase mismatch. On the other hand, plasma mismatches can become significant for this increased propagation (interaction) length.

An important limitation to the enhancement of high harmonic generation is the re-absorption of harmonic light within the generating medium reported by E. Constant et al. [58]. This limitation is the same for both the case of a loosely focused beam or a beam guided in a gas-filled hollow-core fiber or cell. Under optimal conditions, they measured the conversion efficiency of $4 \times 10^{-5}$ (two orders of magnitude enhancement) for the $15^{th}$ harmonic generated in xenon gas using a 40 fs laser pulse of wavelength 800 nm. They demonstrated sufficiently good phase matching that the output was re-absorption limited.

Aside from issues of phase matching, it is worth noting that shorter laser pulse durations lead to increased conversion efficiency because strong ionization occurs only after significant amounts of harmonic radiation is emitted. [11,59]. Special laser pulse temporal profiles can be designed to enhance the atomic response [60]. An improvement to harmonic emission has been demonstrated using a genetic algorithm to adjust the computer-controlled pulse temporal profile [61]. The inter-play between intrinsic and geometrical phase mismatches may play a role for conversion efficiency enhancement [62]. Another scheme that has been implemented produces harmonics from two
superimposed laser foci. The foci are separated longitudinally by a few millimeters. This scheme enhances the phase matching for an improved output [63].

The enhancement of the conversion efficiency is especially important for high-order harmonic wavelengths between 2 and 5 nm, in the so-called “water window” between the carbon and oxygen absorption edges where carbon absorbs and water transmits, permitting the possibility of high-resolution biological imaging. Promising work by Z. Chang et al. [10] shows that high harmonic generation at 2.7 nm may be useful for biological experiments.

In summary, the efficiency of converting laser light into the high harmonics can be enhanced through various methods. However, the high harmonics remain several orders of magnitude dimmer than other EUV sources such as a capillary discharge sources [64] producing 7 mJ pulses of spatially incoherent radiation in a 0.3 nm bandwidth centered at 13.4 nm (comparatively long duration). This wavelength corresponds to high harmonic emission of the order 59. There is no particular method yet that can enhance the HHG conversion efficiency to become comparable with the brightest plasma sources, but there is room for optimism. The quasi phase matching presented in this dissertation is one method that together with other approaches may bring high harmonics into direct competition with plasma sources in terms of absolute yield. Quasi phase matching of HHG with weak interfering light may prove particularly useful for the very high order harmonics (on the order of fifties or higher) produced in neutral gases or generated from ions where phase mismatches can be very severe and difficult to control [50,65].
CHAPTER 2
QUASI PHASE MATCHING WITH WEAK INTERFERING LIGHT

2.1 INTRODUCTION

Shkolnikov and Kaplan [66,67] proposed that if the atomic density of the medium is modulated along the axis of the laser, it might be possible to eliminate harmonic emission from those regions that have destructive phase for an overall dramatic improvement to the efficiency of harmonic production.

Fig. 2.1 Modulated gas density distribution to eliminate harmonic emission from regions with undesirable phase.

The approach, known as quasi phase matching, eliminates harmonic emission from those regions having destructive phase (see Fig. 2.1 and compare with Fig. 1.9). The scheme is
only able to counteract geometric phase mismatches, and does not deal with index phase and intrinsic phase mismatches. In addition, creating such a modulated gas distribution presents significant experimental challenges.

Fig. 2.2 Elimination of harmonic production in (a) the “out-of-phase” regions using (b) a train of counter-propagating pulses. Weak counter-propagating pulses collide with the forward harmonic generating pulse to frustrate emission from undesirable phase zones; (c) harmonic emission from the gas atoms in the remaining zones then constructively interfere.

We propose a new approach for achieving quasi phase matching in a focused laser beam. Rather than attempting to modulate spatially either the density of the atomic
medium or the density of free electrons, we use counter-propagating light to disrupt harmonic emission in specific regions of the focus. In this chapter, we analyze how even relatively weak counter-propagating light can seriously disrupt phase matching at the microscopic level [68].

To achieve quasi phase matching, multiple counter propagating light pulses can be made to collide with the main generating pulse throughout the laser focus, as depicted in Fig. 2.2. The timing and duration of these counter-propagating pulses can be chosen appropriately to frustrate harmonic production in regions with undesirable phase. The emission from the remaining regions then constructively interferes.

![Image of colliding beams](image)

Fig. 2.3 Schematic of experiment on colliding beams (based on the proposal from Ref.[37]).

As discussed in Chapter 1, the geometrical phase mismatches arise primarily from the Gouy shift in a Gaussian laser focus. According to Eq.(1.23), there can be a number
of harmonic (coherent) phase zones along the focus, as depicted in the upper part of Fig. 2.2. The quasi phase matching technique that we propose, shown in Fig. 2.3, can address this problem by directing a series of counter-propagating light pulses into the focus while enabling detection of the high harmonics.

As described by J. Peatross et al. [48], when an annular beam is focused, the center fills in to produce a central peak similar to that of a usual laser focus surrounded by faint rings. The nonlinearity of high-order harmonic generation concentrates the harmonic production in the central peak, and the harmonics emerge as usual, centered along the laser axis so that much of their energy lies inside of the original hole. The holes are necessary because of the lack of suitable materials able to transmit extreme ultraviolet light (EUV).

This approach is fundamentally different from other proposed phase-matching methods using dual laser sources. In other scenarios, either difference-frequency mixing [67] or angular tuning of crossing beams [69] is proposed to allow for phase matching in a positive dispersive medium (i.e., in plasma).

2.2 MICROSCOPIC PHASE DISRUPTION IN SUPERIMPOSED COUNTER-PROPAGATING FIELDS

Consider two plane-wave fields with real amplitudes \( E_1 \) and \( E_2 \) which propagate in opposite directions towards each other. We will assume that the \( E_1 \) is the larger of the fields. We will write the sum of the two fields as a single field having the form of the larger plane wave:

\[
E_1 e^{i(kz-\omega t)} + E_2 e^{-i(kz+\omega t)} = E_1(z) e^{i(kz-\omega t+\phi(z))}
\]  

(2.1)
The total amplitude, which varies spatially, is found to be

\[
E(z) = E_1 \sqrt{1 + \left( \frac{E_2}{E_1} \right)^2 + 2 \frac{E_2}{E_1} \cos 2kz} .
\]  

(2.2)

The phase modulation, which also varies spatially, is found to be

\[
\phi(z) = -\tan^{-1} \left( \frac{\frac{E_2}{E_1} \sin 2kz}{1 + \frac{E_2}{E_1} \cos 2kz} \right) .
\]  

(2.3)

In the limit of a very weak counter-propagating field (i.e., \( E_2 << E_1 \)), the total field reduces to that of the larger plane wave (i.e., \( E_2(z) \rightarrow E_1 \) and \( \phi(z) \rightarrow 0 \)). Nevertheless, even a relatively weak field \( E_2 \) can cause a strong standing intensity modulation. The fringe visibility for the standing intensity modulation is given by

\[
V = \frac{I_{\text{max}} - I_{\text{min}}}{I_{\text{max}} + I_{\text{min}}} = \frac{2E_2/E_1}{1 + (E_2/E_1)^2} .
\]  

(2.4)

If \( E_2 \) is only a tenth as strong as \( E_1 \) (i.e., one hundred times less intense), the standing intensity pattern has fringe visibility of 0.20. In this case, \( I_{\text{max}} \) is 49% more intense than \( I_{\text{min}} \). Fig. 2.4 represents the standing intensity profile for this case. By intensity, we do not refer to net energy propagation (i.e., Poynting vector) but rather to the square of the combined field (i.e., \( \varepsilon_0 cE_t^2(z)/2 \)). Fig. 2.4 also shows the standing phase variation described by Eq. (2.3) as well as the relative intensity \( (E_2(z)/E_1)^2 \) and phase \( \phi(z) \) in the case that the weak field is 100 times less intense (i.e., \( E_2/E_1 = 0.1 \)). The standing phase variation fluctuates over a total range of \( 0.06\pi \), which translates into a phase variation of more than \( \pi \) for harmonics beyond the 15th order. As seen in Fig. 2.4,
both the standing intensity modulation and the standing phase modulation are periodic over a half fundamental laser wavelength.

![Graph of standing intensity and phase variations](image)

**Fig. 2.4** Standing intensity and phase variations resulting when a plane wave is met by a counter-propagating plane wave one hundredth as intense.

The phase variation within the half wavelength interval can seriously dampen harmonic production. It is useful to perform a microscopic phase-matching calculation over this interval, say from $-\lambda/2$ to zero, to find the effective harmonic emission strength to be compared with the case of no counter-propagating light (i.e., $E_z=0$). To perform this calculation, we invoke a simple model: The strength of the $q^{th}$ harmonic is assumed
to follow the laser field strength raised to the $p^{th}$ power [22]. That is, the emission from individual atoms within the interval is proportional to

$$E_q(z,t) \sim E_i^p(z)e^{iq(kz-\omega t+\phi(z))}.$$  \hspace{1cm} (2.5)

Fig. 2.5 Accumulated harmonic field produced from gas atoms in an interval of a half wavelength and exiting this interval at time $t$.

The effective emission from the interval is found by summing up the contributions from all participating dipoles to the net field leaving the interval (i.e., at $z=0$). Fig. 2.5 represents a medium emitting a particular harmonic in a half-wavelength interval. To obtain the net harmonic emission in the direction of the strong beam, one integrates (along $z$) the emission with the time argument adjusted appropriately for a traveling wave. The net field at the edge of the interval is proportional to
\[ \int_{-\lambda/2}^{0} E_q(z, t + z/c)dz \sim e^{-i\phi} \int_{-\lambda/2}^{0} E_1^p(z)e^{iq\phi(z)}dz. \] (2.6)

Notice that the integral contains the standing phase factor \( \exp[\text{i}q\phi(z)] \), which causes the argument to be oscillatory. We may compare the result of Eq. (2.6) with \( \lambda E_1^p / 2 \) which is the result when \( E_2 \) is zero. We will call this comparison the effective emission. By this, we mean the apparent response of the atoms in the interval as though the phase matching had not been disrupted.

![Fig. 2.6](image.png)

Fig. 2.6 The effective emission from a microscopic phase-matching interval for 11\textsuperscript{th}, 31\textsuperscript{st}, and 51\textsuperscript{st} harmonics as a function of relative counter-propagating field strength. The calculation assumes that harmonic emission follows a power law with \( p=5 \).
We define a microscopic phase-matching factor as

\[
\xi = \frac{2}{\lambda E_1^p} \int_{-\frac{\lambda}{2}}^{0} E^p_i(z) e^{i\phi(z)} dz.
\] (2.7)

Note that \( \xi \) is real because \( E_i(z) \) is even and \( \phi(z) \) is odd over the interval of integration.

Fig. 2.6 shows the behavior of the effectiveness factor \( \xi \) for a variety of harmonic orders as a function of counter-propagating field. In all cases, \( p \) is chosen to be equal 5. As is evident, the emitted harmonics are significantly degraded even with a modest counter-propagating field. For harmonic orders in the twenties or higher, a counter-propagating field strength of only one tenth the main field is sufficient to seriously disrupt the harmonic emission, owing to the standing phase factor \( \exp[iq\phi(z)] \) in Eq.(2.6). This means that to realize this situation experimentally, counter-propagating pulses need only have one hundredth the intensity of the main pulse. As is also evident in Fig. 2.6, the effectiveness of emission is zero only at a few specific counter-propagating field strengths. However, where the effectiveness goes negative, the emission is out of phase by \( \pi \). Since the reason for suppressing the harmonic production is to eliminate emission having the incorrect phase, the \( \pi \) phase change is actually more beneficial than if the effectiveness factor were zero at the corresponding counter-propagating field values. It should be noted that the decrease in the microscopic factor with counter-propagating field strength is entirely due to the functional form of \( \phi(z) \); if \( \phi(z) \) is arbitrarily set to zero, the result of Eq.(2.7) continually increases above one, owing to the standing intensity modulation.

In the next chapter, we show experimentally that the microscopic phase mismatch in colliding pulses does indeed effectively extinguish high harmonic
generation. We also demonstrate an overall enhancement to harmonic production when counter-propagating light disrupts a single out-of-phase zone in the focus. The eventual goal of our work is to produce a series of weak counter-propagating pulses with appropriate timing and durations to suppress the emission in many regions in a laser focus that have undesirable phase. Assuming good harmonic suppression in the “out-of-phase” regions, overall harmonic production is expected to increase in proportion to the square of the number of suppressed regions, since the harmonic power is proportional to the square of the electric field. Progress towards these experiments is described in Chapter 6.
CHAPTER 3
EXPERIMENTS ON HIGH HARMONIC GENERATION IN INTERFERING LIGHT

3.1 LASER SYSTEM

The laser for the experiments described in this chapter is a chirped pulse amplification system. A more detailed description of the laser setup is given in the Masters Thesis of Ian Kohl [70]. I helped in the construction the laser system and in the optimization of its performance for experiments. Further discussion of the chirped pulse amplification techniques is addressed in Chapter 5, where the construction of a larger laser system is described in detail.

Fig. 3.1 Block-diagram of the laser setup.
An overview of the laser setup is shown on Fig. 3.1. Short pulses (p-polarized) originate in the oscillator, following the design published by Henry Kapteyn and Margaret Murnane [71]. The beam consists of a train of 18 femtosecond pulses separated by 11 nanoseconds (the repetition rate of 90.9 MHz), which corresponds to the round-trip time in the cavity. The oscillator is a passively mode-locked Ti:Sapphire cavity which uses a Spectra Physics “Millenium” laser (wavelength 532 nm) as the pump source. Fig. 3.2 shows a schematic of the oscillator, stretcher, pulse selector and amplifier portions of the laser setup [72].

![Laser system schematic](image)
The pulses are stretched in duration to avoid pulse intensities that may damage the amplifier crystal and the other optical elements in the system. The pulses enter the stretcher, which expands the 18fs pulses to about 70ps duration. The pulses become negatively chirped (see Chapter 5). In our setup, the direction of p-polarization is parallel to the optical table surface. The pulses proceed to the pulse picker, which consists of a Quantum Technology Pockels cell #1 with two crossed polarizers on its entrance and exit. This Pockels cell selects pulses for amplification and injects them into the amplifier stage.

Pockels cell #1 selects a pulse for amplification at a repetition rate of 1 kHz. This repetition rate matches the rate of available pump energy from a 1 kHz Nd:YAG laser (Lee Lasers, series 800). As shown on Fig. 3.2, the train enters the pulse picker through a polarizer transmitting p-polarized light only. When activated, Pockels cell #1 energizes for only 6 nanoseconds and rotates the polarization of a single pulse to s-polarized. A second polarizer transmits only s-polarized light (the “seed” pulse). This selected pulse encounters another polarizer, which reflects s-polarized light and transmits p-polarized light. A 40 cm focal length lens is used to match the size of the “seed” beam to the most effective amplifier “mode,” which is determined experimentally. The pulse encounters Pockels cell #2, which rotates the polarization back to p-orientation. The role of the second Pockels cell is twofold: it acts as an isolator to prevent premature amplifier lasing as well as a means of capturing the “seed” pulse for amplification. It also provides an avenue for pulses to exit the multi-pass amplifier system.

The lasing in the cavity occurs for p-polarized light, which matches the Brewster cut of the amplifying Ti:Sapphire crystal. It transmits p-polarized light with nearly 100%
efficiency but reflects about 26% of s-polarized light on each side. The beam in the amplifier travels along a counter-clockwise spiral-like progression for 6 passes through the amplifier crystal (see top view on the Fig. 3.2). Then, it gets retro-reflected to make 6 more amplification passes in the reverse direction until it re-encounters Pockels cell #2. Since this Pockels cell is no longer energized, the p-polarized amplified pulses pass freely through it and also through the subsequent polarizer. The measured average power of the beam lies in the range between 0.65 and 0.90 W. This wide spread of amplified power occurs mainly because of necessity of frequently adjusting the laser system alignment. The Ti:Sapphire crystal is cooled to near 0-degrees Celsius, which reduces thermal lensing effects in the Ti:Sapphire amplifying crystal.

The pump energy for the Ti:Sapphire crystal comes from two beams with combined energy ~15 mJ per pulse. The pump beams are focused from either side of the Ti:Sapphire crystal to a common area of diameter ~600 \( \mu \)m Full Width at Half Maximum (FWHM), where they overlap the aligned “seed” pulse injected into the amplifier. The pump pulses are created in an intra-cavity frequency-doubling LBO crystal that produces the second harmonic (532 nm) of the 1064 nm light that lases in the pump laser. The LBO crystal phase matching is dependent on its temperature stability. If the temperature changes by ~1° C, the LBO crystal performance drops ~30%. An external cooler is used to keep the crystal temperature to within the limits of ±0.5° C. Since the amplifier gain after the total of 12 passes through the Ti:Sapphire crystal is on the order of a few million, even a small change in the pump power is followed by a significant change in the
amplified signal. This inherent instability posed some challenges while running experiments, but with patience it was possible to proceed.

3.2 HARMONICS PRODUCTION AND DETECTION SETUPS

For our experiment, we required two laser pulses, a short forward propagating and a longer counter-propagating pulse. Fig. 3.3 shows the schematic of the compression stage of the laser amplifier together with the setup used for harmonic production and detection. After the beam exits the amplifier, it is expanded up to few millimeters in diameter; hence, we significantly reduce its intensity. The beam encounters a 50% beam splitter just before the compression stage. The compression stage consists of two independent parallel grating modules, one to achieve the shortest possible duration of the forward pulse and a second for creating a somewhat longer pulse (typically, 1ps) for the counter-propagating beam. The pulses are sent side-by-side through the compression grating pairs. A delay stage controls the relative arrival timing of the two pulses. The two beams enter the experimental chamber from different directions. The forward (main) strong beam generates high harmonics, and the other beam provides the weak counter-propagating light. The shortest measured duration of the forward pulse that is achieved with this laser system is 30 fs (FWHM).
Fig. 3.3 Compression, harmonics generation (interaction), and detection chambers.

We used a standard auto-correlation technique to measure the forward pulse duration and a cross-correlation technique to measure the counter-propagating pulse duration relative to that of the short forward pulse. For this purpose, it was necessary to disrupt the experiment and divert the beams to the auto- and cross-correlation setups. A more detailed description of the correlation technique is given in Chapter 6.

The experimental chamber consists of a vacuum-tight stainless steel chamber body with attached Varian 500 l/s turbo pump and scroll roughing pump. The laser beams enter through ports. In the case of the forward beam, a 30 cm focal-length lens functions as the vacuum window. The other beam enters the chamber through a flat 800nm AR-coated glass window; the beam passes through a 30cm lens inside the chamber. A gas jet
is placed on an adjustable arm that can move in three dimensions. A CCD camera and a flat mirror enable simultaneous viewing of the colliding beams from the perpendicular direction and at an angle at \(~45^\circ\) from above.

The counter-propagating light is directed to the intersection area through a 30-cm focal-length lens and a flat mirror with a 2.5 mm diameter hole drilled through. The hole provides an avenue for high harmonics emerging in vacuum to be measured while allowing some fraction of counter-propagating beam energy to be reflected towards the focus. The hole is necessary because of the lack of suitable materials able to transmit high harmonic (EUV) light.

The spatial alignment of two colliding pulses is crucial since in order to have the superposition of the beams required for the experiment, the beams must be collinear [73]. Each beam waist is on the order of few tens of micrometers. Fig. 3.4 shows the measured focal dimensions of both beams.

![Fig. 3.4 Focal measurements of the (a) forward-propagating beam and (b) the counter-propagating beam.](image)

The counter-propagating light gets reflected from the center of a flat mirror 25 cm before the focus (5 cm after the focusing lens). The beam’s diameter is 6.3 mm when it
hits the mirror. The mirror has a 2.5 mm hole drilled through its center, which cuts a portion of the beam’s energy. Even though the counter-propagating beam loses its central part to the mirror with the hole, its focused profile is similar to that of the forward beam, as is evident in Fig. 3.4. The hole allows about half of the counter-propagating pulse energy to be directed towards the laser focus.

Fine spatial adjustment of the counter-propagating beam to the forward beam is accomplished with the mirror labeled M in Fig. 3.3. A tilt to this mirror in either dimension moves the beam focus perpendicular to the direction of propagation. For our experimental setup, the foci become misaligned (i.e. the counter-propagating beam’s focus moves laterally by ~15µm), when the knob on mirror M is rotated by only 3 degrees. This makes the alignment difficult and tedious albeit possible.

The jet is approximately 300 µm wide and is backed with argon gas with pressure between 20 to 150 torr. In the experiments, the duration of the chirped, weak counter-propagating pulse was set to approximately 30 times longer than that of the forward pulse. Owing to the duration of the counter-propagating pulse, the two pulses encounter each other in a region with width similar to the thickness of the gas jet (300µm). Since both beams travel with the speed of light in opposite directions near the gas jet, the interaction region length is effectively half the length corresponding to the duration of the counter-propagating pulse. The useful duration of the counter-propagating pulse is about twice the FWHM, so the effective interaction distance is about \( t_{ps} / c = 300\mu m \), where \( c \) is the speed of light in vacuum.

Of critical importance is the temporal alignment between the two beams since they must arrive to the gas jet at the same time. This short interaction length (sub-
millimeter) creates significant challenges in the temporal alignment. Prior to an experiment, we back-fill the chamber with air and temporally re-compress the duration of counter-propagating pulse so that it ionizes a track through the air similar to the track made by the forward pulse. Rayleigh scattering from the resulting free electrons is observed with a CCD camera. As shown in the Fig. 3.5, a distinctively higher amount of Rayleigh scattering occurs from the region where the two pulses collide. From this observation, the timing of the two beams can be set.

![Image of Rayleigh scattering](image)

**Fig. 3.5 Appearance of excess Rayleigh scattering (a bright dot) from colliding pulses in air.**

We explored different search techniques to find the intersection point of the colliding pulses. It was necessary to adjust the delay stage and mirror M of Fig. 3.3 in an iterative search. The Rayleigh scattering from the collision area is monitored
simultaneously in two dimensions with a CCD camera connected to a TV monitor. This assures both spatial and temporal alignment of the two pulses. As the chamber is evacuated, the excess Rayleigh scattering from the collision point can also be observed in the gas jet when backed with the gas pressure of a hundred torr or more. The increased scattering is due to enhanced ionization, owing to the interference between the two pulses. After alignment, the counter-propagating pulse is chirped to the necessary pulse duration. Since this requires the distance between grating pair in the compressor to be changed, the arrival time of the pulse must be simultaneously adjusted using the delay stage. In this way we guarantee that the pulses still meet in the gas jet.

A distinct advantage to using weak counter-propagating light is that it does not harm or alter the medium as it passes through. The weak counter-propagating pulse cannot ionize the medium significantly. Thus, the counter-propagating light can meet the main generating pulse at any point in the medium without affecting regions yet to be encountered by the main generating pulse. The arrival time of the short forward pulse can be scanned to produce harmonics at different locations throughout the medium while interacting with the wide counter-propagating pulse. The disadvantage of using the counter-propagating beam is that, even though its power is relatively small, it travels back to the laser system where it runs the risk of damaging the optics. However, the weak counter-propagating light traveling back to the Ti:Sapphire crystal has an advantage in the case when the amplifier works in a saturation regime. The crystal emits some spontaneous radiation in all directions, but the presence of backward traveling counter-propagating beam takes some of that energy for coherent amplification. Thus, the presence of the counter-propagating beam significantly decreases the amount of
spontaneous radiation emitted from the crystal. If we periodically block and re-open the spatially aligned counter-propagating beam, we may observe the effect of the dimming of the spontaneous emission from the Ti:Sapphire crystal with the same periodicity. This effect is a nice tool for spatial alignment of the colliding beams. After both beams are spatially aligned, the temporal alignment is just a straightforward search for the excess Rayleigh scattering (bright dot) mentioned earlier in this section. We do it by means of adjusting the arrival time of the forward beam while moving the delay stage back and forth.

The spectrometer design is explained in more detail in Natan Terry’s undergraduate thesis [74]. I helped to construct the spectrometer. The high harmonics are detected using a 600-lines-per-millimeter grazing-incidence tungsten-coated grating prepared on BK-7 substrate with 1-meter radius of curvature. The grating has relatively high reflectivity in the EUV spectrum range. It disperses different harmonic orders spatially and focuses them onto a Micro Channel Plate (MCP) placed along a section of the grating’s Rowland circle. The MCP is coupled to a phosphorus screen to visualize the harmonics. A CCD camera is used to observe the light emission pattern on the phosphor screen. The CCD camera is read by a Spiricon beam profiler connected to a PC computer running LabView software. The detection chamber is connected to the harmonic production chamber via an intermediate section necessary for differential pumping of the experimental setup. The intermediate section includes its own 65 liter/sec Balzer turbo pump. The detection setup consists of the following:

1) a vacuum-tight stainless steel chamber body with 70 liter/sec Varian turbo pump and a cold cathode pressure gauge;
2) an EUV diffraction focusing grating;
3) a rotational and translational stage for adjusting the grating angle and distance to the MCP for proper focusing of particular harmonics.

The detection setup is designed to resolve harmonics of orders 7 through 35 focused by the grating onto the MCP. The pressure in the interaction chamber is maintained below $10^{-2}$ torr despite the large “leak” that is the gas jet. From our observations, the pressure in the chamber is low enough to avoid serious re-absorption of the harmonics traveling from the focus to the detection chamber. Nevertheless, the pressure in the detection chamber must be on the order of $10^{-5}$ torr or lower for the MCP to function properly (the reason for the 3-stage differential vacuum-pumping scheme). Small apertures (~3 mm diameter) are used to isolate the three pumping chambers. The grating diffracts the harmonics into a horizontally dispersed series of vertical lines on the phosphor screen. Many harmonic orders can be resolved and observed simultaneously. The lines pattern can be observed by eye and also using a CCD camera. A typical CCD camera image is similar to that shown in Fig. 4.5.

3.3. LASER SYSTEM ANOMALIES

We observed spectral splitting of the high harmonic signal during some of our experiments. We determined that the splitting has origins in the laser oscillator system rather than in the dynamics of the harmonic generation phenomenon [75]. Since the oscillator used in this system is a “home-made” system, many of its parameters need optimization and frequent re-adjustment during operation. For instance, we observed that the oscillator prefers to double-pulse when its pump power is just above 4.25 W. The
double pulsing is especially frustrating in the presence of feedback from the amplification stage or when some mechanical stress is introduced to the optical table. When the oscillator is double-pulsed, each of the two pulses generates harmonics. Typically, the separation between pulses can be a few nanoseconds, or sometimes a few picoseconds. The two pulses have slightly different spectral content centered on different wavelengths, so the pulses produce slightly different harmonic frequencies during experiments. In our detection system, the two lines representing the $q$th harmonic appear close enough to each other to give the appearance of one line, slightly split. The double pulsing is an especially undesirable defect because the amplified pulse energy gets divided into two parts with some unknown proportionality factor. Each of these two pulses has energy lower than that in the ideal case of no double pulsing. This frustrates the production of higher harmonic orders. If this defect occurs randomly or on purpose by introducing some stress into the oscillator, it can be detected by a sudden decrease in the highest harmonic order produced. The double pulsing can also be detected in the focus as multiple bright plasma dots (in the case when both pulses are well compressed). The most obvious detection of multiple pulsing is seen using the cross-correlation technique (see Chapter 6).

The energy of the laser system differs over time by ~10%. This results in the harmonics significantly changing their brightness over short periods of time. There may be more than one factor that contributes to this unstable performance. The Ti:Sapphire crystal might be working in an under-saturation regime. While the gain in our amplifier is ~4 times per pass (12 passes total), the total amplification gain is on the order of few million. Even a slight change in the oscillator pulse energy, or pump pulse energy, can cause large changes in the amplified signal. When the amplifier medium is saturated, the
erratic pulse energy fluctuation is limited and helps the amplified pulse energy to be more stable. In addition, certain modes of the oscillator produce pulses, whose energies differ from shot to shot. The thermal effects in the amplifying Ti:Sapphire crystal also affect its performance. This unstable shot-to-shot operation is mitigated by averaging data from hundreds of laser shots using a computer acquisition program.

3.4 CONTROL OF LASER HIGH HARMONIC GENERATION

Each of the two beams is focused using a 30 cm focal-length lens. The beam diameter at each lens is 7.5mm, providing f/40 focusing. The laser wavelength is centered on 800nm. The two beams have equal energies of ~0.15mJ. The duration of the generating laser pulse is measured by autocorrelation to be 30fs (FWHM), producing a peak intensity of approximately 5×10^{14} W/cm² on a 50μm diameter focal spot (measured from 1/e² intensity). The focused beam is 1.2 times diffraction limited.

The gas nozzle consists of a 3mm section of a syringe needle with 300 μm diameter. It is positioned at the collision point of the two pulses. The temporal alignment of the counter-propagating pulse is marked in a cross-correlation setup between the two pulses. The duration of the counter-propagating light is adjusted by changing the compression grating separation and thereby introducing chirp. The pulse duration gets lengthened to 1ps (FWHM), measured by cross-correlation with the forward-propagating pulse. When the counter-propagating pulse is strongly chirped, the collision of the two pulses cannot be discerned through Rayleigh scattering from ionized electrons. However, the temporal alignment is assured by monitoring the cross correlation of the two beams. The gas nozzle is positioned just at the edge of the colliding laser beams where the
thickness of the gas flow is similar to the thickness of the nozzle opening. The gas flows continuously during the experiment. The backing pressure for the experiment is typically a few tens of torr.

![Graph](image.png)

Fig. 3.6 Harmonic signal as a function of counter-propagating pulse delay with a narrow gas distribution.

The highest harmonic produced in argon observed under our laser conditions is the 31st. Fig. 3.6 shows the relative emission of the 23rd harmonic produced in argon with a jet backing pressure of 20 torr. The relative arrival time of the 1ps counter-propagating pulse has been scanned (plotted as a function of delay stage shift). We checked that when the counter-propagating pulse arrived either well before or after the main generating pulse, the harmonic production was the same with or without the counter-propagating light. This rules out the possibility that the counter-propagating light harms the medium (e.g., through ionization). The intensity of the counter-propagating light is lower than the
main generating pulse intensity by roughly a factor of 60. Thus, its intensity is insufficient to ionize the medium significantly.

As is evident in Fig. 3.6, the harmonic generation process is turned off, nearly to the point of extinction. The contrast for the 23rd harmonic is approximately two orders of magnitude. Similar behavior is seen for the neighboring harmonics, although the lower harmonic orders (i.e. the teens) do not show as strong an extinction. This is expected since a stronger standing phase modulation on the laser field is required to suppress the lower harmonic orders.

In the actual experiment, the envelope of each laser pulse varies in time, in contrast with true plane waves. This is especially true of the well-compressed forward-traveling pulse. Moreover, the counter-propagating pulse we use in the experiments is chirped so that the frequencies of the colliding pulses do not necessarily match everywhere in the collision region. Nevertheless, as evidenced by the experimental results, the details of how the pulses vary somewhat from true plane waves are not critical to achieving strong suppression of harmonic buildup. It is possible that intensity-dependent intrinsic phases also play a role [76].

We created poor phase matching conditions by moving the gas jet away from the laser focus a distance of 1 mm. This produced a gas distribution in the laser focus approximately 1 mm thick. The backing pressure was increased to 50 torr in order to maintain pressure at the interaction area similar to what we had in the previous experiment. In the absence of counter-propagating light, the 23rd harmonic emission with the wider gas distribution was very poor due to a severe phase mismatch, arising mainly from geometrical effects owing to the Gouy shift. The Rayleigh range of each laser beam
is approximately \( z_0 = 2 \) mm. Near the focus, the phase mismatch becomes \( \pi \) over a distance of roughly \( \Delta z = 0.3 \) mm (coherence length for 23rd harmonic). Therefore, we estimate there to be approximately three phase zones within the gas distribution (see Eq.(1.23) in Chapter 1), the outer two zones having the same phase and differing from the center zone by \( \pi \). The result is very poor emission of the 23rd harmonic because the center zone with its higher gas density presumably offsets the emission from the other zones.

Fig. 3.7 shows the emission of the 23rd harmonic as a function of the counter-propagating pulse delay. Since the gas distribution is much thicker, the 1ps counter-propagating pulse is only able to interact with the generating pulse over a fraction of the gas distribution (i.e., approximately one phase zone). As seen in Fig. 3.7, the harmonic emission shows a dramatic increase, as the counter-propagating light is able to suppress harmonic emission in a portion of the gas jet. This restored the signal to about the level when the jet is close to the laser beam in the absence of counter-propagating light. The curve also gives direct evidence for the thickness of the gas jet.
Fig. 3.7 Harmonic signal as a function of counter-propagating pulse delay with a wide gas distribution.

3.5 HIGH-ORDER HARMONIC GENERATION IN CROSSED LASER BEAMS

Our group also disrupted and enhanced harmonic generation with colliding-beam geometries other than counter-propagating [77]. These experiments are described in more detail in John Madsen’s thesis [78]. Rather than sending the interfering light pulses into the focus in the exact counter-propagating direction, we directed the interfering light a few degrees off axis. We also sent the interfering light into the focus from the perpendicular direction. In both cases, the weak interfering light was effective in suppressing harmonic emission (23rd harmonic). An obvious advantage to working off-axis is the avoidance of potentially damaging feedback to the laser amplifier system from the residual energy in the counter-propagating beam. The disadvantage is that we could no longer rely on the dimming effect of the spontaneous emission from the Ti:Sapphire crystal.
For both off axis cases, we created poor phase matching conditions by moving the gas jet away from the laser focus a distance of 1 mm. We estimated that this produced a gas distribution in the laser focus approximately 1 mm thick. The pressure in the wider gas distribution was adjusted to 2-4 torr (with increased backing pressure). Under these conditions, there was more than one phase zone within the gas distribution, which resulted in very poor emission for the 23rd harmonic. Again, we were able to restore the harmonic production with the interfering laser pulse crossing the primary beam in a region of the focus.

3.6 CONCLUSION

We have demonstrated that it is possible to “turn off” harmonic production with relatively weak interfering light [72,77]. The interfering light may intersect the primary laser pulse at virtually any angle (other than near co-propagating with the primary laser pulse). We have shown that our approach is an effective tool for quasi phase matching. However, these results, which constitute the central message of this dissertation, merely show proof-of-principle that the quasi phase-matching approach works. The next step is to apply the technique to much higher harmonic orders (where better phase-matching techniques are most needed) and to utilize multiple counter-propagating pulses for a substantial boost to harmonic output. For this purpose, we have constructed a larger laser system, capable of producing up to two orders of magnitude more energy per pulse. The new laser system is described in Chapter 5. The experimental setup for producing high harmonics is described in Chapter 6. With the new system, we expect to generate harmonics beyond 100 orders from neon. Although, I have constructed the new laser
system, the experiments that will be performed with it are not included in this dissertation.

When considering quasi phase matching with counter-propagating light, one should keep in mind that there is an efficiency cost associated with diverting laser energy into an interfering beam; if the energy is retained in the forward beam, presumably a somewhat larger focus could be used to increase the harmonic output. Nevertheless, because only relatively low intensity is required in the interfering beam, the amount of energy required for even a comparatively long interfering pulse is modest, and the potential for an overall efficiency gain is substantial, especially in situations where the phase matching is otherwise poor (higher order of the observed harmonics, higher free electron concentration in the laser plasma, etc.).
CHAPTER 4

DEVELOPMENT OF COMPACT EXTREME ULTRAVIOLET REFLECTOMETER

4.1 INTRODUCTION

High-order harmonic are polarized, highly directional and coherent (i.e., focusable). As discussed in Chapter 1, the harmonics form a comb of frequencies with nearly uniform intensity throughout the EUV spectral range (the “plateau”) of wavelengths ranging from a few nanometers to hundreds of nanometers. This radiation comb of wavelengths can be utilized to perform EUV spectroscopic reflectometry on various sample surfaces.

As was described in Chapter 3, a compact laser source constructed at Brigham Young University reaches intensity up to $10^{15}$ W/cm$^2$. This is sufficient to produce high-order harmonics of the 800 nm laser light in argon gas. As mentioned in Chapter 1, the efficiency of converting the high-intensity laser light into a particular harmonic order is typically quite low, on the order of $10^{-7}$ ($31^{st}$ harmonic, wavelength 26 nm). However, this provides a flux of about $10^9$ photons per laser shot for a harmonic order in the twenties.

Reflectometry that utilizes the high harmonics has not been previously developed. The commonly used sources of EUV light are synchrotrons and plasma sources. One advantage of the harmonics compared to a plasma source is that they are polarized (similar to a synchrotron source), and the polarization can be easily rotated using a half-wave plate placed in the laser beam before the harmonic production vacuum chamber.
Another advantage of the harmonics is the broad spectral coverage. In this chapter I show that it is feasible to utilize a wide range of harmonics simultaneously.

Spectroscopic reflectometry is an important tool for thin film metrology and for finding optical constants [79]. Optical constants define how light interacts with a material. The optical constants of materials often depend on the fabrication methods. Hence, one must determine the optical constants for the particular methods used rather than to rely on handbook values for the elements involved. Reflectometry measures the power ratio of reflected s- and p- polarizations. Polarized reflectometry measurements typically utilize synchrotron radiation as a light source [80]. However, synchrotron sources are expensive and unavailable to some research teams. The relatively inexpensive plasma source has the disadvantage of not being polarized as well as of sparseness of the available wavelengths. As a consequence, many of the optical constants of materials used in thin films are not well known in the EUV range.

Ellipsometry, in general, is a more powerful method to measure optical constants and thin film layer thicknesses [81]. Ellipsometry reveals the complex reflection coefficients of a surface, whereas reflectometry reveals only the amplitudes of those coefficients. However, ellipsometry requires a means of measuring change of polarization state upon reflection. Thus, special EUV light analyzers of polarization are required [81-85], which are not available at our research group. This chapter discusses preliminary proof-of-concept experiments on EUV polarized reflectometry using the laser high harmonics.
4.2 HIGH-ORDER HARMONICS AS A SOURCE OF POLARIZED EUV LIGHT

The experimental setup for producing and detecting high-order harmonics is shown in Fig. 4.1. The 30fs laser pulses, having focused intensity $10^{14} - 10^{15}$ W/cm$^2$ and repetition rate of 1 kHz, interact with a low-density jet of argon gas in the vacuum chamber. The spectral “comb” of high-order harmonic beams produced in the jet emerges from the focus as coherent beams embedded in the residual laser beam [12-14].

Fig. 4.1 High harmonics are generated in a jet of argon gas in vacuum chamber.

High-order harmonics can only be produced in linearly polarized laser light interacting with an atomic gas. Assuming the classical model for the harmonic generation (see Chapter 1), the harmonics bear the same polarization orientation as the incident light. Thus, we can produce pure s- or p- polarizations, relative to a sample surface. We use a half-wave plate to change the orientation of linear polarization of the harmonics simply
by rotating the polarization of the 800 nm light. Then, the linearly polarized laser light is focused with f/50 geometry to a gas jet with a 300 μm diameter nozzle.

The setup consists of three vacuum chambers: the harmonic production (interaction) chamber, the intermediate chamber (for differential pumping) and the detection chamber. Despite a backing pressure of few tens of torr on the gas jet, the pressure in the harmonics production chamber is on the order of $10^{-2} – 10^{-1}$ torr. This low pressure helps avoid re-absorption of the EUV light in the gas used to produce it. Moreover, we decrease significantly the total path length for the harmonics within the focus by using a special extension to the intermediate chamber. This extension, shown in the Fig. 4.1, has a 3 mm diameter hole (aperture) on its end to reduce the gas flow into the intermediate chamber from the interaction chamber. The exit of the intermediate chamber also has a small aperture for the same purpose. The three-stage differential pump system reduces the pressure near the MCP detector, which is similar to one described in Chapter 3.

In order to evaluate the Polarization Dependent Loss ($PDL$) for the EUV grating, we must calibrate s- and p- polarized light reflectivity in the spectral region of interest. The definition of $PDL$ commonly used in the characterization of its diffraction gratings [86] is

$$PDL = 10 \log_{10} \left( \frac{\epsilon_p}{\epsilon_s} \right), \quad (4.2)$$

where $\epsilon_p$ and $\epsilon_s$ are the measured diffraction efficiencies for p- and s-plane polarized incident light, respectively. The $PDL$ is expressed in decibels.
We produced high-order harmonics of pure s- and pure p- polarizations, and compared the reflected signals. The results, Fig. 4.2, suggest that there is little difference between s- and p-polarized grating diffraction for wavelengths ranging from 27nm to 61nm. This is somewhat surprising. The harmonic signal varies from shot to shot, so that we averaged hundreds of laser shots for each picture. We used a PC computer running a LabView software to collect and process a series of digitized images taken with the CCD camera. As is seen from the figure, the PDL is assumed to be very close to zero since the grating response in the measured spectral range is similar for both s- and p- polarizations.

Fig. 4.2 EUV grating calibration showing similar response for either s- or p- polarized light for the harmonic orders 13-29.
4.3 REFLECTOMETER SETUP

Undergraduate students Greg Harris and Richard Curras helped to construct most of the reflectometer setup. We optimized the setup design and produced the experimental data. The setup, shown on Fig. 4.1, was made such that the EUV grating chamber can be easily turned 90 degrees counterclockwise (as seen on the top view on figure). We install two surfaces for testing in the middle of the chamber at 45° to the incident harmonics. The 45-degree angle is chosen since it is very close to Brewster’s angle for the EUV light spectra. The high-harmonic reflectometer instrument is shown in Fig. 4.3.

![Fig. 4.3 Reflectometer setup.](image)

We installed two thin flat wafers in the detection chamber, one called the “reference,” and the other called the “sample.” Half of the harmonic beam hits each wafer so that the reflectivity of both can be measured simultaneously. The reference and the
sample wafers are aligned such that the harmonics reflected from the wafers are separated spatially as they travel to the MCP (Fig. 4.4).

Fig. 4.4 Reflected from both wafers, the EUV signals are simultaneously resolved using a focusing diffraction grating.

Reflected signals are detected on the MCP (Fig. 4.5). We have chosen a “pure” silicon wafer (with residual silicon oxide film on silicon substrate film, ~18 Angstrom thick) as the “reference” and a second silicon wafer with a ~250-Angstrom silicon oxide film as the “sample.” These wafers were surplus samples that were given to us by Professor David Allred. They were not new and may have been exposed to unknown
contaminants. Nevertheless, they were adequate for performing proof-of-principle reflectometry measurements.

Fig. 4.5 S-polarized high harmonics registered on the MCP after being reflected simultaneously from the “sample” (upper) and the “reference” (lower) wafers.

Fig. 4.5 shows a digitized snapshot taken from the CCD camera for s-polarized light. Harmonics reflected from the reference and the sample film are both displayed. Harmonics ranged from the 13th to the 27th order are observed. In order to evaluate the feasibility of using the high order harmonics in this reflectometer scheme, we have measured signals for both s- and p-polarizations reflected from the wafers. The comparison is shown on Fig. 4.6.

The harmonic signals are measured simultaneously on each sample for a given polarization, which is an attractive feature of the EUV reflectometer design. Fig. 4.6 indicates significant differences between the reflectivities of s- and p- for most wavelengths. Figures 4.6 (a) and (b) also indicate that the signals reflected from each wafer vary, presumably because the wafers have different thicknesses of SiO₂ coatings.
Harmonics reflected from the “reference” (thinner oxide film) appear to reflect low-order spectral components better while harmonics reflected from the “sample” appear to reflect higher spectral components better.

![Graphs showing S- and p-polarized high harmonic signals](image)

Fig. 4.6 S- and p-polarized high harmonic signals reflected at 45-degrees angle from (a) the “reference” wafer and (b) the “sample” wafer.

Some anomalous effects occur if the pressure for producing harmonics is too high. As seen in Fig. 4.7, harmonics of higher orders have higher intensity if the gas density (backing pressure) is kept lower. This may happen in part because of pressure-dependent harmonic beam deflection so that the test and reference substrates are not illuminated the same. In fact, whether each wafer is illuminated equally is general concern that should be investigated. To eliminate this potential problem, we could raster the wafers up and down such that whole harmonics beams hit each wafer in an alternating fashion. Also, we might use a gas cell instead of the jet since the gas flowing from jet into
the vacuum can exhibit non-uniform density distributions leading to non-uniform emission of harmonics across the beam.

Fig. 4.7 Harmonic signal reflected from the sample and reference surfaces as a function of gas pressure.

4.4 CONCLUSION

We have demonstrated feasibility of using the high-order harmonics as a polarized light source to perform reflectometry measurements in the EUV spectral range. The response of our EUV grating (tungsten-coated, 2-degree blaze angle, 6-degree angle of incidence) is similar for s- and p-polarizations. The preliminary experiments represent a potentially important application of high harmonics. Reflectometry can reveal important parameters of thin films and optical surfaces. To use high-harmonic spectra below 27 nm, i.e. the higher order harmonics, it is important to improve the brightness of the harmonics, which is the goal of my dissertation research. Quasi phase matching of the much-higher-order harmonics may enable reflectometry measurements with them as well.
A future goal is the development of ellipsometric capabilities. It will be necessary to develop an analyzer capable of measuring the orientation of the polarization ellipse to determine the phase differences between the complex reflection coefficients for s- and p-polarized light.
CHAPTER 5
CONSTRUCTION OF HIGHER-POWER LASER SYSTEM

5.1 INTRODUCTION

The experiments reported in Chapter 3 showed feasibility of using weak interfering light as an effective tool for achieving the quasi-phase matching. Using the theory and techniques explained in Chapters 2 and 3, we showed that we are able to increase the efficiency of converting laser light into a particular high harmonic. However, our investigations were limited to relatively low-order high harmonics because of energy limitations in our laser system. This chapter discusses the principles and design of a new high-power laser system. The new system produces not only higher pulse energy but also a train of counter-propagating pulses to disrupt harmonic emission in many zones with undesirable phase. The new system accomplishes the following objectives:

1) the system must produce sufficient pulse energy to achieve intensities as high as \( \sim 10^{16} \text{W/cm}^2 \) (necessary for ionizing Ne and He gases) in a relatively large focal volume (i.e., f/50 focusing geometry);

2) the temporal separation between individual pulses in the counter-propagating train must be independently adjustable to enable finding the optimal configuration for suppressing zones with out-of-phase harmonic emission (refer to Fig. 2.2 in Chapter 2).
5.2 CHIRPED PULSE AMPLIFICATION

A. Chirped pulse amplification with Ti:Sapphire

The technique of Chirped Pulse Amplification (CPA) was introduced by Mourou and Strickland [87] in 1987. The technique allows one to increase the energy of short pulses while avoiding very high peak powers that can damage to the amplifier material or other optics in the laser system. As shown by Martinez et al., the pulse can be stretched in time by introducing a frequency chirp caused when the pulse undergoes a large amount of dispersion [88]. This frequency chirp (or stretch) must be reversible since we desire to re-compress the duration of the pulse after amplification.

The CPA scheme is illustrated on Fig. 5.1. Ultrashort (100 fs or less) light pulses are generated at low pulse energy in a modelocked laser oscillator. Typically, an oscillator produces light pulses at high repetition rate (~ 10^8 Hz) corresponding to the round-trip time of the cavity with pulse energies in the range of ~10^-9 J, and with pulse duration of a few, or a few tens of femtoseconds. These ultrafast pulses are then chirped using a diffraction-grating stretcher. The pulses are typically stretched to the duration of about 100 picoseconds, decreasing their peak power by three or four orders of magnitude. The stretched pulses then enter amplification stages where their energy increases by six to nine orders of magnitude, reaching the level of a few tens of millijoules per pulse.
Current state-of-the-art chirped pulse laser systems often employ a Ti:Sapphire (Ti:Al₂O₃) crystal as an amplifying medium. The high damage fluence (9-12 J/cm²) of Ti:Sapphire allows one to pump the gain medium with high energy pulses. In addition, the high saturation fluence permits one to extract relatively high energy to allow the use of amplifier crystals of modest size. Owing to the relatively short exited-state lifetime (3.2 μsec), Ti:Sapphire is usually pumped with pulsed lasers (10-100ns) emitting in the blue-green region of the spectrum [89].

To obtain a total gain of ~10⁸ typically requires many passes through the amplifier crystal (or through several amplifier crystal stages) with gain between 2 to 100 times per pass. It is important to avoid loss during amplification process because the pulse
spectrum narrows with each amplification pass. Fortunately, Ti:Sapphire lases over a wide spectral range centered around 800 nm. This is important since, for example, a 45nm spectral bandwidth is required to produce a 25fs pulse in the system I describe later in this Chapter. However, the gain in Ti:Sapphire is not identical for each of the spectral components that may be present in a pulse. Thus, during amplification, a pulse can suffer modest narrowing of its spectrum, which adds to the pulse duration upon compression.

After amplification, the pulse enters a grating compressor, which undoes the effect of the stretcher and restores the pulse to its short duration. In our experiments, the amplified beam is split into the so-called “forward” and “counter-propagating” beams. These enter separate re-compression stages, allowing individual adjustment of pulse duration and, hence, peak power. We typically compress the main (forward) generating pulse to its shortest possible duration.

B. Stretcher

A traditional stretcher design [88] is shown in Fig. 5.2. The idea is to vary the path lengths between Fourier-spectrum frequency components of the pulse.

![Design of a traditional stretcher.](image)

Fig. 5.2 Design of a traditional stretcher.
The typical design consists of two identical lenses forming a 1:1 telescope between a pair of gratings with the same groove density and a retro-reflectors mirror. The first grating G1 is imaged to G1*. The grating G2 and the image G1* are parallel while G1 and G2 are anti-parallel. Different frequency components travel different path lengths by means of the retro-reflector (flat mirror). The frequency components retrace their paths back through the system. The purpose of the telescope is to cause a negative distance between G1* and G2. Then, the effect can later be undone by bouncing the pulse between two parallel gratings with no telescope in place.

Fig. 5.3 All reflective, single-grating stretcher design.
The higher-frequency components (“blue”) experience longer path lengths (i.e., they emerge from the system at the back of the chirped pulse). In this case, the pulse is called “positively chirped” since the frequency of the pulse increases in time. The pulse is called “negatively chirped” if the “red” frequencies occur at the back of the pulse.

In our laser system, we followed the design given in Ref.[90] to eliminate refractive optics, which can introduce undesirable dispersion and chromatic aberration. A folded reflective concave mirror setup substitutes for the two lenses. One advantage is that only one grating is required, but the principle is the same as in Fig. 5.2. Fig. 5.3 depicts this stretcher design.

C. Compressor

In the compressor, the spectral components of the previously stretched (chirped) pulse undergo the reverse process. That is, the pulse is made short again. In the compression system (Fig. 5.4), the initial pulse diffracts from the first diffraction grating, which sends different frequency components along different paths. The difference in path lengths depends on the spacing between the gratings, the angle between the beam and each grating and the grating groove density. The spectral components diffract from a second grating, which causes them to travel parallel to each other again before being retro-reflected from a flat mirror. By using the right choice of the above-mentioned parameters, the pulse chirp can be undone. To ensure optimal compression, fine adjustment to the grating spacing must be done while measuring the pulse duration.
5.3 COUNTER-PROPAGATING TRAIN OF PULSES SOLUTION

As explained in Chapter 3, we showed experimental feasibility of colliding multiple beams for an overall harmonic emission enhancement. Using only one interfering counter-propagating pulse, we were able to enhance harmonic emission by eliminating emission from one out-of-phase zone in a relatively wide generating medium (especially at poor phase matching conditions). In order to see more significant harmonic emission enhancement, we should frustrate harmonic emission from several out-of-phase zones. We investigated the possibility of creating a sequence of counter-propagating pulses in the grating compression setup. In the compressor (see Fig. 5.4), the frequency components of the pulse diffracted from the gratings are spatially separated horizontally along the retro-reflector mirror (see side view of the mirror on the upper portion of Fig. 5.5). If a mask is placed in front of the retro-reflector mirror, thus blocking particular portions of pulse spectra dispersed on the retro-reflector mirror, it appears possible to modulate the amplitude of pulse upon reflection from the mirror. Then, as the beam retraces its path, the spectral components are put back together. If the “re-compressed”
pulse has some residual chirp, the missing spectral regions create voids in the chirped pulse upon exiting from the compressor.

Fig. 5.5 Elimination of harmonic production (harmonic order in the upper thirties) in the “out-of-phase” regions using a train of counter-propagating pulses created by removing spectral regions of a chirped pulse.
As depicted on Fig. 5.5, the timing and durations of these counter-propagating pulses can be chosen to frustrate harmonic production in regions with undesirable phase, which is the goal of the technique. The remaining regions then constructively interfere.

As described in Jamie Titensor’s thesis [91], this technique suffers from limitations of the Uncertainty Principle:

\[
\Delta \omega_{\text{FWHM}} \Delta t_{\text{FWHM}} \geq \text{Const}. \tag{5.1}
\]

The Uncertainty Principle requires the pulse structure to be on the order of picoseconds or longer before the multiple pulses can be clearly separated.

We elected to utilize a different scheme to create a train of amplified pulses with variable durations between individual pulses. The pulse train is produced by a series of beam splitters followed by amplification of the train of pulses. The scheme offers complete adjustability of the timing of pulses in the train. However, the scheme does not permit the adjustment of duration individual pulses in the train since the counter-propagating pulses all encounter the same compression setup, in sequence one after another.

5.4 LASER SYSTEM LAYOUT AND CHARACTERISTICS

A block-diagram of our new laser setup is shown in Fig. 5.6. The essentials of stretching the pulse, amplifying, and then recompressing are represented.
Fig. 5.6 Block-diagram of new laser system.

A more detailed schematic of the new laser setup is shown on Fig. 5.7. The new system has built into it a separate channel producing a train of up to five independent counter-propagating pulses.

Fig. 5.7 New laser setup.
To begin with, we designed a protective enclosure for the laser system. A high-efficiency (HEPA) air filter blows clean air into the enclosure to prevent dust from contaminating the sensitive optics. The air inlet into the laser room is also equipped with a filtration system. As a general rule, we try to keep the lab clean as clean as possible at all times by wearing special shoes, etc.

Fig. 5.8 Measured spectrum of the K-M Labs “MTS” femtosecond oscillator.
Our pulses originate from a K-M Labs “MTS” oscillator (2), a passively mode-locked Ti:Sapphire laser producing a train of ~15 fs (FWHM) p-polarized pulses. The wavelength is centered near 800 nm with a spectral bandwidth approaching 100nm. The pulses are separated by 11 ns (repetition rate 90.9 MHz), and the total output power is around 600 mW. The oscillator is pumped by a continuous 4.5Watt, 532nm beam, produced by a Coherent “Verdi” laser (1). The Verdi laser produces s-polarized light. Mirror periscope #1 switches the polarization of the oscillator pump beam to p-type. In our setup, the direction of p-polarization is parallel to optical table surface. The measured spectrum of the oscillator is shown on Fig. 5.8.

The train of oscillator pulses travels through a collimation lens (3) and enters the pulse stretcher (4), which expands the 15 fs pulse duration to about 70 ps. The concave mirror in the stretcher setup has radius of curvature 75 cm. The focal distance F (see Fig. 5.3) is 37.5 cm. The distance between the centers of the grating and the concave mirror is 23 cm. The Thermo-RGL grating’s reflectivity in the spectral range of the pulse is below 70% despite our expectations. In the stretcher, the pulse becomes negatively chirped. After the stretcher, the 800 nm light is diverted into the pulse picker system (5). The diagram of the pulse picker setup is shown on Fig. 5.9.
The pulse picker consists of periscope #2, which switches polarization of light from p- to s- type, and Pockels cell #1 with two crossed polarizers on its entrance and exit. A collimating lens is placed just before the entrance cube polarizer #1.

A Pockels cell is a KDP crystal that becomes a wave-plate when an electric field is applied. The electrically induced birefringence rotates the polarization of the light that passes through it. The length of our KDP non-linear crystal is made such that it rotates linear polarized 800 nm infrared (IR) light by 90 degrees when a high voltage (~4.8 kV) is applied across the crystal (this happens when ~7kV is applied to the driver).

The entrance polarizer is necessary to clean the s-polarization state of the incoming light. The crossed polarizer on the exit is used to block s-polarized light and transmit p-polarized light. When the Pockels cell is not active, the pulses are extinguished on the second polarizer. When the Pockels cell is energized for approximately 10ns, it selects a single pulse to become p-polarized (the “seed” pulse) from the oscillator’s train of pulses. This is done at a repetition rate of 10 times per second since this is how often our pump laser fires.
After the first Pockels cell, the p-polarized 800 nm “seed” pulses enter the first amplification stage, which consists of triangle-style multi-pass setup [1] as shown on Fig. 5.10. The beam is parallel to the optical table surface. The height of the beam is 4 inches above the optical table.

![Triangle-type multi-pass Chirped Pulse Amplifier stage #1 (top view).](image)

The beam enters at the turning mirror M1. The mirrors M2 and M3 are concave dielectric mirrors of focal length 50 cm each. They reflect 800 nm light and transmit 532nm light. The mirrors are located 100 cm from each other, forming a triangle arrangement together with a flat wide mirror M4. The Ti:Sapphire crystal (produced by Saint Gobain Crystals) is a 0.25% doped, Brewster-cut, 7 mm thick crystal. The Ti:Sapphire crystal is centered between the mirrors M2 and M3. The “seed” pulse makes 9 passes counter clockwise (see top view in Fig. 5.10) around the triangle formed by the mirrors M2-M3-M4. On each pass, the beam hits different places on each of the mirrors but passes through the same point in the amplifier crystal. On the ninth pass, the
amplified pulse is extracted from the amplifier with a pick-off mirror (7a). In order to control spatial divergence in the amplified beam, each pass of the beam goes through a hole in an aperture array (AA) (Fig. 5.11). The beams at the aperture are separated by 2.5mm.

![Fig. 5.11 Aperture array AA (side view).](image)

The distance between each hole in the aperture array is the same and is made to match the natural separation between the beams on the different passes. This separation is controlled by the distance between the amplifier crystal and mirror M4 (in our case-10.0cm). The diameter of the holes has been optimized to improve the amplifier performance in terms of output pulse energy. Without the aperture holes, the beam tends to grow in width on each pass. The wider beam focuses to smaller spots in the crystal, which can lead to less amplification as well as to crystal damage. To find the optimal aperture size, we first used no aperture array and then introduced other arrays with smaller and smaller holes to determine which gave maximum output pulse energy. With no aperture array in place, the structure of the output mode resembled that shown in Fig. 5.12 (a). With the aperture array depicted in Fig. 5.11, which gave the highest output energy, the spatial structure of the output beam resembled Fig. 5.12 (b).
We synchronized the timing of Pockels cell #1 with the pump laser repetition rate. We have placed an avalanche photodiode near the oscillator output to detect spurious light. The signal from the photodiode triggers the timing circuits for the laser system, which consists of a pair of interlinked Stanford Research System (SRS, model DG 535) delay boxes. The two boxes are configured such that one is slave to the other. The delay boxes have sub-nanosecond fine delay steps and variety of triggering options. One channel of the coupled delay boxes is used solely for introducing the repetition rate of 10Hz in the laser system. The boxes wait for that channel to finish executing its 0.1s delay before accepting a new trigger. The delay boxes controls the firing of the pump laser and the various Pockels cells in the system. The pump pulse activates the Ti:Sapphire crystal a few tens of nanoseconds prior to the arrival of the “seed” pulse, which is sent into the amplifier stage by the pulse picker. The exited-state lifetime in the Ti:Sapphire crystal is about 3μs, which is long compared to all timing intervals except for

Fig. 5.12 The observed beam profile from the triangle amplifier in the case of (a) no aperture array and (b) an aperture array with optimized hole diameter.
the delay channel that enforces the 10Hz repetition rate. The timing sequencing is explained in Appendix A2.

The amplifier is pumped by a 532nm Spectra-Physics “Quanta-Ray” laser, which produces s-polarized light pulses with 12ns duration and energy up to 700mJ. The repetition rate of the laser is internally/externally adjustable over the range of 10±1 Hz. The “Quanta-Ray” (Spectra Physics) laser is a two-stage Q-switched Nd:YAG amplification system with extracavity frequency-doubling. The two amplifier stages are pumped by krypton flash lamps. The Q-switch of “Quanta-Ray” laser receives a signal from the delay controller that is correlated with the other timing circuits in the laser system. The timing boxes also supply a pre-trigger to initiate the flash lamps in the “Quanta-Ray” laser about 180μs before its Q-switch is fired.

We switch the polarization of the green pump laser to p-polarized light by means of mirror periscope #3. We then split the pump beam into two parts at the beamsplitter (BS1). About 45% of the beam energy goes through it and is sent directly to the second Ti:Sapphire amplifier stage. The 55% beam then arrives at an 80% reflecting beamsplitter (BS2), which sends most of the energy also to the second Ti:Sapphire amplifier stage from the opposite direction. This leaves about 12% of the original green pump beam energy, which goes to the first Ti:Sapphire amplifier stage. Another beamsplitter (BS3) divides this weaker beam into two equal beams that are sent into the first Ti:Sapphire amplifier stage from both directions. These pump beams travel by means of mirrors M8-M9-M10 (on the left-hand side) and M6-M7 (on the right-hand side). The beams transmit through mirrors M2 and M3, which are reasonably transparent to the 532 nm pump light.
Telescopes focus each pump beam in the first amplifier stage to points about 10cm beyond the Ti:Sapphire crystal, in each case. Each telescope consists of a negative lens followed by a positive lens (L2 and L3) creating a controllable focal length that we set to approximately to 72cm. This focal length was experimentally chosen to create the highest possible pump intensity for which no damage occurs to the Ti:Sapphire crystal when the Quanta Ray laser is set to maximum.

After a series of performance optimizations, we are able to achieve as high as 8mJ from the first amplifier stage, although we usually operate at 5mJ or less to avoid damaging optics in the system. The overall gain of this triangle amplifier stage is therefore near $10^7$. In order to regulate the output power of the first stage, we use a half-wave-plate in the beam path of the pump beam for that amplifier stage. Since the Ti:Sapphire crystal is Brewster-cut, s-polarized light experiences a $\sim 26\%$ reflection as it enters the crystal. By turning the linear polarization axis of the pump light with the wave plate, we regulate the amplifier’s output power by restricting pump energy. The diameter of the diverging beam emerging from the first amplifier stage is $\sim 2$ mm. This beam grows in size at a rate of an additional 2mm per meter of propagation. Thus, the diameter of the beam at the second amplification stage is close to 5mm.

The first amplifier stage is similar to the amplifier we used in our earlier laser system (see Chapter 3), but has several advantages:

1) a mutual alignment of the “seed” beam and pump beams is much simpler since the pump beams have a wider diameter in the crystal;

2) the pump beam power never reaches the damage threshold value at the crystal;
3) we are able to regulate output power of the amplifier by changing the polarization state of the pumping beams.

The amplified pulses exit the first stage and enter the beam isolator setup (8) (see side view on Fig. 5.13).

![Diagram of beam isolator setup](image)

Fig. 5.13 Side view of the beam isolator setup.

The beam from the first-stage amplifier enters cube polarizer #2, where p-polarized light is cleaned for polarization purity. The p-polarized light then enters Pockels cell #2, which rotates the polarization of the pulse to s-type. This Pockels cell is energized only at the time when the amplified pulse goes through. Cube polarizer #3 deflects undesirable p-polarized light to the extent that Pockels cell #2 does not perfectly rotate the polarization of the laser pulse. The now s-polarized pulse exits cube polarizer #3. A subsequent plate polarizer transmits the s-polarized light while rejecting any p-polarized light.

The pulse then goes through Pockels cell #3, which functions as follows:

1) Pockels cell #3 becomes energized when the incoming s-polarized pulse goes through it, turning the polarization back into p-type;
2) it turns off when retro-reflected light pulses return to it from the opposite direction at a later time (described later).

While traversing the entire selector setup, the laser beam loses about 40-45% of its energy. However, most of the lost energy corresponds to spontaneous emission and satellite pulses that we wish to eliminate from the beam.

The beam leaves the pulse isolator setup and enters the second amplification stage. The beam makes two passes through the second Ti:Sapphire crystal (9) for an overall additional gain of ~10. An extraction mirror (10) sends the amplified pulse to the setup that creates counter-propagating pulses (11). The beam looses ~30% of its power on a series of beamsplitters that are used to create the counter-propagating pulses. The beam emerges from beamsplitter array (11) whereupon a mirror (12) sends the pulse to a long delay stage (13) to compensate for timing. The delay stage (13) plays the same role as the delay stage we used in former experiments. The stage movement mechanism is operated by Melles-Griott “Nanomover” driver controlled from a PC computer running a LabView software. The delay scans the forward beam arrival timing relative to that of the counter-propagating pulses that are created (11). The forward beam (14) then exits the table and enters the pulse compression chamber.

The setup for creating the train of counter-propagating pulses is shown in Fig.5.14. The beam hits several 45° beamsplitters (BS4, BS5, BS6, BS7, and BS8) that each reflects 95% of the incident power. In each case, 5% of the pulse energy goes through to a retro-reflecting mirror (RM1, RM2, RM3, RM4, and RM5). In each case, the retro-reflecting light transmits again through the corresponding beamsplitter with 5% efficiency. The retro-reflected pulses all are very weak with slightly different energies.
depending on which channel creates each of them in the system. The counter-propagating pulse created by RM1 has 0.250% of the original energy, by RM2 – 0.226%, by RM3 – 0.204%, by RM4 – 0.184% and by RM5 – 0.166%. The ratio of energy between the first and last created pulses is 1.36. It is to our advantage that there is a slight discrepancy in the quality of the CVI beamsplitters we use; we put the most reflective ones in first (BS4) to help make all of the retro-reflected beams be similar in pulse energy.

![Diagram](image)

Fig. 5.14 The stage creating the counter-propagating train of pulses.

The distance between beamsplitter BS4 and any of the retro-reflection mirrors RM1, RM2, RM3, RM4 and RM5 is approximately 30.5cm. We are able to vary the position of each retro-reflecting mirror RM1 – RM4 over several centimeters with sub-micron precision using the New Focus “Picomotor” fine-alignment drivers on which the
mirrors are placed. Mirror RM5 cannot be moved. The positions of the rest of the retro-
reflection mirrors are compared to it. With this system, we are able to precisely regulate
the relative timing of the retro-reflected pulses to form an arbitrary train of counter-
propagating pulses.

The second amplification stage is designed to amplify the initial beam up to pulse
ergies around 30 mJ in the first double pass. A fraction of that (~30%) is used to create
the train of counter-propagating pulses. The same amplifier stage is used subsequently to
amplify the train of counter-propagating pulses created by the sequence of beam splitters.
The train of weak retro-reflected pulses therefore makes two passes exactly along the
initial beam path in the second amplifier stage. When the counter-propagating train of
pulses arrives back to the pulse isolator setup, Pockels cell #3 has been turned off so that
the p-polarized pulses get rejected by the plate polarizer into periscope #4. About 1-2 %
of p-polarized light leaks through the plate polarizer, but cube polarizer #3 prevents this
leaking light from re-entering the first amplification stage.

Fig. 5.15 The second amplification stage (side view).
The pulse train is then sent for two more passes through the same Ti:Sapphire crystal along new beam paths. All together, the gain is sufficient to increase the energy in each counter-propagating pulse to over 10mJ. Fig. 5.15 shows a side view of the second amplification stage with the path of the counter-propagating beam explicitly shown. The counter-propagating beam is sent by M15 for its final two amplification passes through the Ti:Sapphire crystal.

The lenses L3 and L4 in Fig. 5.5 are positive 75 cm focal length lenses. They focus the 532 nm pumping light from the “Quanta Ray” laser. The focal points of the lenses are located in each case behind the crystal such that the 3mm diameter beam strikes the crystal from each direction. Again, we want to prevent damage at the crystal while maintaining the highest possible pump fluence. The beam exits at the mirror M19, which sends it to the turning mirror M20, shown on the Fig. 5.7.

The measured spectrum of the forward beam (14) is shown in Fig. 5.16. If we compare this amplified spectrum with the spectrum of the Ti:Sapphire oscillator (Fig. 5.8), we see that the amplified pulse spectrum is no longer centered at 800 nm, but rather at 790 nm. Also, the spectrum gets narrower during the amplification process as is expected. The pulse width is ~32 nm (FWHM). The Uncertainty Principle (Eq.(5.1)) predicts that the shortest available pulse duration after re-compression will be about 46 fs. This is much longer than the 15 fs pulse duration we have in the oscillator where the pulse spectral bandwidth was about 100 nm. This is due mainly to the bandwidth narrowing effect mentioned earlier in this chapter. Also, the spectral response characteristic of the stretcher grating (Thermo-RGL) might significantly shift and narrow the spectrum.
Fig. 5.16 Measured spectrum of the forward beam on exit from the second amplification stage.

To avoid damage to the compression gratings, the high-energy beams are expanded to ~1.5 cm diameter (measured from $1/e^2$ intensity) by means of the telescopes T1 and T2 (see Fig. 5.7). The forward beam (14) goes through a negative lens with focal length 20cm followed by a positive lens with focal length 50cm located at the distance of
30.4 cm after the negative lens. The counter-propagating beam (15) goes through a negative lens with focal length 20 cm followed by a positive lens with focal length 75 cm located at a distance of 56.6 cm after the negative lens. The telescope separations collimate the two beams (compensating for any previous divergence). The counter-propagating beam (15) and the forward beam (14) then exit the laser setup and enter the pulse compression chamber.

The Ti:Sapphire crystals in the oscillator and both amplification stages are temperature stabilized at 18.0°C, using the water-cooling stage (Coherent Laser) connected in parallel to each of the crystal mounts. This stabilizes the temperature and helps avoid long-term over-heating of the crystals that can cause beam distortion and deflection.

5.5 PULSE COMPRESSION

A. Overview of pulse compression chamber and experimental setup

The pulse compression setup, cross- and auto-correlation setup, and overall view of the experiment setup are shown in Fig. 5.17. The pulse compression chamber is a stand-alone vacuum-tight chamber with a large optical breadboard mounted inside. However, for our current experiments we do not require the pulse compression chamber to be evacuated. The vacuum compatibility is available for future experiments where all available energy is put into a single well-compressed pulse. The forward beam (14) and the counter-propagating beam (15) enter the pulse compression setup where each beam is temporally re-compressed independently. After the re-compression, the pulses reflect from 95% beamsplitters (BS9) and (BS10) directing most of the power towards the
experimental setup. The 5% of the light that leaks through the beamsplitters is used to analyze the pulse duration in the auto- and cross-correlation setup (see Chapter 6).

![Diagram of experimental setup](image)

Fig. 5.17 Pulse compression chamber (left) and overview of the experimental setup (right).

Following the compression chamber, the beams (14) and (15) emerge onto an optical table that holds the experiment setup for high harmonic production. Each beam is focused into the vacuum chamber using a 75cm lens (L8 and L9). Lens L8 in the counter-
propagating beam is placed on the New Focus “Picomotor” X-Y-Z stage. We use the stage to adjust the alignment of the focused counter-propagating beam in the collision chamber. The overall path lengths of the forward and the counter-propagating beams are \(~15\) meters, measured from the beamsplitter BS4 (see Fig. 5.14). The path lengths must be equal to each other while the two beams meet either in cross-correlation setup and in the interaction chamber in order to achieve temporal alignment of the pulses within tens of femtoseconds precision. This precision corresponds to a spatial alignment of the colliding beams to within few of micrometers.

**B. Compressor**

An expanded view of the compressor setup is shown on the Fig. 5.18. The setup consists of two independent compressor stages. The alignment of the gratings is critical to achieve the best possible compression of the pulse. This is especially important for the forward pulse, which has to be re-compressed to the shortest possible pulse duration to excite the highest harmonic orders as possible. The forward beam (14) performs an extra delay path between mirrors M24 and M25 to accommodate for unaccounted path difference. The height of the beam is 4 inches above the optical breadboard in the pulse compression chamber. The grating G2 is placed on a manual translation stage to adjust the distance between gratings G1 and G2. The gratings’ grooves are set to be parallel to each other and perpendicular to the optical table. The retro-mirror set (M26, see Fig. 5.18) consists of two plane mirrors placed one above the other, forming a right angle between them. The mirrors lift the retro path to a height of 5.5 inches above the optical
table surface, so that the re-compressed beam passes freely above mirror M25 towards the beam splitter BS9.

Fig. 5.18 Compressor setup.

The angle between the beam pathways M25-G1 and G1-G2 as well as between pathways M27-G3 and G3-G4 has been chosen to be the minimum available (about 15°) to achieve a quasi-Littrow angle re-compression. The beams traveling from M25 to G1
and from G2 to mirror set M26 are set to be parallel to each other and to the optical table.
After careful alignment and optimization, the pulse re-compression of the amplified pulse is as short as 45±4 fs duration. This is the shortest available pulse duration we may achieve upon compression according to the Uncertainty Principle (see section 5.4). However, this is about 15fs longer than we would like, so more work will need to be done on this in the future. The counter-propagating pulse compression setup is similar to the one for the forward pulse. The only difference is that we use a Melles-Griot “Nanomover” driver controlled from a PC computer running a LabView software. The delay stage (13) (see Fig. 5.7) and the grating G4 must move together in such a way that the relative timing of the two beams is preserved while adjusting the compression of the counter-propagating pulses. The optimal distance between the centers of the gratings G1 and G2 is found to be 32.5 cm, which produces the shortest available forward pulse duration. The distance between G3 and G4 is 34.4 cm when the duration of each counter-propagating pulse is 0.83 ps.

C. Auto- and cross-correlation

The auto- and cross-correlation setup uses a 20cm-focal-length lens (L7) for 1:1 imaging of light produced in a KDP crystal (Quantum Technology). The second harmonic light is generated by mixing two laser pulses in the crystal. The light is observed by a CCD camera (CCD camera 1) placed after the lens. This CCD camera is read by a Spiricon beam profile analyzer. The distance between lens L7 and CCD camera 1 is 40 cm, which matches the distance from the crystal to the lens.
The counter-propagating pulse train is temporally characterized through a traditional cross-correlation technique, relying on our short forward-traveling pulse. The short forward-traveling pulse is analyzed through an auto-correlation technique [91,92]. An expanded view of the auto- and cross-correlation setup is shown on the Fig. 5.19.

Compared to our former setup (see Fig. 3.3 in Chapter 3), the main advantage of our new auto- and cross-correlation setup is that we can do pulse duration measurements without interrupting the pulses that go to the experiments. The purpose of our setup is:

1) to measure the duration of the forward beam using an auto-correlation technique;

Fig. 5.19 Auto- and cross-correlation setup.
2) to measure the duration and relative timing of the counter-propagating pulse train using a cross-correlation technique;

3) to ensure temporal alignment of the colliding pulses while altering the compression of the counter-propagating pulses and simultaneously moving the translation stage (13) and the grating G4.

As was previously mentioned, the beams for the auto- and cross-correlation setup are obtained from beam splitters BS9 and BS10. Each transmits 5% of the incident power. The forward beam (14) reflects from the manual translation stage formed by the mirrors M29 and M30, which control the path length of the beam. This beam and the counter-propagating beam are both diverted towards the KDP crystal, which we use also for cross-correlation measurements (with slightly different alignment). The KDP crystal is only 0.1 mm thick to minimize phase mismatch of the second order harmonic produced within crystal’s volume. The crystal is cut in a specific crystallographic orientation for near-normal-incidence auto-correlation measurements of a femtosecond pulse. Typically, the angular separation between the correlating beams is 10° or less. Each beam sent into the crystal produces second harmonic light (blue color, 400 nm). When two beams have temporally overlapping pulses within the crystal volume, a third beam of the second harmonic light emerges in a direction between the two original beams. The orientation of the crystal influences the strength of the second harmonic emission through phase matching. This second harmonic light emerging from the crystal reflects from mirror M23 mounted on a removable magnetic mount and is sent towards the CCD camera 1.

The auto-correlation setup for measuring the duration of the forward-propagating pulse utilizes a 50% beam splitter BS11 to create a replica of that pulse. The beam splitter
is placed on a removable magnetic mount such that it intersects the beam going to the manual stage (M29, M30). This beam splitter is installed when it is desired to make an autocorrelation measurement. The two beams are then sent to mix in the crystal. Extra reflections in the setup (M33, M34) ensure that the relative path length of the two beams is similar. The second harmonic light is reflected from the mirror M22 mounted on a removable magnetic mount, and imaged at the CCD camera 1. More description of the auto- and cross-correlation techniques is given in Appendix A1.

5.6 SUMMARY OF LASER SYSTEM

A pulse from the femtosecond oscillator (“MKS” oscillator, K-M Labs) having energy of ~1 nJ is amplified in the first amplification stage to an energy of about 5mJ (gain ~10^7 times). The oscillator tends to double-pulse if its pump power is about 5 Watts or more. We avoid this double pulsing by pumping the oscillator with 4.5 Watts. The oscillator produces on average about 450 mW of power. The amplified pulse loses about 50% of its energy while traveling through the pulse isolator setup. However, the pulse reaches an energy of ~30 mJ after two passes in the second amplification stage (gain 12 times). Part of this pulse is used to create additional counter-propagating pulses with typical initial energy of ~75 μJ. After a total of four passes, these pulses individually reach intensities up to 11 mJ (gain ~147 times).

The spectral bandwidth of the forward pulse at the exit from the laser setup (before the compressor) is measured to be only ~32 nm (FWHM), which is one-third of the pulse bandwidth produced by the oscillator. The forward pulse is re-compressed to a measured duration of 45±4 fs. The measurement error is due to the resolution of our auto-
correlation setup. The amplified pulse spectrum is centered at 790 nm instead of 800 nm as expected. This may give us a clue as to why the spectral bandwidth is disappointingly narrow. In the future, we plan to perform further investigations of the amplified spectrum narrowing phenomena due to stretcher and compressor gratings, and due to the amplification narrowing effect.

We have examined the intensity profiles in two dimensions of the forward beam before and after re-compression. The intensity profiles are very close to the ideal Gaussian distributions in both dimensions.

We have evaluated the operation of the laser system for a regime that does not damage laser components too easily. For best results, we adjust the pump power at the first amplification stage such that its output is near ~5 mJ per pulse. The forward beam pulse energy is about 20 mJ following the series of beam splitters. The pulse energy of beam #5 in the counter-propagating train of pulses ends up being 10 mJ. The sum of the energies for all five counter-propagating pulses reaches 55 mJ.

We re-compress all pulses in the counter-propagating train in a single grating setup. This means that they all have the same pulse duration (which we leave somewhat chirped). The relative temporal alignment of the colliding forward and counter-propagating beams is not affected by re-compression or chirping of the counter-propagating beams. We can adjust the relative timing of the counter-propagating pulses while observing their temporal alignment in the cross-correlation setup. This makes it easy to coordinate the timing of the forward-propagating pulse via its delay stage (13) with the counter-propagating pulses when their compression is adjusted. We use a
computer running a LabView software to keep track of the simultaneous movements of the stages. The software accuracy is calibrated by cross-correlation measurements.
CHAPTER 6

EXPERIMENTAL RESULTS

6.1 SUPPRESSION OF HARMONICS AND SEARCH FOR ENHANCEMENT

A. Limitations of former experimental setup designs

The apparatus we designed and constructed for our new experiments is similar in organization to that used in the earlier experiments (see Chapters 3 and 4), but it is more advanced. There are several major differences between the current and old designs. Some criticisms of our old approaches are presented here in order to appreciate the capabilities of the new setup.

In the earlier experiments we experienced difficulties with spatial misalignment of counter-propagating beams when the atmospheric gas from the chamber got evacuated. Even though the beams were originally well aligned in atmosphere, the alignment would shift seriously when the chamber was evacuated. The phenomenon was likely due to the movement of vacuum chamber walls during pumping. One of the lenses was installed directly on a chamber port (see Fig. 3.3 in Chapter 3). In addition, the optical breadboard inside the interaction chamber was attached to the chamber walls. Thus, any deformation of the chamber could also cause misalignment of the internal optics. To preserve alignment of the colliding beams, we had to pump the chamber gradually in several iterative steps, lowering the pressure inside the chamber by a few hundred torr at each step while re-aligning the beams to keep a bright dot of plasma visible (the result of colliding pulses if sufficiently well-aligned). When the chamber was evacuated completely, we relied on the dimming of the spontaneous emission from the Ti:Sapphire
crystal (owing to beam feedback) to indicate the closeness of the alignment (see description of this effect in Chapter 3).

We had two separate roughing pumps, one connected to the interaction chamber turbo-pump, and another connected to both turbo pumps of the intermediate and the detection chambers. The former had a much higher pump rate than the latter, which caused the apertures between differential pumping chambers to move out of alignment, owing to momentary high pressure differentials during initial evacuation.

In the old setup, there was no way to adjust the spacing between the lens that focuses the forward beam and the lens that focuses the counter-propagating beam while the chamber was under vacuum. The self-focusing of both beams in air caused the apparent focus to occur at a shifted position (~0.5cm). Thus, the alignment of the focal positions found in air became obsolete in vacuum. John Madsen’s thesis [78] explains the difficulty of troubleshooting this type of misalignment.

A single CCD camera observed the alignment of the beams in two dimensions. The CCD was positioned horizontally from the intersecting beams, while vertical images of the interaction area were observed with the same camera by means of a 45° mirror installed just above the collision region a few millimeters away. The laser ablated the end of the steel nozzle during experiments, which quickly degraded the reflectivity of the near-by mirror with a layer of debris. Cleaning and re-installing the mirror was a difficult task. In addition to the problems in the experimental setup, we also had to deal with the unstable operation of the laser system as mentioned in Chapter 3.
B. Experimental setup

Most of the problems that we encountered in the former experimental and laser setups (see Chapter 3) have been fixed. A detailed view of the new experimental setup is shown on the Fig. 6.1. The forward beam (14) is directed to the interaction chamber by mirrors M35-M37 and focused by a 75 cm focal length lens L9 to the center of the chamber. The diameter of the beam-traversing aperture A1 is 15 mm (measured from $1/e^2$ intensity), providing the f/50 focusing geometry. The measured focal spot diameter is ~70μm (measured from $1/e^2$ intensity). The counter-propagating beam (15) is reflected from mirrors M38 and M39. The counter-propagating beam is focused into the interacting chamber by means of a 75 cm focal length lens L8 and a 45° mirror M40. The beam enters the vacuum chamber through an AR-coated window placed on the chamber vacuum flange between L8 and M40. The beam diameter at the aperture A2 is identical to one in the forward beam providing f/50 focusing. The lens L8 is installed on a motorized translation stage, which can adjust the position of the lens in three dimensions with sub-micron precision. The X and Y adjustment is used to spatially align aiming of the colliding beams. The Z coordinate adjustment moves the counter-propagating beam waist towards or away from the forward beam as necessary, even when the chamber is evacuated.
Fig. 6.1 Experimental setup.
We observe plasma streaks in the focus of the intense pulses, seen when the chamber is backfilled with air, using CCD cameras 3 and 4 facing along X and Y coordinates. The CCD cameras are connected to separate TV monitors. When the chamber is back-filled with atmospheric pressure air, each beam nominally should focus to positions 75cm after their respective lenses, but instead they each focus to ~74.5cm due to the self-focusing effect [78]. When the chamber is evacuated, they focus again at 75cm (because the self-focusing effect goes away). We calibrated the positions of the beam waists using the CCD cameras observing the plasma streaks in air. We then determined the necessary displacement for the Z-translation of lens L8 to compensate for alignment as the chamber was evacuated. For this purpose, lens L8 is moved ~1 cm towards the focus. The alignment technique is shown in Fig. 6.2.

![Diagram](image)

**Fig. 6.2** The misaligned foci (a) in air and (b) under vacuum.

Mirror M40 has a 3 mm diameter hole drilled through its center. The counter-propagating beam diameter is 9.3 mm upon reflecting from M40. This means that about
10% of the beam’s area on the mirror is cut. The energy loss because of the hole is much less than the loss we incurred for the experiments described in Chapter 3, about 20%. The focal diameter of the counter-propagating beam is similar that of the forward beam (see measurement results in Fig. 3.4 in Chapter 3).

The vacuum chamber consists of four sections forming a common vacuum volume. Each section of the chamber is clamped tight to the optical table and connected via flexible bellows to the other sections. This provides stability of optical alignment while the chamber is evacuated. All turbo pumps in the setup are connected to a single oil-free scroll-roughing pump, which ensures that all sections of the chamber are evacuated together.

The detection chamber is the same one used in former experiments [72,77]. It houses either a 1200 lines/mm or a 600 lines/mm tungsten coated grating deposited on BK-7 glass substrate of focal length 1 meter. The 1200 lines/mm grating can resolve harmonic orders in the forties or higher, but it is ineffective for harmonic orders in the twenties. The 600 lines/mm grating is effective for harmonic orders in the twenties and teens, but the maximum order it can resolve reliably is about the 35\textsuperscript{th}. The detection setup produces sharp images of distinguished harmonic orders on the phosphor screen. A turbo pump (70 liter/sec) maintains $\sim10^{-6}$ torr vacuum in the detection chamber. This is important for the MCP to function properly. The image produced by the MCP can be observed by eye or on CCD camera 2 connected to a Spiricon beam profiler, coupled with a PC computer running a LabView software. The entrance to the chamber has a 5mm diameter aperture to restrict the flow of gas from the interaction chamber.
An intermediate section of the vacuum chamber setup enables the counter-propagating beam to enter. Mirror M40, which has a hole through it, is mounted on a small optical breadboard inside this section. This section is equipped with a turbo pump (60 liters/sec) for the purpose of differential pumping.

The section of the vacuum chamber where harmonic generation occurs is equipped with a positioning arm connected to the top flange. The positioner moves the gas jet in three dimensions within the center of the chamber. At the exit from this chamber section, a 3mm-diameter aperture restricts gas flow into the intermediate and detection chamber sections. Two CCD cameras, #3 and #4, are mounted on the bottom in the middle of this chamber section. They are used to align the colliding beams both spatially and temporally. The chamber is equipped with a large turbo pump designed to withstand high gas loads.

The forward beam entrance chamber has AR-coated lens L9 mounted on a flange. The lens serves as the entrance window for the forward beam. This chamber is designed not to move when the chamber is evacuated.

The gas jet consists of an electrical pulsed valve (The Lee Company, model VHS-SP) triggered from the SRS laser-timing box. An oval-shaped nozzle (width 1 mm, thickness 0.3 mm, length 1 cm) is connected to the valve. We estimate that the width of the nozzle should correspond to approximately 4 phase zones for the highest-order harmonics that we observe (see Eq.(1.23)). As seen in Fig. 6.3, the laser notched the nozzle end-piece since the nozzle was placed as close as possible to the focus during the preliminary experiments. However, the shape of the eroded nozzle probably helps maintain a more uniform and concentrated gas density distribution in the region of
harmonic generation. In the future, our research group plans to incorporate a gas cell rather than a pulsed jet, since the cell can provide a more uniform gas distribution in the interaction region.

Fig. 6.3 The nozzle end-piece eroded (notched) during preliminary experiments.

The repetition rate for the pulsed valve is set to that of the laser, i.e. 10 Hz. The valve opens ~180 μs prior to the laser pulse arrival. The valve stays open for a regulated duration as short as 250μs. The pulsed operation of the gas jet helps preserve expensive gases and decreases the load on the vacuum pumps. The area of the jet’s opening is about four times that used in former experiments (we used a 300μm diameter nozzle constantly
supplying gas into the collision area). If we used a constant flow of gas through this larger new nozzle opening, it would represent a significant increase in gas usage. Most of the gas would be wasted between the 10 Hz laser shots. The jet is connected through flexible tubing to a swage-lock jack mounted on top of the interaction chamber. Outside of the chamber, the flow is controlled with a leak valve and pressure gauge (species insensitive). A cylinder of neon or argon gas is hooked up to the valve with flexible tubing and a gas regulator.

6.2 PRELIMINARY EXPERIMENTAL RESULTS

We generated high-order harmonics in neon gas in the final days before our laboratory had to be closed for building renovation. The maximum observed harmonic order was 65\textsuperscript{th}, but only harmonics in the mid-forties were distinguishable on the MCP after being dispersed from the 1200 lines/mm EUV grating. The detection has to be modified to effectively resolve harmonic orders higher than mid-forties. Fig. 6.4 represents the image produced on the phosphor screen. Harmonic orders 39\textsuperscript{th} through 45\textsuperscript{th} are resolved with CCD camera 2.
A curious phenomenon we observed was that the harmonic yield improved slightly when the aperture on the forward beam was closed slightly. This was done in the absence of the counter-propagating beam (i.e., it was blocked). Closing the diaphragm (aperture A1) placed just before the lens L9 means both a decrease in the laser energy and an increase in the focal spot size. Thus the focal intensity should decrease. Nevertheless, we observed a marked increase to the harmonic signal in neon gas (meaning an even...
stronger increase of the conversion efficiency). Of course the harmonic production eventually decreased if the aperture size became too small. Under our conditions, we observed that the optimal aperture size was 15 mm in diameter. This effect was observed at several research groups [8,53,93,94]. They used different laser parameters and gases. We intend to do further study of this phenomenon.

Fig. 6.5 Train of five short counter-propagating pulses colliding with the forward pulse in air (atmospheric pressure).
We created a train of five counter-propagating pulses. The pulses were compressed to ~80 fs duration and collided with the short forward pulse in the middle of the air-filled interaction chamber. The pattern of the resulting five bright plasma dots was observed using CCD camera 3. The image is shown on Fig. 6.5. The vertical axis indicates the relative brightness. The other two dimensions indicate the positions of the interacting pulses along the axis of the beam.

Pulse #5 occurs before pulses #4 through #1 in the train. Using the cross-correlator, we set the distance between two neighboring pulses to be 0.5 mm, which corresponds to a timing interval of 1.67 ps. Since the forward pulse and the counter-propagating train of pulses collide with a relative speed of $2c$, the distance between the dots is 0.25 mm (timing 0.83 ps), which is also observed in the cross-correlation setup by CCD camera 1. We blocked both beams and installed a millimeter-graded ruler at the place where the beams collided, and confirmed the data obtained by cross-correlation.
Fig. 6.6 Superposition of the total of five individual snap-shots of short counter-propagating pulses (number 1 through 5) colliding with the short forward pulse in air shows relative brightness of collision dots.

We also took five snap shots of individual pulses in the counter-propagating train colliding with the forward beam. For this purpose, we blocked four pulses at the retro-mirror set RM1-RM5 (see Fig. 5.14 in Chapter 5), leaving one of the mirrors unblocked for each picture. We combined all five data arrays of the snap shots into one, represented on Fig. 6.6. This figure suggests that each of the collision dots #5 through #1 has a similar brightness, i.e., the energies of pulses #5 through #1 in the train are similar. Nevertheless, each of the five dots in Fig. 6.5 has a different brightness. This can be
explained by the fact that the forward pulse gets dispersed more and more as it produces each plasma dot. Thus, it loses its energy before it has a chance to produce the final dots. That is why the dot #1, the most intense one in the Fig. 6.6, happens to be the weakest one in the Fig. 6.5.

If the pulses in the counter-propagating train overlap in time, we observe a “beat” pattern (interference) using the cross-correlation setup. These “beats” occur when the separation between two neighboring pulses is on the order of the pulse width. The periodicity of the interference varies depending on the amount of overlap and the pulse chirp. In the case where two counter-propagating pulses were nearly exactly overlapped in time, we measured the energy of the output (Molelectron detector). A significant increase in pulse energy was observed, which may be explained by the high degree of constructive interference between individual pulses. In this case when the pulse power is high (10 mJ for each pulse in the train before entering the compressor), the optical elements in the laser and compression setups can be seriously endangered. If these “beats” are inadvertently formed while adjusting pulse timing during optimization scans to maximize high harmonics, the experimental results might be misinterpreted. However, there is a possibility of using the “beats” produced from a pair of pulses to create a rapid pulse train with many peaks. This might be useful for dealing with severe phase mismatches where a short period for quasi phase match would be ideal. The period of the interference peaks could be chosen to align with the period of the “out-of-phase” zones.
6.3 HIGH-ORDER HARMONIC GENERATION IN COLLIDING PULSES

We suppressed emission of the harmonic order 39 produced in neon. We collided the short (~45 fs) forward pulse with a single 970 fs long counter-propagating pulse (pulse #5 from the train). The other pulses in the train were blocked. The relative arrival time of the counter-propagating pulse was scanned (plotted as a function of delay stage shift).

Fig. 6.7 Emission of the 39\textsuperscript{th} harmonic produced in neon is suppressed with a weak counter-propagating pulse of light.
The intensity of the counter-propagating light is lower than the main generating pulse intensity by roughly a factor of 60. Thus, its intensity is insufficient to ionize the medium significantly. We checked that when the counter-propagating pulse arrived either well before or after the main generating pulse, the harmonic production was the same with or without the counter-propagating light.

As is evident in Fig. 6.7, the harmonic generation process is turned off in part. The delay stage (13) displacement is correspondent to the relative arrival of forward beam. The emission of 39\textsuperscript{th} harmonic is reduced by a factor of 3. The figure shows the emission of the 39\textsuperscript{th} harmonic produced in neon with a jet backing pressure of 225 torr. This is a relatively high backing pressure despite the use of the pulsing jet. This backing pressure reveals the fact that the jet nozzle opening is much wider than that used in former experiments (see Chapter 3).

6.4 INTERPRETATION OF EXPERIMENTAL RESULTS

Our new laser system does not require frequent re-alignment. The shot-to-shot energy fluctuation of the laser system is \(~10\%\). The forward and counter-propagating beams are identically p-polarized with f/50 focusing. The measured forward beam waist diameter is 70 \textmu m (FWHM, measured from 1/e\textsuperscript{2} intensity) indicating 1.3 times the diffraction limit. The energies of the forward and counter-propagating pulses before entering the compressor setup are 21 mJ and 10 mJ, 4mJ and 1.8 mJ after the compressors, respectively. The forward pulse can be compressed to a duration of 45\pm4 fs,
reaching a focused intensity of $2.4 \times 10^{15}$ W/cm$^2$ (see Eq.(1.8)). The error in the pulse duration measurement is due to the resolution in our auto-correlator setup.

We can produce high harmonics up to the 65th order in neon. However, currently we can reliably distinguish the harmonics up to the 45th order; our detection setup has to be upgraded to accommodate higher orders. The laser intensity necessary to produce harmonics from neon ions is around $1.1 \times 10^{16}$ W/cm$^2$ (see Table 1.2). If the energy loss in the main pulse re-compression stage was a factor of 2 instead of 5 and if it had a duration of 25fs, we could, in principle, produce the harmonics from neon ions past the 119th order. The gratings (Thermo-RGL) in the stretcher and in the main beam compressor are the first objective of our future experiment optimization investigations.

We created a train of five counter-propagating pulses. The distance between neighboring pulses in the train can be varied with sub-femtosecond precision (much more accuracy than needed). The pulses in the counter-propagating train can be chirped all together to any desired duration, so that they can be collided with the strong forward harmonics-generating pulse throughout a wide jet of gas. We hope to control emission of particular harmonic orders using several counter-propagating pulses through a trial-and-error procedure in order to find particular “out-of-phase” zones in laser focus where harmonic emission should be diminished. This technique should lead to overall enhancement of high-harmonic production. A computer program running the translation stages at the mirrors RM1-RM4, coupled with the CCD camera 2 (see Chapter 5) and using a genetic algorithm [95] would be ideal for future experiments.

We suppressed harmonic emission produced in neon. The contrast for the 39th harmonic is approximately 3 times. This is not a significant suppression compared with
the cases discussed in Chapter 3. However, these trial runs represent an initial step
towards a series of new experiments at higher pulse energy. They demonstrate that the
necessary equipment is in place and functioning. The earlier proof-of-principle
experiments (see Chapter 3) give reason for optimism that it should be possible to
improve output of high-order harmonics by as much as two orders of magnitude.
REFERENCES


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A1. Auto- and cross-correlation techniques

The cross-correlation uses a CCD camera and a beam profile analyzer (Spiricon) to view the second harmonic light (400 nm) generated by mixing of two crossing pulses in the crystal. The path lengths of both beams must be matched so that the pulses arrive simultaneously in the crystal. The crystal is installed on a rotation stage, which may tilt the crystal to an arbitrary angle relative to the crossing beams. The crystal is a square piece of KDP crystal with 0.1 mm thickness and 4.5x4.5 mm² surface area. It is mounted vertically in a 1-inch round holder that can rotate. The holder can be turned within this mount around an axis normal to the crystal’s surface. The crystal surface plane stays perpendicular to the optical table surface at all times. For cross-correlation, the beamsplitter BS11 must be removed from its kinematic mount and set away. The mirror M23 must be installed in its place on the same kinematic mount. The image of the cross-correlation stripe in the crystal is projected in 1:1 proportion onto the CCD camera 1 using the 20 cm focal length lens L7.

For auto-correlation, the beamsplitter BS11 must be installed on its magnetic mount. The distances “BS11-M33-M34-crystal” and “BS11-M29-M30-M31-crystal” must be equal to each other in order to cause an interaction between the pulses (making a stripe of 400 nm light (blue color) from the crystal). The tilt of the crystal relative to the auto-correlated beams directions is not necessarily similar to the cross-correlation case tilt. Since the angle between the auto-correlated beams is different (usually, smaller) from the cross-correlation case, the tilt must be adjusted to achieve the best phase match of the
two correlated beams, i.e., the best second harmonic generation efficiency. The mirror M22 shares the same magnetic mount with the mirror M23. Mirror M23 must be substituted for the mirror M22 for auto-correlation.

A2. Timing sequencing in the system

A periodic signal of positive polarity with the threshold amplitude of 0.04-0.20 V and frequency 100 MHz arrives at the SRS timing box from an avalanche photodiode installed in the oscillator. The timing box triggers on one of these peaks after which it ignores input until all timing delays have been executed. The signal gives a timing mark to the rest of the delay channels in the box.

SRS delay box #1.

1. Channel C. Timing C=T+0.100000000 s. This is a “dummy” channel, which creates the repetition rate of 10Hz for the rest of delay circuits. The SRS delay box is triggered from one of the peaks in the signal arriving from an avalanche photodiode (50 Ohm load). The channel C allows triggering of the rest of the delays in the system only at the repetition rate set at C, i.e., at 10 Hz. The signal from channel C triggers the flashlamps in the “Quanta Ray” pump laser.

2. Channels A and B. Timing A=T+181.800 μs and timing B=A+0.300 μs. These two channels produce a rectangular signal at the “positive” AB output. This AB output is coupled to the Q-switch trigger of the “Quanta Ray” laser. The Q-switch can release the stored energy from that laser at the rising edge of the “positive” AB signal. The 12ns pulse from the “Quanta Ray” laser occurs at the repetition rate set at channel.
The laser produces up to 700 mW of average power. The timing B is necessary to reset the triggering logic in the laser.

3. Channel D. Timing D=T+0.001 s. This channel is coupled with the pulsing gas jet electronics. The jet starts opening at the time T for the duration 0.001 s.

The delay box #2 is triggered by positive signal of 1.9±0.1 V from box #1 output.

The two boxes work as if they were one, the repetition rate being directed by the channel C of the box #1.

SRS delay box #2.

1. Channel A. Timing A=T+181.938 μs. This channel is the only channel that must be adjusted ±5 ns or less at any start-up of the oscillator (the maximum value of amplified pulse peak power is observed using a photodiode connected to an oscilloscope and placed after the first amplification stage). The signal from A arrives at trigger port #1 of Pockels cell #1. A 6-foot coaxial cable is connected between trigger ports #1 and #2, providing a rectangular window of the 6ns duration during which the cell is energized.

2. Channel B. Timing B=A+0.042 μs. The signal from B arrives at trigger port #1 of Pockels cell #2. The 6-foot coaxial cable is connected between trigger ports #1 and #2 providing a window of the 6ns duration during which the cell is energized.
3. Channel C. Timing C=A+0.043 \( \mu \)s. The signal from C arrives at trigger port #1 of Pockels cell #3. A difference of 1 ns between B and C naturally corresponds to a distance of 1 foot that light travels between Pockels cells #2 and #3.

4. Channel D. Timing D=A+0.059 \( \mu \)s. The signal from D arrives at trigger port #2 of Pockels cell #3. The signals from C and D provide a 16 ns window during which the cell is energized allowing the amplified pulse travel from amplifier stage #1 to amplifier stage #2. By the time the cell later becomes de-energized, the counter-propagating train returns (about 15ns later). The counter-propagating beam is ejected from the plate polarizer towards periscope #4.