

Modeling Nonlinear Optical Response in 2D materials from Nonequilibrium Quantum Dynamics

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To my

Contents

Acknowledgements	i
Contents	iii
Summary	v
List of Tables	vi
List of Figures	viii
List of Abbreviations	xiii
1 INTRODUCTION	1
1.1 Nonlinear Optical Response	2
1.2 Photocarrier Injection	4
1.3 High-order Harmonic Generation	5
1.4 Review on Theoretical Models	6
1.5 Structure of the Thesis	8
2 THEORETICAL FOUNDATIONS	11
2.1 Crystallography Properties	11
2.1.1 Structural Parameters	12
2.1.2 Reciprocal Space	13
2.2 Tight-binding Approach	13
2.3 Electron Dynamics	15
2.3.1 Time-dependent Schrödinger Equation	16
2.3.2 Quantum Master Equation	18

3 CO-LINEAR POLARIZATION INDUCED PHOTOVOLTAIC EFFECT	21
3.1 Time-dependent Perturbative Analysis on QuI	22
3.2 Third-order Nonlinear Regime: $1 + 2$ QuIC	30
3.3 Deeply Off-resonant Highly-nonlinear Regime	40
4 THZ-INDUCED HHG AND NONLINEAR CHARGE TRANSPORT	49
4.1 Fully Dynamical Simulations for THz Field	50
4.2 Quasi-static Approximation	53
4.3 Nonlinear Charge Transport in Nonequilibrium Steady-state	58
4.4 Comparison with Thermodynamic Model	63
5 ENHANCEMENT OF MIR-INDUCED HHG BY COHERENT COUPLING WITH THZ FIELD	71
5.1 MIR-induced HHG in Graphene under THz Fields	72
5.2 Orientational Dependence of HHG	80
5.3 Comparison of Nonequilibrium Steady State and Thermodynamic Model	83
5.4 Contribution of Nonequilibrium Population	84
6 CONCLUSION AND OUTLOOK	91
A ADIABATIC BASIS REPRESENTATION	93
Bibliography	97

Summary

Modeling Nonlinear Optical Response in 2D materials from Nonequilibrium

Quantum Dynamics

by

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Through the exploration of high-order harmonic generation, this research contributes to the broader scientific community's knowledge and paves the way for new discoveries and technological breakthroughs. The study of HHG not only deepens our comprehension of fundamental physics but also holds great promise for revolutionizing various fields, ranging from materials science and chemistry to biology and quantum technologies.

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List of Tables

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List of Figures

2.1	Lab coordinate system	12
3.1	The time profiles of the electric field given by Eq. (3.47) are shown for (a) $\phi = 0$ and (b) $\phi = \pi/2$	32
3.2	(a, b) The conduction population distribution $n_c(\mathbf{k})$ computed with (a) $\phi = 0$ and (b) $\phi = \pi/2$. (c, d) The population imbalance distribution $\Delta n_c(\mathbf{k})$ computed with (c) $\phi = 0$ and (d) $\phi = \pi/2$	33
3.3	34
3.4	The dc components of the currents $\mathbf{J}_{dc}(t)$ computed from Eq. (3.50) are shown as a function of time. The results using the relative phase of $\phi = 0$ are shown in panel (a), while those using $\phi = \pi/2$ are shown in (b)	35
3.5	The persistent current $J_{dc}(t_f)$ as a function of the relative phase, ϕ . The results are computed by setting E_0 to 2.57 MV/cm and $\hbar\omega$ to 3 eV.	37
3.6	The current after the laser irradiation is shown as a function of the fundamental photon energy $\hbar\omega$. The results computed different field strengths: (a) $E_0 = 1.03$ MV/cm, (b) 51.43 MV/cm, and (c) $E_0 = 102.84$ MV/cm.	38
3.7	(a) The conduction population distribution $n_c(\mathbf{k})$ after the irradiation of the laser field, and (b) the population imbalance distribution $\Delta n_c(\mathbf{k})$ are shown. The results are computed by setting E_0 to 10^{10} V/m.	41

3.8	The dc components of the currents, $\mathbf{J}_{dc}(t)$, are shown as a function of time. The results are computed with the deeply off-resonant condition, $\hbar\omega = 1.0$ eV.	42
3.9	The persistent current, $J_{dc}(t_f)$, is shown as a function of the relative phase ϕ . The results are computed with the deeply off-resonant condition, $\hbar\omega = 1.0$ eV.	43
3.10	(a) The persistent current, $ J_{dc}(t_f) $, is shown as a function of the field strength, E_0 . (b) The number of conduction population after the laser irradiation is shown as a function of the field strength E_0	44
4.1	Computed harmonic spectra $I_{\text{HHG}}(\omega)$ with Eq. (4.2) for different chemical potentials, $\mu = 0, 70$ and 170 meV.	52
4.2	Electric current in graphene under a static electric field, $E_0 = 8.5$ MV/m.	54
4.3	Steady current $J_S(E_0)$ as a function of field strength E_0 . The results of the fully dynamical calculation are shown as the red points, while the interpolated result is shown as the blue-solid line.	55
4.4	Comparison of the THz-induced current computed with the fully dynamical calculation and the quasi-static approximation.	56
4.5	Comparison of the HHG spectra computed with the fully dynamical simulations in Sec. ?? and the quasi-static approximation Sec. ?? . Here, the chemical potential is set to $\mu = 170$ meV.	57
4.6	Nonlinear effective conductivities of graphene as a function of the static field strength E_0 evaluated with the total currents (solid lines) and intraband currents (dashed lines) for different values of the chemical potential, $\mu = 0, 70$ and 170 meV.	59
4.7	(a) The equilibrium population distribution in the conduction band $f^{\text{FD}}(\epsilon_{c,k})$. (b-d) The field induced conduction population change for different field strengths, (b) 0.01 MV/m, (c) 3 MV/m, and (d) 10 MV/m. (e) The population distribution in the conduction band in the nonequilibrium steady-state under a static field, $E_0 = 10$ MV/m.	68

4.8 Computed effective conductivities are shown as a function of the excess energy. The results for the nonequilibrium steady-state (red-solid), the thermodynamic model (green-dashed), and the thermodynamic model plus the single-band approximation (blue-dotted) are shown.	69
5.1 The angular dependence of the harmonic yield obtained from the electron dynamics calculations in the presence of the MIR field. The third, fifth, and seventh harmonic yields are scaled by factors of 60, 800, and 1000, respectively.	74
5.2 (a, b) The current $J(t)$ induced by THz and MIR fields, $E_{THz}(t) + E_{MIR}(t)$. The x -component of the current is shown as the blue line, whereas the y -component is shown as the red line. The inset is the panel (a) shows the time profile of the applied THz field. (c, d) The current $J(t)$ induced by the static and MIR fields, $E_{dc}(t) + E_{MIR}(t - \tau_{MIR})$. The x component of the current is shown as the orange line, whereas the y -component is shown as the green line. In the panels (a) and (c), the polarization of all the fields is parallel to the Γ - M direction (the x -direction in the present setup) as $e_{THz} = e_{dc} = e_{MIR} = e_x$. In the panels (b) and (d), the polarization of THz and static fields is parallel to the x -direction as $e_{THz} = e_{dc} = e_{MIR} = e_x$, while that of the MIR field is perpendicular as $e_{MIR} = e_y$. (e) The power spectra $I_{HHG}(\omega)$ computed using the current in (a) and (c). (f) The power spectra $I_{HHG}(\omega)$ computed using the current in (b) and (d).	75
5.3 The harmonic yields are shown as a function of the static field strength E_{dc} . In each panel, the results obtained using the different relaxation times, T_1 and T_2 , are compared. The results of the third harmonics are shown in the panels (a) and (b), whereas those of the fifth harmonics are shown in the panels (c) and (d). The results obtained using the parallel configuration ($e_{MIR} = e_x = e_{THz}$) are shown in the panels (a) and (c), whereas those obtained using the perpendicular configuration ($e_{MIR} = e_y \perp e_{THz}$) are shown in the panels (b) and (d).	78

- 5.4 The power spectra of emitted harmonics, $I_{HHG}(\omega)$, are shown. The results obtained using a weak THz field ($E_{THz} = 0.1$ MV/cm) are shown in the panels (a) and (b), while those obtained using a strong THz field ($E_{THz} = 1.0$ MV/cm) are shown in the panels (c) and (d). The results obtained using the parallel configuration ($e_{MIR} = e_x = e_{THz}$) are shown in the panels (a) and (c), whereas those obtained using the perpendicular configuration ($e_{MIR} = e_y \perp e_{THz}$) are shown in the panels (b) and (d). 79
- 5.5 The angular dependence of the harmonic yield in the nonequilibrium steady-states under a static field along the Γ - M direction is shown for different static field strengths, E_{dc} . The angle θ denotes the relative angle between the static field and the *MIR* field. (a–d) The total intensity I_{total}^{nth} is shown for the second, third, fourth, and fifth harmonics. (e–h) The component of the intensity parallel to e_{MIR} is shown for each harmonic. (i–l) The component of the intensity perpendicular to e_{MIR} is shown for each harmonic. The results are normalized by the maximum total intensity I_{total}^{nth} for each harmonic. 80
- 5.6 The emitted light intensity, I^{nth} , is shown as a function of the excess energy for (a) third (b) fifth, and (c) seventh harmonics. The results for the nonequilibrium steady-states induced by a static field parallel (red solid line) and perpendicular (blue dashed line) to the MIR field are compared with the thermodynamic model (green dotted line). In each panel, the field strength of the static field parallel to the MIR field is shown as the secondary axis. 89
- 5.7 (a) The calculated conduction population distribution, $n_{ck}^{\text{neq-steady}}$ for the nonequilibrium steady-state is shown. Here, the Dirac point is indicated by the blue circle. (b–e) The angular dependence of the emitted harmonic intensity is shown for the (b) second, (c) third, (d) fourth, and (e) fifth harmonics. The results obtained using the nonequilibrium population model and the nonequilibrium steady-state are shown by the blue and green solid lines, respectively. 90

List of Abbreviations

h-BN	Hexagonal boron nitride.	12, 26, 30
HHG	High-Order Harmonic Generation.	1, 5
QuI	Quantum Interference.	4, 28
QuIC	Quantum Interference Control.	30, 31, 36, 37, 39, 40, 43, 45
SBE	Semiconductor Bloch equations.	7, 8
TDDFT	Time-dependent Density-functional Theory.	7, 8
TDSE	Time-dependent Schrödinger equation.	7, 16, 30

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Chapter 1

INTRODUCTION

In the conventional understanding of linear optics, the response of a material to an incident electromagnetic wave is linearly proportional to the strength of the electric field. However, when the intensity of the incident light exceeds a certain threshold, the nonlinear optical response becomes significant. This regime reveals a rich variety of phenomena, including the photon carrier injection, phase modulation, and harmonic generation. After the first observation of nonlinear optical phenomena by Franken [1], based on recent advancements in laser technology, groundbreaking research in the field of nonlinear optics have been driven [2–4], have conducted in a new era of intense light generation. These developments have paved the way for in-depth exploration of light-matter interactions in highly nonlinear regimes. One of the most captivating nonlinear optical phenomena made accessible by these advances is High-order Harmonic Generation (**HHG**), a process characterized by its extreme photon upconversion and remarkable nonlinear characteristics.

This thesis aims to provide a comprehensive overview of the nonlinear response phenomena including photocarrier injection and high-order harmonic generation in the context of nonlinear optics. We will explore the theoretical foundations of light induced time-dependent quantum dynamics evolution in solid systems, including the quantum mechanical description of the light-matter interaction process and the role of excited electron dynamics. By investigating photocarrier injection and HHG, we seek to deepen our understanding of the nonlinear optical response of materials and unlock the potential for applications in

fields such as ultrafast spectroscopy, attosecond science, and advanced imaging techniques.

1.1 Nonlinear Optical Response

In studies of optical response theory within solid systems, the dielectric function serves as a centered concept, the dielectric function characterizes how the material's polarization, induced by the external field, evolves with the field's frequency and intensity. Mathematically, the dielectric function connects the material's polarization density to the electric field through Maxwell's equations, forming the basis for understanding its optical properties. Linear response theory, predicated on the assumption of weak perturbations, asserts that the induced polarization is directly proportional to the strength of the applied field. Linear response theory is most applicable when the perturbations are weak. In other words, the system's behavior is approximately linear in the vicinity of its equilibrium or initial state, the behavior of a system when it reacts proportionally to an applied perturbation. In such cases, the system's behavior is described by linear susceptibility ($\chi^{(1)}$).

The electric susceptibility (χ_e) describes the response of a material to an applied electric field (\mathbf{E}). The relationship between the induced polarization (\mathbf{P}) and the applied electric field can be expressed as:

$$\mathbf{P}(\mathbf{t}) = \epsilon_0 \chi_e \cdot \mathbf{E}(t)$$

Where:

- \mathbf{P} is the induced polarization vector,
- χ_e is the electric susceptibility tensor, and
- \mathbf{E} is the applied electric field vector.

In nonlinear response, the relationship can be expressed using a power series expansion:

$$\mathbf{P}(\mathbf{t}) = \epsilon_0 (\chi^{(1)} \cdot \mathbf{E}(t) + \chi^{(2)} \cdot \mathbf{E}(t)^2 + \chi^{(3)} \cdot \mathbf{E}(t)^3 + \dots)$$

In nonlinear response theory, when considering the time-domain response of a material to an external perturbation, one often encounters frequency-dependent susceptibilities. These susceptibilities can be expressed in terms of tensor notation and integrated over frequency to account for the material's response over a range of frequencies.

Let's use the electric susceptibility tensor as $\chi^{(2)}$ as an example, representing the second-order susceptibility. In the frequency domain, the induced polarization (P) can be expressed in terms of the applied electric field (E) and the susceptibility tensor as:

$$P_i(\omega) = \sum_{j=1}^3 \sum_{k=1}^3 \int \int \chi_{ijk}^{(2)}(\omega, \omega_1, \omega_2) E_j(\omega_1) E_k(\omega_2) d\omega_1 d\omega_2$$

Where:

- $P_i(\omega)$ is the induced polarization component at frequency ω along direction i ,
- $\chi_{ijk}^{(2)}(\omega, \omega_1, \omega_2)$ is the frequency-dependent second-order susceptibility tensor,
- $E_j(\omega_1)$ and $E_k(\omega_2)$ are the components of the applied electric field at frequencies ω_1 and ω_2 respectively, and
- The integral is taken over all possible frequencies.

In summary, the nonlinear response of a material to external perturbations can be described using tensor notation, incorporating frequency or time integrals to capture the material's response over a range of frequencies or times. The high-order susceptibility tensor ($\chi^{(n)}$) plays a crucial role in characterizing this nonlinear response.

Nonlinear response often involves multiphoton processes, where multiple photons are absorbed simultaneously, lead the progress to be complicated. The mul-

tiphoton absorption and the harmonic generation are different nonlinear optical phenomena associated with different components of nonlinear susceptibilities. Frequency and Intensity-dependent absorption is a key feature of nonlinear response and is often exploited in applications such as laser-induced material processing.

1.2 Photocarrier Injection

In the realm of bulk photovoltaic injection, the second-order nonlinear optical effect, as explored in [5], has garnered considerable attention for its potential in achieving highly efficient light-to-current conversion. Investigations into shift-current, detailed in references [6–9], exemplify the significance of this phenomenon.

Another noteworthy aspect of second-order nonlinear current is the "injection current" [5, 10–13]. This current can be induced by the disruption of time-reversal symmetry through elliptically or circularly polarized light, in addition to the breaking of intrinsic spatial inversion symmetry. The injection current results from the population imbalance induced by quantum interference (**Qui**) between various excitation paths, arising from the interference between absorption pathways associated with orthogonal components of polarization. Consequently, this leads to a polar distribution of electrons or holes in momentum space, generating a current injection that temporally aligns with optical intensity. Remarkably, the non-oscillating current induced by quantum interference may persist even after the perturbing laser irradiation stops.

It is noteworthy that, unlike the shift-current, which occurs solely during laser irradiation, the injection current exhibits persistence even after the conclusion of laser irradiation. This enduring quality underscores the unique and sustained contribution of the injection current in the context of nonlinear optical effects.

Going, beyond second-order nonlinear effects, researchers have investigated into the realm of photovoltaic effects induced by intense few-cycle laser pulses [14–19]. Notably, such laser pulses possess the capability to extrinsically break spatial inversion symmetry. In addition to this, the presence of a strong field introduces

highly nonlinear excitation channels, including pathways such as tunneling excitation. The combination of extrinsic spatial inversion symmetry breaking and intense nonlinear interactions between light and matter opens the possibility for an intense few-cycle laser pulse to induce a direct current (dc-current) even in a material with intrinsic inversion symmetry.

The uniqueness of the photovoltaic effect with a few-cycle pulse lies in its dependence on breaking the inversion symmetry of the incident light fields. This intrinsic connection allows for the manipulation of the induced current by controlling both the intensity and carrier-envelope phase of the pulse [14, 17]. The exploration of these intense few-cycle laser pulses not only expands the understanding of nonlinear optical phenomena but also unveils avenues for precise control and manipulation of induced currents through tailored light-matter interactions.

In recent research, an effective method for manipulating valley population has surfaced, centered around the interplay of two circularly polarized lights with different frequencies, denoted as ω and 2ω . This investigation is particularly fitting in the context of two-dimensional systems [20, 21]. Individually, each circularly polarized light breaks time-reversal symmetry, and when combined, the fields exhibit the added capability of breaking spatial inversion symmetry. This coupled contravention upon time-reversal and spatial inversion symmetries gives rise to a population imbalance in momentum space upon laser excitation. This, in turn, appears as a sustained charge flow persisting even after the laser irradiation. Notably, this methodology has extended beyond theoretical exploration, with numerical studies incorporating first-principles calculations applied to bulk solids [22]. The multifaceted approach of combining circularly polarized lights at different frequencies not only enriches our understanding of valley population dynamics but also holds promise for diverse applications in the realm of materials and quantum phenomena.

1.3 High-order Harmonic Generation

Traditional high-order harmonic generation (HHG) is a phenomenon where intense laser light interacts with a gas, causing the generation of very high-frequency

light waves. This process typically occurs when a strong laser field ionizes a material, causing electrons to be ripped away from atoms. These electrons then undergo complex dynamics, involving acceleration, coherent motion, and recombination with their parent ions. Through this intricate dance, the electrons emit high-energy photons with frequencies much higher than that of the incident laser, often extending into the extreme ultraviolet (XUV) and X-ray regions of the electromagnetic spectrum [23].

Since first observed in 1987 using rare gases as target specimens [24, 25], gas-phase HHG has been intensively utilized to generate ultrashort attosecond light pulses [26–28] for investigating ultrafast dynamics in matter in the time domain which is a typical time scale for the motion of electrons.[29–32]. HHG provides a unique window into electron dynamics and allows us to investigate processes occurring on attosecond timescales. The emitted harmonics carry valuable information about the electronic structure, band gaps, and transient states of the material, offering a powerful tool for probing and controlling ultrafast processes. In recent years, there have been significant advancements in experimental techniques for studying high-order harmonic generation. The use of intense femtosecond laser pulses, pulse shaping methods, and advanced detection schemes have enabled precise control and characterization of the generated harmonics. These experimental advances have led to breakthroughs in attosecond science, providing tools for investigating ultrafast phenomena in a wide range of atoms [30, 33, 34], molecules [35–37], and solids [31, 32, 38–40]. The HHG in solid-state systems was first observed in ZnO in 2011 in mid-infrared (MIR) laser field [41], and it has since garnered significant attention, both from a fundamental research perspective and due to its technological potential, as evidenced by recent studies in solids [42–46].

1.4 Review on Theoretical Models

The generation of high-order harmonics involves complex quantum mechanical processes and intricate interplays between the laser field and the electronic structure of atoms. The fundamental processes involved in HHG in a gaseous medium

can be understood within the framework of the well-known three-step model [47, 48]. This semi-classical framework delineates the gas HHG process through three stages:

- 1. Ionization: Initially, an electron is ionized by the intense laser field and electron tunneling excitation from the valence band to the conduction band.
- 2. Acceleration: Subsequently, the strong laser field imparts energy to the liberated electron (holes, propelling its acceleration away from the ionized molecule.
- 3. Recombination: In the final step, the oscillatory force of the laser field drives the electron back toward the ionized parent molecule. During this process, the electron undergoes recombination with the molecule, releasing the surplus kinetic energy acquired in the second step in the form of a high-energy photon.

However, this model is not directly applicable to solid-state HHG due to differences in density, structure, band structure, Coulomb interactions, and surface effects between gases and solids. Solid-state HHG involves more complex mechanisms, including interactions with the crystalline lattice and surface effects, requiring sophisticated theoretical models for accurate description.

The understanding and control of HHG have been greatly advanced by the development of sophisticated theoretical models, the primary and widely used numerical methods are based on time-dependent Schrödinger equation (TDSE) , semiconductor Bloch equations (SBE) and time-dependent density-functional theory (TDDFT). We won't go in detailed about the theoretical foundations, but just summarize current status and some challenges of present numerical simulations which are yet to be improved or solved.

- The time-dependent-Schrödinger equation (TDSE) model, utilizing both Bloch state basis and Houston state basis, is proficient in examining electronic dynamics within periodic potentials. Nonetheless, simulating the HHG of real solid materials proves challenging, primarily due to the method's reliance on idealized model potentials.

- The semiconductor Bloch equations (**SBE**), improving it from a two-band model to a multiband model allows for a more comprehensive study of real systems, effectively capturing the main features of High-Order Harmonic Generation (HHG) in solids. This enhancement involves incorporating accurate energy bands and transition dipole moments derived from first-principles calculations. Future endeavors should focus on obtaining the correct phase of transition dipole elements. However, a significant challenge persists as current first-principles codes often yield random phases for transition dipole moments.
- The time-dependent density-functional theory (**TDDFT**) model appears to be the ideal approach for straightforwardly studying High-Order Harmonic Generation (HHG) in solids within real coordinate space. However, its main drawback lies in its significant computational time requirements. Additionally, directly and intuitively analyzing the physical processes or mechanisms within the solid-state energy band picture can pose challenges within the TDDFT framework.

Furthermore, properly incorporating Berry curvature in models and understanding its role in solid High-Order Harmonic Generation (HHG) requires more attention in future theoretical simulations. Despite lingering open questions in this evolving field, it's hoped that this review can offer valuable reference points, and ongoing efforts from the relevant community will gradually illuminate the intricacies of solid HHG studies.

1.5 Structure of the Thesis

This thesis is organized as follows: Chapter. 2 first introduces the crystallographic description for typical hexagonal lattice we are studying, then we study the light-induced electron dynamics in 2D materials based on the tight-binding model by time-dependent Schrodinger Equation and quantum master equation. In Chapter. 3 we investigate light-induced electron dynamics in monolayer hexagonal boron nitride under the influence of two-color linearly-polarized laser fields at

frequencies ω and 2ω , by solving the time-dependent Schrödinger equation with a tight-binding model. We start from time-dependent perturbative analysis in the weak field regime, then we expand our results to third-order nonlinear regime and deeply off-resonant highly-nonlinear regime. In Chapter 4, we study THz-induced HHG in graphene with the method described by quantum master equation. The microscopic mechanism of HHG with the quasi-static approximation and the population distribution in the Brillouin zone is described in detail together with its numerical implementation in Chapter 5. We further elucidate the role of the nonequilibrium nature of THz-induced electron dynamics by comparing the nonequilibrium picture in the present work and the thermodynamic picture in the previous work [49]. We explore the possibility of using a THz field to enhance MIR-induced HHG in graphene based on the knowledge gained from Chapter 4. We investigate the dynamics under MIR and THz fields and evaluate the emitted harmonic spectra. As a result of the analysis, we find that coupling via the induced coherence by THz and MIR fields plays an essential role in enhancing MIR-induced HHG, clarifying the importance of the field-induced coherence beyond the simple population effect. Finally, the conclusion of this thesis is summarized in Chapter 6.

CHAPTER 1. INTRODUCTION

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Chapter 2

THEORETICAL FOUNDATIONS

2.1 Crystallography Properties

The hexagonal lattice nanostructure, a fundamental Bravais lattice, manifests as a distinctive geometric arrangement prevalent across a spectrum of materials, owing to its highly efficient packing characteristics. This lattice's spatial configuration profoundly influences the mechanical, electrical, and thermal properties of materials. Understanding lattice structures is crucial for deciphering material behavior across diverse conditions, spanning from semimetals to topological insulators. This significance is particularly noteworthy in the realm of two-dimensional materials. We have selected graphene, exemplifying a semimetal, and hexagonal boron nitride (h-BN), recognized as an insulator, for our discussion on nonlinear optical response on 2d materials.

Graphene, an extraordinary carbon allotrope, showcases a captivating atomic arrangement within a hexagonal lattice nanostructure, depicted in Figure 2.1 (a). Carbon atoms meticulously align in a single layer, forming an exceptional two-dimensional material. The unique atomic-scale hexagonal lattice structure involves each carbon atom intricately bonding through σ -bonds with its three nearest neighbors and a delocalized π -bond. This precise arrangement plays a pivotal role in the formation of a valence band elegantly spanning the entirety of the graphene sheet, making monolayer graphene an outstanding conductor of electricity, and finding applications in electronic devices, sensors, and various fields [50].

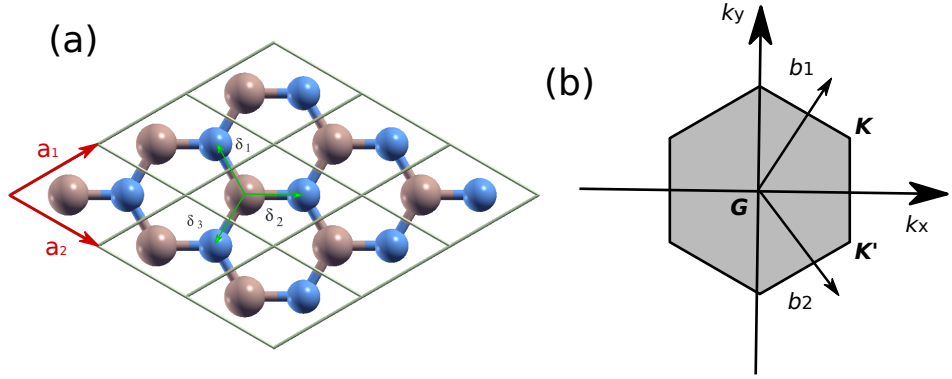


Figure 2.1: (a) Hexagonal lattice, the two triangular sublattices are shown in two different colors. (b) Brillouin zone in momentum space.

Similar to graphene, h-BN also features a hexagonal lattice structure, but with alternating boron and nitrogen atoms forming the hexagons, making it a wide-gap insulator due to inversion symmetry breaking, which is used as a dielectric material in electronics, a substrate for graphene-based devices, and as a solid lubricant.

2.1.1 Structural Parameters

We define the basis of hexagonal lattice primitive vectors $E = (\vec{a}_1, \vec{a}_2)$ as shown in Fig 2.1 (a):

$$\vec{a}_1 = a \begin{pmatrix} \frac{\sqrt{3}}{2} \\ \frac{1}{2} \end{pmatrix}, \vec{a}_2 = a \begin{pmatrix} \frac{\sqrt{3}}{2} \\ -\frac{1}{2} \end{pmatrix}$$

Where a is the lattice constant, for graphene $a = 1.42\text{\AA}$ [50], and for h-BN $a = 2.5\text{\AA}$ [51]. Generate only A sites while sites in B sublattice are generated by $n_1\vec{a}_1 + n_2\vec{a}_2 + \vec{\delta}$, where $\vec{\delta}$ has to be chosen as one of the three nearest-neighbor vectors,

$$\vec{\delta}_1 = a \begin{pmatrix} -\frac{1}{2\sqrt{3}} \\ \frac{1}{2} \end{pmatrix}, \vec{\delta}_2 = a \begin{pmatrix} \frac{1}{\sqrt{3}} \\ 0 \end{pmatrix}, \vec{\delta}_3 = a \begin{pmatrix} -\frac{1}{2\sqrt{3}} \\ -\frac{1}{2} \end{pmatrix}$$

2.1.2 Reciprocal Space

The reciprocal basis $B = (b_1, b_2, b_3)$ is generated using the formula:

$$\vec{b}_k = \frac{2\pi \cdot \vec{a}_i \times \vec{a}_j}{V}$$

i, j, k are circular permutations, $V = \vec{a}_1 \cdot (\vec{a}_2 \times \vec{a}_3)$, presents the mix product between the three vectors, i.e. the volume of the unitary cell. By assuming the lattice primitive vector in the vertical direction of the two-dimensional material plane is infinit, $\vec{a}_3 \rightarrow \infty$, we get the 2D reciprocal vectors as shown in Fig 2.1 (b):

$$\vec{b}_1 = k_D \begin{pmatrix} \frac{1}{2} \\ \frac{\sqrt{3}}{2} \end{pmatrix}, \vec{b}_2 = k_D \begin{pmatrix} \frac{1}{2} \\ -\frac{\sqrt{3}}{2} \end{pmatrix}$$

And with $k_D = \frac{4\pi}{\sqrt{3}a}$. The corresponding Brillouin zone is depicted together with the two high-symmetry points K and K' Fig 2.1 (b). Two inequivalent corners of the Brillouin zone K and K' can be chosen as follows:

$$K = k_D \left(\frac{1}{2}, \frac{1}{2\sqrt{3}} \right), \quad K' = k_D \left(\frac{1}{2}, -\frac{1}{2\sqrt{3}} \right)$$

2.2 Tight-binding Approach

In this section, we delve into the fundamental principles of the tight-binding approach, with a particular focus on the nearest neighbor tight-binding model. This approach is essential for understanding the electronic properties of materials and is a crucial component of graphene's electronic structure analysis.

The foundation of the tight-binding approach is rooted in the Bloch theorem, which is satisfied by the tight-binding function

$$\Phi_{\alpha,\beta}(\mathbf{r}, \mathbf{k}) = \frac{1}{\sqrt{N}} \sum_{\mathbf{R}} e^{i\mathbf{k}\cdot\mathbf{R}} \phi_{\alpha,\beta}(\mathbf{r} - \mathbf{R}_{\alpha,\beta}), \alpha = A \text{ or } B \quad (2.1)$$

Here, β is the general atomic orbitals index, For simplicity, we only consider the contribution from the outermost valence band electron's orbital and omit β in the following discussion. N represents the number of unit cells, and $\phi_{\alpha}(\mathbf{r} - \mathbf{R}_{\alpha})$

denotes the orbital function of an electron at cell \mathbf{R} in sublattice α . In the context of a honeycomb lattice, we focus on the nearest neighbor approximation. This approximation asserts that an atom in sublattice A only interacts with its three closest neighbor atoms in sublattice B. This simplification is particularly useful for understanding the interactions between electrons bound to non-equivalent atoms.

The Hamiltonian for this nearest-neighbor interaction is expressed as:

$$\hat{H}_{AB} = \frac{1}{N} \sum_{\mathbf{R}_A} \sum_{\mathbf{R}_B} e^{i\mathbf{k}(\mathbf{R}_B - \mathbf{R}_A)} \langle \phi_A(\mathbf{r} - \mathbf{R}_A) | \hat{H} | \phi_B(\mathbf{r} - \mathbf{R}_B) \rangle \quad (2.2)$$

Due to the translational invariance in a Bravais lattice, the summation over each atom in a sublattice occurs N times, simplifying the expression to:

$$\hat{H}_{AB} = \sum_{\mathbf{R}_A} e^{i\mathbf{k}(\mathbf{R}_B - \mathbf{R}_A)} \langle \phi_A(\mathbf{r} - \mathbf{R}_A) | \hat{H} | \phi_B(\mathbf{r} - \mathbf{R}_B) \rangle \quad (2.3)$$

We transform the Hamiltonian of real space into the momentum space representation, and then the tight-binding Hamiltonian under the matrix representation is:

$$\hat{H} = \begin{pmatrix} \epsilon_A & t_0 f(\mathbf{k}) \\ t_0 f(\mathbf{k})^* & \epsilon_B \end{pmatrix} \quad (2.4)$$

Here ϵ_A and ϵ_B are the on-site energies of electrons on the nearest neighbor atoms, t_0 presents the hopping parameter:

$$t_0 = \langle \phi_A(\mathbf{r} - \mathbf{R}_A) | \hat{H} | \phi_B(\mathbf{r} - \mathbf{R}_A - \vec{\delta}_i) \rangle \quad (i = 1, 2, 3)$$

The three nearest neighbor hoping strength here are identical here due to the hexagonal lattice symmetry. For graphene, we set ϵ_A and ϵ_B to 0, and $t_0 = 2.8 \text{ eV}$ in accordance with the previous work [50]. For h-BN ϵ_B and ϵ_N denote the on-site energies for boron and nitrogen sites, respectively.

$$\hat{H}(\mathbf{k}) = \begin{pmatrix} \epsilon_B & t_0 f(\mathbf{k}) \\ t_0 f(\mathbf{k})^* & \epsilon_N \end{pmatrix}, \quad (2.5)$$

We set ϵ_B to 3.34 eV and ϵ_N to -2.56 eV and t_0 to 2.6 eV computed with the first-

principles calculations [52], the band gap $E_g = \epsilon_b - \epsilon_n$ equals 5.9 eV.

The off-diagonal terms of the tight-binding Hamiltonian 2.4:

$$\begin{aligned} f(\mathbf{k}) &= e^{i\mathbf{k}\vec{\delta}_1} + e^{i\mathbf{k}\vec{\delta}_2} + e^{i\mathbf{k}\vec{\delta}_3} \\ &= e^{-\frac{ia k_x}{\sqrt{3}}} + 2e^{\frac{ia k_x}{2\sqrt{3}}} \cos\left(\frac{a}{2}k_y\right) \end{aligned} \quad (2.6)$$

Solving the stationary Schrödinger equation using matrix diagonalization:

$$\hat{H}_{\mathbf{k}}|\phi_{b\mathbf{k}}\rangle = \epsilon_{b\mathbf{k}}|\phi_b\rangle, \quad (2.7)$$

We get the eigenenergy of Hamiltonian from 2.4, where b is a band index, $|\phi_{b\mathbf{k}}\rangle$ is an eigenstate, and $\epsilon_{b\mathbf{k}}$ corresponds to the eigenenergy. As the Hamiltonian is a 2-by-2 matrix in this work, the band index b denotes either a conduction ($b = c$) or valence ($b = v$) state.

$$\epsilon_{b\mathbf{k}} = E_0 \pm \frac{1}{2} \sqrt{E_g^2 + 4t_0^2|f|^2} \quad (2.8)$$

$E_0 = \frac{\epsilon_A + \epsilon_B}{2}$ and $E_g = \epsilon_b - \epsilon_n$ is the energy gap. For graphene, $\epsilon_A = \epsilon_B = 0$ the band gap equals 0. Their corresponding eigenvectors are:

$$|\phi_b\rangle = \begin{pmatrix} \frac{E_g \pm \sqrt{E_g^2 + 4t_0^2|f|^2}}{2t_0 f^*} \\ 1 \end{pmatrix} \quad (2.9)$$

2.3 Electron Dynamics

Consider the crystal under the homoheneous electric field \mathbf{E} , the wavelength of the fields is much longer than the spatial scale of the electron dynamics, so-called long wavelength approximation, or dipole approximation. The electric field enter the solid system through a uniform vector potential $\mathbf{A}(t)$ under Peierls substitution[53]. The time-dependent Hamiltonian is written as

$$\hat{H}(t) = \frac{[\hat{\mathbf{p}} + e\mathbf{A}(t)]^2}{2m} + V(\mathbf{r}) \quad (2.10)$$

Transforming to the \mathbf{k} -space representation, we have

$$\hat{H}(\mathbf{k}, t) = \hat{H}\left(\mathbf{k} + \frac{e}{\hbar} \mathbf{A}(t)\right) \quad (2.11)$$

where \mathbf{k} denotes the Bloch wavevector. The vector potential $\mathbf{A}(t)$ is related to the applied electric field $\mathbf{E}(t)$ as $\mathbf{E}(t) = -d\mathbf{A}(t)/dt$, and it is included in the Hamiltonian as the wavevector shift $\mathbf{k} \rightarrow \mathbf{k} + e\mathbf{A}(t)/\hbar$ via the Peierls substitution [53].

To study the nonlinear photocarrier injection, we need to introduce time-dependent Schrödinger equation (**TDSE**) for close quantum systems since the injection carriers need to be discussed without considering dissipation. On the other stage, HHG induced by intensified THz or MIR laser has strong relaxation which can not be ignored so we introduce quantum master equation for HHG progress under intensified long-term pulse. We will discuss the progress solving quantum dynamics in close and open system separated in the next sections to get nonlinear response current, which is the main observable for both theory and experiment.

2.3.1 Time-dependent Schrödinger Equation

The light-induced electron dynamics can be described by solving the following **TDSE** at each \mathbf{k} -point:

$$i\hbar \frac{d}{dt} |\psi_{\mathbf{k}}(t)\rangle = \hat{H}\left(\mathbf{k} + \frac{e\mathbf{A}(t)}{\hbar}\right) |\psi_{\mathbf{k}}(t)\rangle, \quad (2.12)$$

$|\psi_{\mathbf{k}}(t)\rangle$ is a single-particle electronic wavefunction at \mathbf{k} . Solving this time-dependent Schrödinger equation (**TDSE**) is an initial value problem. In two-band systems, usually the ground state $\psi_{\mathbf{k}}(0)$ is used as the initial state occupied at the valence band. Various numerical methods can be chosen for doing the time propagation, here we split the propagation into short-time propagation using the composition property rely on a sufficiently small Δt , $t' = t + \Delta t$:

$$|\psi_{\mathbf{k}}(t')\rangle = \exp\left[-i \int_t^{t'} d\tau \hat{H}(\tau)\right] |\psi_{\mathbf{k}+\mathbf{A}(t)}\rangle \quad (2.13)$$

Since Δt is sufficiently small, the exponential mid-point propagator:

$$\exp \left[-i \int_t^{t'} d\tau \hat{H}(\tau) \right] \approx \exp \{ -i \Delta t \hat{H}(t + \Delta t/2) \} \quad (2.14)$$

We approximate the exponential using a Taylor expansion to fourth-order:

$$\exp \{ A \} = \sum_{k=0}^{\infty} \frac{1}{k!} A^k \quad (2.15)$$

Once the time-evolution of the wavefunctions, $|\psi_{\mathbf{k}}(t)\rangle$ is computed, the current induced in the matter can be further evaluated with

$$\mathbf{J}_{\mathbf{k}}(t) = \frac{1}{(2\pi)^2} \int_{BZ} d\mathbf{k} \langle \psi_{\mathbf{k}}(t) | \hat{\mathbf{J}}_{\mathbf{k}}(t) | \psi_{\mathbf{k}}(t) \rangle. \quad (2.16)$$

Here, $\hat{\mathbf{J}}_{\mathbf{k}}(t)$ is the current operator, and it is defined as

$$\hat{\mathbf{J}}_{\mathbf{k}}(t) = \frac{\partial}{\partial \mathbf{k}} \hat{H} \left(\mathbf{k} + \frac{e\mathbf{A}(t)}{\hbar} \right) = -t_0 \begin{pmatrix} 0 & \frac{\partial f(\mathbf{k}+\mathbf{A})}{\partial \mathbf{A}} \\ \frac{\partial f^*(\mathbf{k}+\mathbf{A})}{\partial \mathbf{A}} & 0 \end{pmatrix}, \quad (2.17)$$

where $\frac{\partial f(\mathbf{k})}{\partial \mathbf{k}}$ is given by

$$\frac{\partial f(\mathbf{k})}{\partial \mathbf{k}} = i\vec{\delta}_1 e^{i\mathbf{k}\cdot\vec{\delta}_1} + i\vec{\delta}_2 e^{i\mathbf{k}\cdot\vec{\delta}_2} + i\vec{\delta}_3 e^{i\mathbf{k}\cdot\vec{\delta}_3}. \quad (2.18)$$

We can also analyze the population distribution of photocarriers induced by the laser fields. To achieve this, we compute the conduction population distribution by projecting onto the eigenstates of the Hamiltonian defined as:

$$\hat{H}_k |\phi_{bk}\rangle = \epsilon_{bk} |\phi_{bk}\rangle, \quad (2.19)$$

where b is a band index, $|\phi_{bk}\rangle$ is an eigenstate, and ϵ_{bk} corresponds to the eigenvalue. As the Hamiltonian is a 2-by-2 matrix in this work, the band index b denotes either a conduction ($b = c$) or valence ($b = v$) state.

Using eigenstates defined with Eq. (2.19), the conduction population distribution n_{ck} after the laser irradiation can be evaluated as:

$$n_{ck} = |\langle \phi_{ck} | \psi_k(t_F) \rangle|^2, \quad (2.20)$$

where t can be an instantaneous time during or after the laser field. By imposing the normalization of $|\phi_{bk}\rangle$ and $|\psi_k(t)\rangle$, the computed conduction population satisfies $0 \leq n_{ck} \leq 1$. It is important to note that, in the present theoretical setup, the conduction population is a constant of motion after laser irradiation since any relaxation processes are not considered.

2.3.2 Quantum Master Equation

In contrast to closed quantum systems, which are entirely isolated from external influences and can be adequately described by the Schrödinger equation, the quantum master equation is typically used in the context of the time evolution of an open quantum system, where the system of interest is susceptible to exchanges of energy, particles, or information with its external environment. To expound processes such as relaxation, dephasing, and thermalization of the nonlinear response experiments, the quantum master equation is predominantly employed which is salient in the realm of open quantum systems.

We describe the light-induced electron dynamics in graphene with the following quantum master equation [54–57]:

$$\frac{d}{dt}\rho_{\mathbf{k}}(t) = \frac{1}{i\hbar} [\hat{H}_{\mathbf{k}+e\mathbf{A}(t)/\hbar}, \rho_{\mathbf{k}}(t)] + \hat{D}[\rho_{\mathbf{k}}(t)], \quad (2.21)$$

$\rho_{\mathbf{k}}(t)$ is the reduced density matrix at \mathbf{k} . The quantum master equation delineates the dynamical evolution of the density matrix associated with the quantum system, which encompasses both pure and mixed quantum states. To elucidate the impact of dissipation, we formulate the relaxation operator, denoted as $\hat{D}[\rho_{\mathbf{k}}(t)]$, within the framework of Eq.(2.21) employing the relaxation time approximation[58] and employing the Houston basis [59, 60]. The Houston states, $|u_{b\mathbf{k}}^H(t)\rangle$, are characterized as eigenstates of the instantaneous Hamiltonian, expressed as:

$$\hat{H}_{\mathbf{k}+e\mathbf{A}(t)/\hbar}|u_{b\mathbf{k}}^H(t)\rangle = \epsilon_{b,\mathbf{k}+e\mathbf{A}(t)/\hbar}|u_{b\mathbf{k}}^H(t)\rangle \quad (2.22)$$

The expansion of the reduced density matrix can then be carried out using the Houston states.

$$\rho_{\mathbf{k}}(t) = \sum_{bb'} \rho_{bb',\mathbf{k}}(t) |u_{b\mathbf{k}}^H(t)\rangle \langle u_{b'\mathbf{k}}^H(t)|, \quad (2.23)$$

where $\rho_{bb',\mathbf{k}}(t)$ are the expansion coefficients. On the basis of the Houston state expansion, we define the relaxation operator [54, 55, 61] as

$$\hat{D}[\rho_{\mathbf{k}}(t)] = - \sum_b \frac{\rho_{bb,\mathbf{k}}(t) - f^{FD}(\epsilon_{b,\mathbf{k}+e\mathbf{A}(t)/\hbar}, T_e, \mu)}{T_1} |u_{b\mathbf{k}}^H(t)\rangle \langle u_{b\mathbf{k}}^H(t)| \quad (2.24)$$

$$- \sum_{b \neq b'} \frac{\rho_{bb',\mathbf{k}}(t)}{T_2} |u_{b\mathbf{k}}^H(t)\rangle \langle u_{b'\mathbf{k}}^H(t)|, \quad (2.25)$$

T_1 is the longitudinal relaxation time, T_2 is the transverse relaxation time, and $f^{FD}(\epsilon)$ is the Fermi–Dirac distribution:

$$f^{FD}(\epsilon, T_e, \mu) = \frac{1}{e^{(\epsilon-\mu)/k_B T_e} + 1}. \quad (2.26)$$

When $t = 0$, the electron system are identical fermions in thermodynamic equilibrium, the average number of fermions in a single-particle state is given by the Fermi–Dirac distribution. μ is the chemical potential, and T_e is the electron temperature.

In the following discussion, we set the longitudinal relaxation time T_1 to 100 fs and the transverse relaxation time T_2 to 20 fs in accordance with the previous works [54–57]. The electron temperature T_e is set to 300 K unless stated otherwise. The chemical potential μ is treated as a tunable parameter to study the effect of doping.

We directly solve the quantum master equation, Eq. (2.21), in the time domain by employing the Runge–Kutta method without any approximation. The electric current is obtained by employing the time-dependent density matrix $\rho_{\mathbf{k}}(t)$, which evolves according to Eq. (2.21):

$$\mathbf{J}(t) = \frac{2}{(2\pi)^2} \int d\mathbf{k} \text{Tr} [\hat{\mathbf{J}}_{\mathbf{k}}(t) \rho_{\mathbf{k}}(t)], \quad (2.27)$$

where $\hat{\mathbf{J}}_{\mathbf{k}}(t)$ is the current operator defined as

$$\hat{\mathbf{J}}_{\mathbf{k}}(t) = -\frac{\partial H(\mathbf{k} + e\mathbf{A}(t)/\hbar)}{\partial \mathbf{A}(t)}. \quad (2.28)$$

The intraband component of the current is the dominant component for nonlinear current contribute HHG progress which will be discussed in the Chapter 4, we can get the intraband current by:

$$\mathbf{J}_{\mathbf{k}}^{\text{intra}}(t) = \sum_{b=v,c} \frac{(-2)}{(2\pi)^2} \frac{e}{\hbar} \times \int d\mathbf{k} \frac{\partial \epsilon_{b,\mathbf{k}+e\mathbf{A}(t)/\hbar}}{\partial \mathbf{k}} n_{b,\mathbf{k}+e\mathbf{A}(t)/\hbar},$$

where the band population $n_{b,\mathbf{k}+e\mathbf{A}(t)/\hbar}$ is defined with the Houston states of the Hamiltonian $|u_{b,\mathbf{k}}^H(t)\rangle$ computed from Eq 2.22, which can be useful for the microscopic analysis in the folowing chapters:

$$n_{b,\mathbf{k}+e\mathbf{A}(t)/\hbar}(t) = \langle u_{b,\mathbf{k}}^H(t) | \rho_{\mathbf{k}}(t) | u_{b,\mathbf{k}}^H(t) \rangle \quad (2.29)$$

Chapter 3

CO-LINEAR POLARIZATION INDUCED PHOTOVOLTAIC EFFECT

Addressing the photovoltaic effect within the perturbative regime has garnered significant attention, particularly in the exploration of dc current injection using two-color linearly polarized light [62–68]. Early investigations have underscored the complex interaction between a fundamental frequency, denoted as ω , and its second harmonic, 2ω [62–64]. A notable study by Jimenez-Galan et al. [20] utilized deeply off-resonant bi-circular laser fields with ω and 2ω to generate a substantial population imbalance in the Brillouin zone. However, it is worth noting that, in principle, linearly polarized light with two frequencies is sufficient to break time-reversal symmetry.

In this chapter, we first theoretically explore the phenomenon of dc-current injection and the generation of population imbalance through the application of two-color linearly polarized laser fields with frequencies ω and 2ω based on time-dependent perturbation analysis. Then we look into light-induced electron dynamics in a typical two-dimensional insulator, *h*-BN, based on a simple tight-binding approximation in a perturbative resonant regime.

In our quantum dynamics simulations, we further uncover that ballistic current can be induced even in the deeply off-resonant regime with two-color linearly polarized light. Consequently, efficient injection of dc-current and the creation of a substantial population imbalance can be realized by employing two-color linearly polarized laser fields with frequencies ω and 2ω , without relying

on the ellipticity of light. These findings offer a potential pathway for achieving ultrafast and efficient control of electron population in matter using multi-color linearly polarized light, opening new avenues for exploring the frontiers of quantum dynamics and optoelectronic applications.

3.1 Time-dependent Perturbative Analysis on QuI

Time-dependent perturbation theory provides a form for describing the response of quantum systems to time-varying external fields, makes it well-suited for analyzing the interaction of materials with intense electromagnetic radiation. Exploring the photovoltaic effect within the perturbative regime has led to a notable focus on elucidating the injection of direct current (dc) through the use of two-color linearly polarized light [62–68]. Here, we investigate the nonlinear photocarrier injection process via the time-dependent perturbation analysis. Under adiabatic basis representation described in Appendix A, one can rewrite the equation of motion for the coefficient vector $c_{\mathbf{k}}(t)$ after expansion as

$$i \frac{d}{dt} \mathbf{c}_{\mathbf{k}}(t) = \mathcal{H}(t) \mathbf{c}_{\mathbf{k}}(t). \quad (3.1)$$

$$\mathcal{H}(t) = i \mathbf{E}(t) \cdot \begin{pmatrix} 0 & M_{12} \\ M_{21} & 0 \end{pmatrix} \quad (3.2)$$

$$M_{12} = e^{-i \int_0^t dt' \Delta \epsilon_{cv, \mathbf{k}+\mathbf{A}(t')} + i \Delta \phi_{cv, \mathbf{k}}^g(t)} \left\langle u_{v, \mathbf{k}+\mathbf{A}(t)} \left| \frac{\partial u_{c, \mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right. \right\rangle \quad (3.3)$$

$$M_{21} = e^{-i \int_0^t dt' \Delta \epsilon_{vc, \mathbf{k}+\mathbf{A}(t')} + i \Delta \phi_{vc, \mathbf{k}}^g(t)} \left\langle u_{c, \mathbf{k}+\mathbf{A}(t)} \left| \frac{\partial u_{v, \mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right. \right\rangle \quad (3.4)$$

$\mathbf{c}_{\mathbf{k}}(t) = \begin{pmatrix} c_{v, \mathbf{k}}(t) \\ c_{c, \mathbf{k}}(t) \end{pmatrix}$ is the coefficient vector, $\Delta \epsilon_{bb', \mathbf{k}+\mathbf{A}(t)}$ is defined by the difference of the single-particle energies as $\epsilon_{b, \mathbf{k}+\mathbf{A}(t)} - \epsilon_{b', \mathbf{k}+\mathbf{A}(t)}$. For simplicity, here assume

3.1. TIME-DEPENDENT PERTURBATIVE ANALYSIS ON QUI

that the contribution from the geometric phases, $\Delta\phi_{cv,\mathbf{k}}^g(t)$ is zero. Then expand the Hamiltonian in Eq. (3.1) up to the second order of the field $\mathbf{A}(t)$ as

$$\mathcal{H}(t) \approx \mathcal{H}^{(1)}(t) + \mathcal{H}_{dyn}^{(2)}(t) + \mathcal{H}_{dip}^{(2)}(t), \quad (3.5)$$

$$\mathcal{H}^{(1)}(t) = i\mathbf{E}(t) \cdot \begin{pmatrix} 0 & e^{-i\Delta\epsilon_{cv,\mathbf{k}}t} \left\langle u_{v,\mathbf{k}} \middle| \frac{\partial u_{c,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle \\ e^{-i\Delta\epsilon_{vc,\mathbf{k}}t} \left\langle u_{c,\mathbf{k}} \middle| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle & 0 \end{pmatrix}, \quad (3.6)$$

$\mathcal{H}_{dyn}^{(2)}(t)$ originates from the modification of the dynamical phase factor:

$$e^{-i \int_0^t dt' \Delta\epsilon_{vc,\mathbf{k}+\mathbf{A}(t')}} \quad (3.7)$$

$\mathcal{H}_{dyn}^{(2)}(t)$ is given by:

$$\mathcal{H}_{dyn}^{(2)}(t) = \mathbf{E}(t) \cdot \begin{pmatrix} 0 & M_{dyn,12} \\ M_{dyn,21} & 0 \end{pmatrix}, \quad (3.8)$$

$$M_{dyn,12} = \frac{\partial \Delta\epsilon_{cv,\mathbf{k}}}{\partial \mathbf{k}} \cdot \left(\int_0^t dt' \mathbf{A}(t') \right) e^{-i\Delta\epsilon_{cv,\mathbf{k}}t} \left\langle u_{v,\mathbf{k}} \middle| \frac{\partial u_{c,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle \quad (3.9)$$

$$M_{dyn,21} = \frac{\partial \Delta\epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \cdot \left(\int_0^t dt' \mathbf{A}(t') \right) e^{-i\Delta\epsilon_{vc,\mathbf{k}}t} \left\langle u_{c,\mathbf{k}} \middle| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle \quad (3.10)$$

and $\mathcal{H}_{dip}^{(2)}(t)$ originates from the modification of the dipole matrix elements:

$$i \left\langle u_{v,\mathbf{k}+\mathbf{A}(t)} \middle| \frac{\partial u_{c,\mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right\rangle \quad (3.11)$$

$\mathcal{H}_{dip}^{(2)}(t)$ is given by:

$$\mathcal{H}_{dip}^{(2)}(t) = \begin{pmatrix} 0 & M_{dip,12} \\ M_{dip,12} & 0 \end{pmatrix}. \quad (3.12)$$

$$M_{dip,12} = e^{-i\Delta\epsilon_{cv,\mathbf{k}}t} \mathbf{A}(t) \cdot \frac{\partial \left\langle u_{v,\mathbf{k}} \middle| i\mathbf{E}(t) \cdot \frac{\partial u_{c,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle}{\partial \mathbf{k}} \quad (3.13)$$

$$M_{dip,21} = e^{-i\Delta\epsilon_{vc,\mathbf{k}}t} \mathbf{A}(t) \cdot \frac{\partial \left\langle u_{c,\mathbf{k}} \middle| i\mathbf{E}(t) \cdot \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle}{\partial \mathbf{k}} \quad (3.14)$$

Hereafter, we analyze the photocarrier injection process based on this perturbative expansion of the Hamiltonian. Use perturbation expansion of Eq. (3.1):

$$i\frac{d}{dt}(\mathbf{c}_{\mathbf{k}}^{(0)}(t) + \mathbf{c}_{\mathbf{k}}^{(1)}(t) + \mathbf{c}_{\mathbf{k}}^{(2)}(t)) = (\mathcal{H}_{\mathbf{k}}^{(0)} + \mathcal{H}_{\mathbf{k}}^{(1)}(t) + \mathcal{H}_{\mathbf{k}}^{(2)}(t))(\mathbf{c}_{\mathbf{k}}^{(0)}(t) + \mathbf{c}_{\mathbf{k}}^{(1)}(t) + \mathbf{c}_{\mathbf{k}}^{(2)}(t)) \quad (3.15)$$

Under initial condition $\mathcal{H}_{\mathbf{k}}^{(0)}(t=0) = 0$, $\mathbf{c}_{\mathbf{k}}^{(0)}(t=0) = \begin{pmatrix} 1 \\ 0 \end{pmatrix}$, the first ($c_{c,\mathbf{k}}^{(1)}(t)$) and second-order coefficient vectors ($c_{c,\mathbf{k},dyn}^{(2)}(t)$, $c_{c,\mathbf{k},dip}^{(2)}(t)$) for the conduction band can be written as:

$$c_{c,\mathbf{k}}^{(1)}(t) = \left\langle u_{c,\mathbf{k}} \middle| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle \int_0^t dt' e^{-i\Delta\epsilon_{vc,\mathbf{k}}t'} \mathbf{E}_1(t') \quad (3.16)$$

$$c_{c,\mathbf{k},dyn}^{(2)}(t) = \frac{1}{i} \frac{\partial \Delta\epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left\langle u_{c,\mathbf{k}} \middle| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle \int_0^t dt' e^{-i\Delta\epsilon_{vc,\mathbf{k}}t'} \mathbf{E}_2(t') \int_0^{t'} dt'' \mathbf{A}_2(t'') \quad (3.17)$$

$$c_{c,\mathbf{k},dip}^{(2)}(t) = \frac{\partial \left\langle u_{c,\mathbf{k}} \middle| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle}{\partial \mathbf{k}} \int_0^t dt' e^{-i\Delta\epsilon_{vc,\mathbf{k}}t'} \mathbf{E}_2(t') \mathbf{A}_2(t') \quad (3.18)$$

Under two-color linearly polarized light, here we consider the perturbation by the external linearly polarized vector potential for \vec{e} -direction, assuming that the perturbation only exists time between 0 and $2T_0$ under the Gaussian distribution:

$$f(t) = e^{-\frac{(t-T_0)^2}{2\sigma^2}} \quad (3.19)$$

3.1. TIME-DEPENDENT PERTURBATIVE ANALYSIS ON QUI

$$\mathbf{A}_1(t) = A_1 \vec{e} \cos[2\omega(t - T_0) + \phi] e^{-\frac{(t-T_0)^2}{2\sigma^2}}, \quad (3.20)$$

$$\mathbf{A}_2(t) = A_2 \vec{e} \cos[\omega(t - T_0)] e^{-\frac{(t-T_0)^2}{2\sigma^2}} \quad (3.21)$$

w is the carrier frequency of the field, \vec{e} represents a unit vector along the polarization direction of the laser field. We assume $T_0 \gg 0$ and $\sigma \gg 0$ to make the approximation:

$$\int_0^{t'} A_2 \vec{e} \cos[\omega(t - T_0)] e^{-\frac{(t-T_0)^2}{2\sigma^2}} dt = \frac{A_2 \vec{e}}{\omega} \sin[\omega(t' - T_0)] e^{-\frac{(t'-T_0)^2}{2\sigma^2}} \quad (3.22)$$

The corresponding external electric field $\mathbf{E}(t) = -d\mathbf{A}(t)/dt$ can be written as the following pulsed form:

$$\mathbf{E}_1(t) = 2\omega A_1 \vec{e} \sin[2\omega(t - T_0) + \phi] e^{-\frac{(t-T_0)^2}{2\sigma^2}}, \quad (3.23)$$

$$\mathbf{E}_2(t) = \omega A_2 \vec{e} \sin[\omega(t - T_0)] e^{-\frac{(t-T_0)^2}{2\sigma^2}} \quad (3.24)$$

Now, let's express this in terms of the first-order coefficient vector is:

$$\begin{aligned} c_{c,\mathbf{k}}^{(1)}(t) &= \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \int_0^t dt' e^{-i\Delta\epsilon_{vc,\mathbf{k}} t'} 2\omega A_1 \vec{e} \cdot \sin[2\omega(t' - T_0) + \phi] e^{-\frac{(t'-T_0)^2}{2\sigma^2}} \\ &= \frac{A_1 \vec{e} \omega}{i} \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \int_0^t dt' e^{-i\Delta\epsilon_{vc,\mathbf{k}} t' - \frac{(t'-T_0)^2}{2\sigma^2}} \cdot (e^{i[2\omega(t' - T_0) + \phi]} - e^{-i[2\omega(t' - T_0) + \phi]}) \end{aligned} \quad (3.25)$$

We consider the population distribution after the laser pulse, so we replace the integra \int_0^t by $\int_{-\infty}^{\infty}$. Under the Gaussian integral,

$$\int_{-\infty}^{\infty} e^{-x^2} dx = \sqrt{\pi} \quad (3.26)$$

we get:

$$c_{c,\mathbf{k}}^{(1)}(t) = \frac{A_1 \vec{e} \omega \cdot \sigma \sqrt{2\pi}}{i} \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \cdot e^{-i\Delta\epsilon_{vc,\mathbf{k}} T_0} [e^{-\frac{1}{2}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2 - i\phi} - e^{-\frac{1}{2}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2 + i\phi}] \quad (3.27)$$

Similarly to the first-order perturbation coefficient vector's derivation, the second-

order coefficient $c_{c,\mathbf{k}}^{(2)}(t)$ can be written as:

$$c_{c,\mathbf{k},dyn}^{(2)}(t) = \frac{(A_2 \vec{e})^2 \cdot \sigma \sqrt{\pi}}{2i} \frac{\partial \Delta \epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \cdot e^{-i \Delta \epsilon_{vc,\mathbf{k}} T_0} \\ \cdot [e^{-\frac{\sigma^2}{4} \Delta \epsilon_{vc,\mathbf{k}}^2} - \frac{1}{2} (e^{-\frac{\sigma^2}{4} (\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} + e^{-\frac{\sigma^2}{4} (\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2})] \quad (3.28)$$

$$c_{c,\mathbf{k},dip}^{(2)}(t) = \frac{(A_2 \vec{e})^2 \omega \cdot \sigma \sqrt{\pi}}{4i} \cdot \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} \cdot e^{-i \Delta \epsilon_{vc,\mathbf{k}} T_0} [e^{-\frac{\sigma^2}{4} (\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4} (\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2}] \quad (3.29)$$

This completes the derivation of the population of the conduction band after the laser pulse:

$$|c_{c,\mathbf{k}}(t)|^2 = |c_{c,\mathbf{k}}^{(1)}(t)|^2 + |c_{c,\mathbf{k},dyn}^{(2)}(t)|^2 + |c_{c,\mathbf{k},dip}^{(2)}(t)|^2 \\ + c_{c,\mathbf{k}}^{(1)*} c_{c,\mathbf{k},dyn}^{(2)}(t) + c.c. \\ + c_{c,\mathbf{k}}^{(1)*} c_{c,\mathbf{k},dip}^{(2)}(t) + c.c. \\ + c_{c,\mathbf{k},dyn}^{(2)*} c_{c,\mathbf{k},dip}^{(2)}(t) + c.c. \quad (3.30)$$

c.c. represents complex conjugate. We consider a prototypical two-dimensional insulator, monolayer hexagonal boron-nitride (**h-BN**). For the 2-band **h-BN**-tight-binding model, the inversion symmetric is breaking:

$$u_{\mathbf{k}}(-\vec{r}) \neq u_{-\mathbf{k}}(\vec{r}) \quad (3.31)$$

We apply the time-reversal relation for the derivation:

$$u_{\mathbf{k}}^*(\vec{r}) = u_{-\mathbf{k}}(\vec{r}) \quad (3.32)$$

To compare the $|c_{c,-\mathbf{k}}(t)|^2$, the absolute value of the coefficient becomes:

$$|c_{c,\mathbf{k}}^{(1)}(t)|^2 = \left| \frac{A_1 \vec{e} \omega \cdot \sigma \sqrt{2\pi}}{i} \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \right|^2 \\ \cdot [e^{-(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2} + e^{-(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2} - 2e^{-\frac{1}{2}((\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2 + (\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2)} \cos(2\phi)] \\ = |c_{c,-\mathbf{k}}^{(1)}(t)|^2 \quad (3.33)$$

3.1. TIME-DEPENDENT PERTURBATIVE ANALYSIS ON QUI

$$\begin{aligned}
|c_{c,\mathbf{k},dyn}^{(2)}(t)|^2 &= \left| \frac{(A_2 \vec{e})^2 \cdot \sigma \sqrt{\pi}}{2} \frac{\partial \Delta \epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \right. \\
&\quad \cdot [e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - \frac{1}{2}(e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} + e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2})] |^2 \\
&= |c_{c,-\mathbf{k},dyn}^{(2)}(t)|^2
\end{aligned} \tag{3.34}$$

$$\begin{aligned}
|c_{c,\mathbf{k},dip}^{(2)}(t)|^2 &= \left| \frac{(A_2 \vec{e})^2 \omega \cdot \sigma \sqrt{\pi}}{4} \cdot \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} [e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2}] \right|^2 \\
&= |c_{c,-\mathbf{k},dip}^{(2)}(t)|^2
\end{aligned} \tag{3.35}$$

The interference terms are:

$$\begin{aligned}
c_{c,\mathbf{k}}^{(1)}(t)^* c_{c,\mathbf{k},dyn}^{(2)}(t) + c.c. &= \sqrt{2} A_1 A_2^2 \vec{e} \omega \cdot \sigma^2 \pi \frac{\partial \Delta \epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left| \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \right|^2 \\
&\quad \cdot [e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - \frac{1}{2}(e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} + e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2})] \\
&\quad \cdot [e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2} - e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2}] \cos \phi \\
&= -(c_{c,-\mathbf{k}}^{(1)}(t)^* c_{c,-\mathbf{k},dyn}^{(2)}(t) + c.c.)
\end{aligned} \tag{3.36}$$

$$\begin{aligned}
c_{c,\mathbf{k}}^{(1)}(t)^* c_{c,\mathbf{k},dip}^{(2)}(t) + c.c. &= \frac{\sqrt{2} A_1 A_2^2 \vec{e} \omega^2 \cdot \sigma^2 \pi}{4} \cdot (e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2}) \\
&\quad \cdot (\left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \Big| u_{c,\mathbf{k}} \right\rangle \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} [e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2 + i\phi} - e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2 - i\phi}] \\
&\quad + \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \Big| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} \left\langle u_{c,\mathbf{k}} \Big| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle [e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2 - i\phi} - e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2 + i\phi}])
\end{aligned} \tag{3.37}$$

Here under the time-reversal relation, $u_{\mathbf{k}}^*(\vec{r}) = u_{-\mathbf{k}}(\vec{r})$, so for $-\mathbf{k}$, we have:

$$\left\langle \frac{\partial u_{v,-\mathbf{k}}}{\partial(-\mathbf{k})} \Big| u_{c,-\mathbf{k}} \right\rangle \frac{\partial \left\langle u_{c,-\mathbf{k}} \left| \frac{\partial u_{v,-\mathbf{k}}}{\partial(-\mathbf{k})} \right. \right\rangle}{\partial(-\mathbf{k})} = \left\langle \frac{\partial u_{v,\mathbf{k}}^*}{\partial(-\mathbf{k})} \Big| u_{c,\mathbf{k}}^* \right\rangle \frac{\partial \left\langle u_{c,\mathbf{k}}^* \left| \frac{\partial u_{v,\mathbf{k}}^*}{\partial(-\mathbf{k})} \right. \right\rangle}{\partial(-\mathbf{k})} \tag{3.38}$$

$$= - \left\langle u_{c,\mathbf{k}} \Big| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right\rangle \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \Big| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} \tag{3.39}$$

The interference terms for central symmetry point $-\mathbf{k}$ can be written as:

$$\begin{aligned}
 c_{c,-\mathbf{k}}^{(1)}(t)^* c_{c,-\mathbf{k},dip}^{(2)}(t) + c.c. &= \frac{\sqrt{2}A_1 A_2^2 \vec{e} \omega^2 \cdot \sigma^2 \pi}{4} \cdot (e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2}) \\
 &\cdot \left(- \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} [e^{-\frac{1}{2}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2 + i\phi} - e^{-\frac{1}{2}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2 - i\phi}] \right. \\
 &\left. - \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle [e^{-\frac{1}{2}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2 - i\phi} - e^{-\frac{1}{2}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2 + i\phi}] \right) \\
 \end{aligned} \tag{3.40}$$

$$\begin{aligned}
 c_{c,\mathbf{k},dyn}^{(2)}(t)^* c_{c,\mathbf{k},dip}^{(2)}(t) + c.c. &= \frac{(A_2 \vec{e})^4 \omega \cdot \sigma^2 \pi}{8} \cdot [e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2}] \\
 &\cdot [e^{-\frac{\sigma^2}{4}\Delta\epsilon_{vc,\mathbf{k}}^2} - \frac{1}{2}(e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2} + e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2})] \\
 &\cdot \frac{\partial \Delta\epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left(\left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} + \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \right) \\
 \end{aligned} \tag{3.41}$$

$$\begin{aligned}
 c_{c,-\mathbf{k},dyn}^{(2)}(t)^* c_{c,-\mathbf{k},dip}^{(2)}(t) + c.c. &= \frac{(A_2 \vec{e})^4 \omega \cdot \sigma^2 \pi}{8} \cdot [e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2}] \\
 &\cdot [e^{-\frac{\sigma^2}{4}\Delta\epsilon_{vc,\mathbf{k}}^2} - \frac{1}{2}(e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} - 2\omega)^2} + e^{-\frac{\sigma^2}{4}(\Delta\epsilon_{vc,\mathbf{k}} + 2\omega)^2})] \\
 &\cdot \frac{\partial \Delta\epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left(\left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} + \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle \right) \\
 &= c_{c,\mathbf{k},dyn}^{(2)}(t)^* c_{c,\mathbf{k},dip}^{(2)}(t) + c.c. \\
 \end{aligned} \tag{3.42}$$

To summarize the steps, the asymmetric population distribution between \mathbf{k} and $-\mathbf{k}$ can be understood by the quantum interference (**Qui**) of different excitation paths:

3.1. TIME-DEPENDENT PERTURBATIVE ANALYSIS ON QUI

$$\begin{aligned}
|c_{c,\mathbf{k}}(t)|^2 - |c_{c,-\mathbf{k}}(t)|^2 &= 2c_{c,\mathbf{k}}^{(1)}(t)^* c_{c,\mathbf{k},dyn}^{(2)}(t) + c_{c,\mathbf{k}}^{(1)}(t)^* c_{c,\mathbf{k},dip}^{(2)}(t) - c_{c,-\mathbf{k}}^{(1)}(t)^* c_{c,-\mathbf{k},dip}^{(2)}(t) + c.c. \\
&= 2\sqrt{2}A_1 A_2^2 \vec{\epsilon} \omega \cdot \sigma^2 \pi \frac{\partial \Delta \epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left| \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \right|^2 \\
&\quad \cdot [e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - \frac{1}{2}(e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} + e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2})] \\
&\quad \cdot [e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2} - e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2}] \cos \phi \\
&\quad + \frac{\sqrt{2}A_1 A_2^2 \vec{\epsilon} \omega^2 \cdot \sigma^2 \pi}{2} \cdot (e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2}) \\
&\quad \cdot (\left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} + \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle) \\
&\quad \cdot [e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2} - e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2}] \cos \phi \\
&= \sqrt{2}A_1 A_2^2 \vec{\epsilon} \omega \cdot \sigma^2 \pi [e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2} - e^{-\frac{1}{2}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2}] \\
&\quad \cdot [\frac{\partial \Delta \epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left| \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \right|^2 (2e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2} - e^{-\frac{\sigma^2}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2}) \\
&\quad + \frac{\omega}{2} (\left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} + \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle) \\
&\quad \cdot (e^{-\frac{1}{4}(\Delta \epsilon_{vc,\mathbf{k}} - 2\omega)^2 \sigma^2} - e^{-\frac{1}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2})] \cos \phi
\end{aligned} \tag{3.43}$$

We consider the band gap between valence and conduction bands close to twice the field's frequency:

$$\Delta \epsilon_{vc,\mathbf{k}} + 2\omega \approx 0 \tag{3.44}$$

Because $\sigma \gg 0$. In a nutshell, the population imbalance becomes:

$$\begin{aligned}
|c_{c,\mathbf{k}}(t)|^2 - |c_{c,-\mathbf{k}}(t)|^2 &\approx \sqrt{2}A_1 A_2^2 \vec{\epsilon} \omega \cdot \sigma^2 \pi e^{-\frac{3}{4}(\Delta \epsilon_{vc,\mathbf{k}} + 2\omega)^2 \sigma^2} [\frac{\partial \Delta \epsilon_{vc,\mathbf{k}}}{\partial \mathbf{k}} \left| \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \right|^2 \\
&\quad + \frac{\omega}{2} (\left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle \frac{\partial \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle}{\partial \mathbf{k}} + \frac{\partial \left\langle u_{c,\mathbf{k}} \left| \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \right. \right\rangle}{\partial \mathbf{k}} \left\langle \frac{\partial u_{v,\mathbf{k}}}{\partial \mathbf{k}} \middle| u_{c,\mathbf{k}} \right\rangle)] \cos \phi
\end{aligned} \tag{3.45}$$

The utilization of two-color fields, such as ω and 2ω , has opened a potential to break the time-reversal symmetry of the systems, even when the combined field is linearly polarized. This symmetry breaking leads to a population imbalance

induced by laser irradiation, consequently resulting in dc-current injection. The population imbalance in this scheme is caused by quantum interference between two excitation paths: One is the two-photon absorption process with photons at the frequency ω , while the other is the one-photon absorption process with photons at the frequency 2ω . Hence, this protocol for dc-current injection is known as quantum interference control (**QuIC**). By manipulating the relative phase of the optical fields at frequencies ω and 2ω , **QuIC** can be applied to achieve control over one- and two-photon absorption processes, often referred to as (1 + 2 QuIC), we will further discuss in the next section.

3.2 Third-order Nonlinear Regime: 1 + 2 QuIC

From perturbation analysis in Sec 3.1, the disruption of time-reversal symmetry can be achieved through the use of linearly polarized light featuring two distinct frequencies. This implies that the injection of dc-current and the generation of a substantial population imbalance can be efficiently realized without relying on the ellipticity of light. This principle is exemplified by employing two-color linearly polarized laser fields with frequencies ω and 2ω . In this configuration, the intrinsic properties of linear polarization and the dual frequencies are sufficient to break time-reversal symmetry, facilitating the desired outcomes of dc-current injection and the establishment of a pronounced population imbalance. This approach provides a adjustable and simplified means to manipulate quantum interference and achieve specific optical responses in the system without the need for elliptically polarized light.

For practical simulation, we examine the light-induced electron dynamics in a typical two-dimensional insulator, monolayer **h-BN**, using a simple tight-binding approximation and **TDSE** introduced in Chapter 2. In the quantum dynamics simulation, we employ the following expression for the vector potential of the applied two-color fields within the interval $-\frac{\tau}{2} < t < \frac{\tau}{2}$ and zero outside this range:

$$\mathbf{A}(t) = -\mathbf{e}_p \frac{E_0}{\omega} \left[\cos(\omega t) + \frac{1}{4} \cos(2\omega t + \phi) \right] \times \cos^4\left(\frac{\pi}{\tau} t\right) \quad (3.46)$$

\mathbf{e}_p represents a unit vector along the polarization direction of the laser field, E_0 denotes the peak field strength, ω is the fundamental frequency, and τ is the duration of the laser field pulse. To illustrate dc current injection according to perturbation derivation in Section 3.1, we simulate electron dynamics using the vector potential from Eq. (3.49) with ω set to 3 eV for practical calculations. Note that the photon energy satisfies the condition ($\omega \leq E_g \leq 2\hbar\omega$) for the 1+2 QuIC process [63]. Furthermore, we set the laser polarization direction, \mathbf{e}_p , with the $\Gamma-K$ direction, the pulse duration, τ , is set to 40 fs. We introduce a relative phase ϕ between the two-color fields. The relative phase governs quantum interferences among different excitation paths induced via ω and 2ω laser fields, while the global phase is utilized to extract a dc-like response from the quantum dynamics. By manipulating the relative phase ϕ in the electric field described in Eq. (3.47):

$$\mathbf{E}(t) = -\mathbf{e}_p E_0 \left[\cos(\omega t) + \frac{1}{2} \cos(2\omega t + \phi) \right]$$

one can induce a population imbalance and, consequently, a dc-current by extrinsically breaking the time-reversal symmetry by utilizing two-color fields with frequencies ω and 2ω . Figure 3.1 (a) illustrates the electric field generated by the vector potential with a relative phase of $\phi = 0$, while Figure 3.1 (b) presents the field with a relative phase of $\phi = \pi/2$. It is evident that the field with $\phi = 0$ in Fig. 3.1 (a) breaks the time-reversal symmetry, as $\mathbf{E}(t) \neq \mathbf{E}(-t)$, while the field with $\phi = \pi/2$ in Fig. 3.1 (b) maintains the symmetry $\mathbf{E}(t) = \mathbf{E}(-t)$. Hence, the time-reversal symmetry of the Hamiltonian is broken when $\phi = 0$ and preserved when $\phi = \pi/2$. Consequently, a population imbalance and resulting dc current injection are expected when $\phi = 0$, while symmetric excitation population and the absence of net residual current are anticipated when $\phi = \pi/2$.

To comprehensively determine the persistent dc-current following laser irradiation, we conduct an in-depth analysis utilizing the photo-excited conduction population n_{ck} computed from Eq. (2.20):

$$n_{ck} = |\langle \phi_{ck} | \psi_k(t_F) \rangle|^2, \quad (3.47)$$

where t_F is a time after the laser field ends ($t_F > \tau/2$). Employing a weak enough laser field in the perturbation region with a strength of $E_0 = 2.57$ MV/cm and fix-

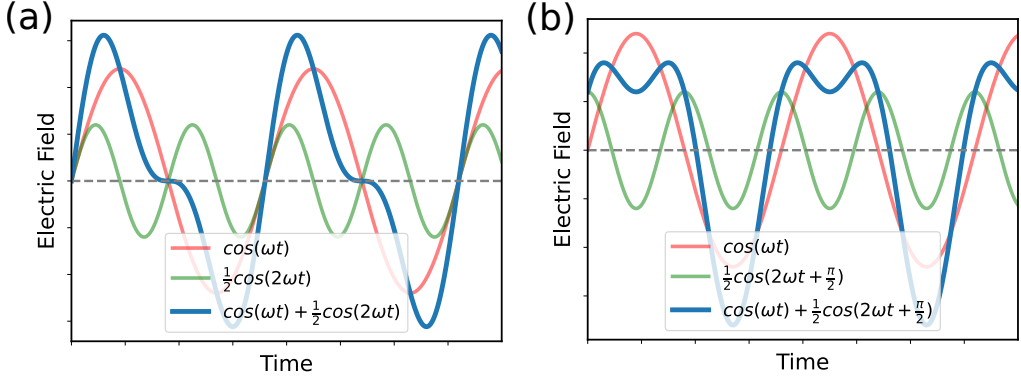


Figure 3.1: The time profiles of the electric field given by Eq. (3.47) are shown for (a) $\phi = 0$ and (b) $\phi = \pi/2$.

ing the relative phase at $\phi = 0$, the resulting conduction population is illustrated in Fig. 3.2 (a). For comparative purposes, Fig. 3.2 (b) presents the conduction population $n_{c\mathbf{k}}$ computed under the same field strength ($E_0 = 2.57$ MV/cm) but with a distinct relative phase ($\phi = 0$). In both instances, the conduction populations exhibit notable excitations centered around the K - and K' -points. This observation implies that the photo-absorption process is primarily governed by a one-photon absorption at the photon energy of $2\hbar\omega$ and a two-photon absorption at the photon energy of $\hbar\omega$. The consistency in the excitation patterns further underscores the dominance of these absorption mechanisms in the system under the specified laser conditions.

While the population distributions in Fig. 3.2(a) and (b) may initially appear similar, a closer examination reveals distinctions. In the case of the time-reversal symmetry-broken field ($\phi = 0$) illustrated in Fig. 3.2(a), the population distribution must show an imbalance between time-reversal pairs (e.g., \mathbf{k} and $-\mathbf{k}$, or K and K'). On the contrary, in the case of the time-reversal field ($\phi = \pi/2$) showed in Fig. 3.2(b), the population distribution $n_{c\mathbf{k}}$ is anticipated to lack such a population imbalance. This discrepancy arises from the absence of a persistent current under these conditions. This analysis deepens our understanding of the complicated relationship between population distributions and the underlying time-reversal symmetry characteristics, providing crucial insights into the dynamic behavior of the system.

To delineate the population imbalance across the Brillouin zone, we introduce the population imbalance distribution $\Delta n_{c\mathbf{k}}$, defined as the disparity in popula-

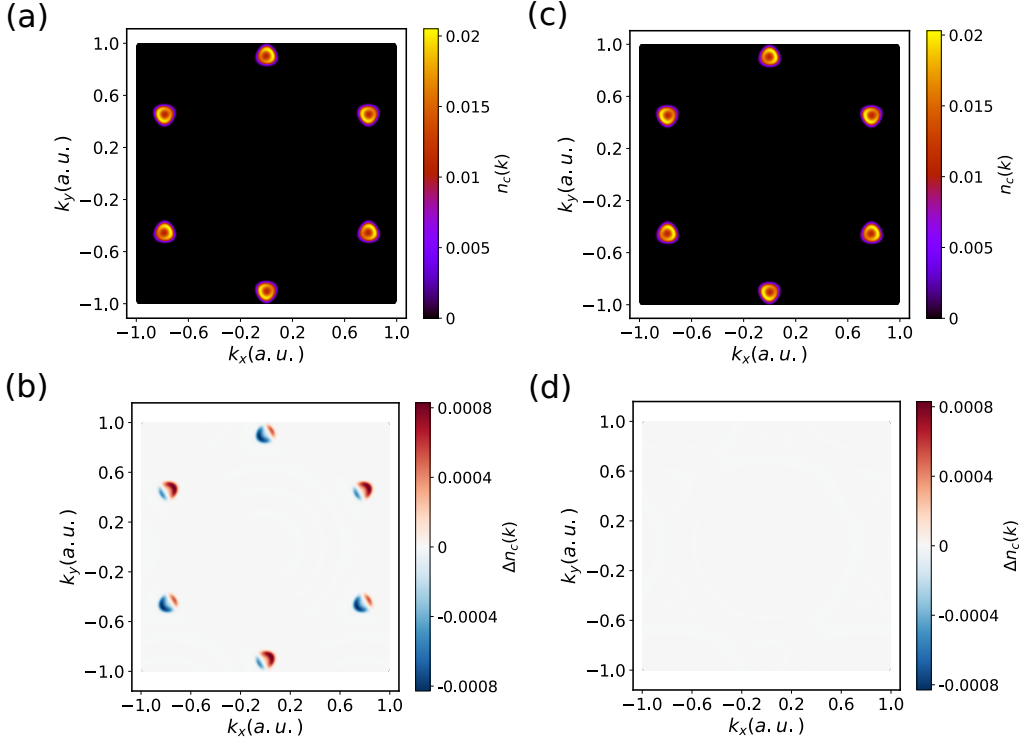


Figure 3.2: (a, b) The conduction population distribution $n_c(\mathbf{k})$ computed with (a) $\phi = 0$ and (b) $\phi = \pi/2$. (c, d) The population imbalance distribution $\Delta n_c(\mathbf{k})$ computed with (c) $\phi = 0$ and (d) $\phi = \pi/2$.

tion between the time-reversal pair k -points, expressed as:

$$\Delta n_{c\mathbf{k}} = n_{c\mathbf{k}} - n_{c,-\mathbf{k}} \quad (3.48)$$

Given the constraint $0 \leq n_{c\mathbf{k}} \leq 1$, the population imbalance distribution is bounded by $-1 \leq \Delta n_{c\mathbf{k}} \leq 1$. In situations where external fields maintain time-reversal symmetry, the populations at \mathbf{k} and $-\mathbf{k}$ are equivalent, resulting in a population imbalance distribution of zero. Conversely, in instances where time-reversal symmetry is broken, non-equivalent populations can be induced at \mathbf{k} and $-\mathbf{k}$, giving rise to a finite population imbalance distribution $\Delta n_{c\mathbf{k}}$. Figures 3.2(c) and (d) illustrate the population imbalance distribution, $\Delta n_{c\mathbf{k}}$, derived from the population distributions presented in Figs. 3.2 (a) and (b), respectively. The figures clearly illustrate that when the external field disrupts time-reversal symmetry ($\phi = 0$), a discernible finite population imbalance is induced. In contrast, when the field preserves time-reversal symmetry ($\phi = \pi/2$), the population imbalance diminishes entirely. This comprehensive analysis of the population imbalance

distribution provides a detailed insight into the complicated interplay between external field characteristics and the resulting population asymmetry within the Brillouin zone.

The temporal evolution of the corresponding electric current, denoted as $\mathbf{J}_{total}(t)$, can be computed using Eq.(2.16). This equation represents a functional dependence on the vector potential $\mathbf{A}(t)$, as illustrated in Fig.3.3. The total current encompasses multiple components and noises, often overshadowing the relatively small value of the dc-component following the laser pulse. To examine and isolate the dc-current generated by the fields, we introduce the global phase θ into the fields as described by the Eq.(3.49), as done in our prior study[69].

$$\mathbf{A}(t) = -\mathbf{e}_p \frac{E_0}{\omega} \left[\cos(\omega t + \theta) + \frac{1}{4} \cos(2\omega t + 2\theta + \phi) \right] \times \cos^4\left(\frac{\pi}{\tau}t\right) \quad (3.49)$$

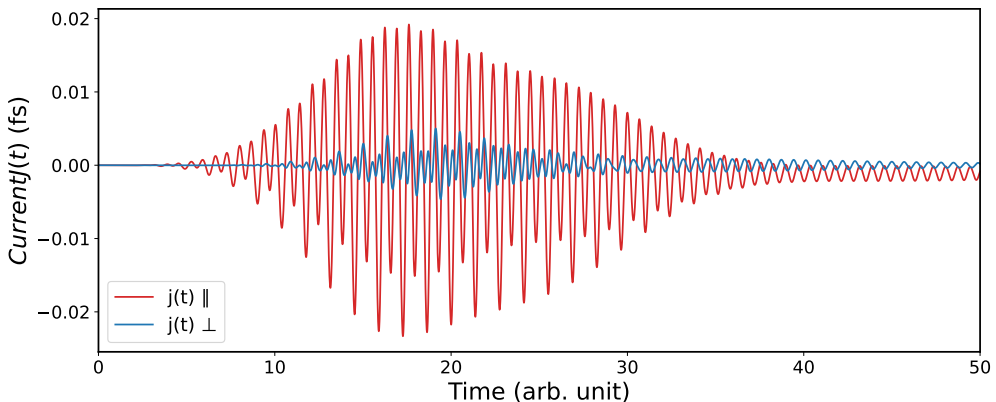


Figure 3.3

The time profiles of the current computed from Eq. (2.16) induced by the electric field given by Eq. (3.47) as shown in Fig. 3.1(a) with relative phase $\phi = 0$.

The current, expressed as a function of the global phase θ in accordance with the vector potential given by Eq.(3.49), is explicitly denoted as $\mathbf{J}(t, \theta)$. By maintaining all laser parameters constant in Eq.(3.49) except for the global phase θ , we can isolate the direct current (dc)-like component of the induced current through the following integral:

$$\mathbf{J}_{dc}(t) = \frac{1}{2\pi} \int_0^{2\pi} d\theta, \mathbf{J}(t, \theta). \quad (3.50)$$

In this formulation, the integral averages out the higher-frequency components, enabling the extraction of the clean dc-like slow-frequency component of the induced current.

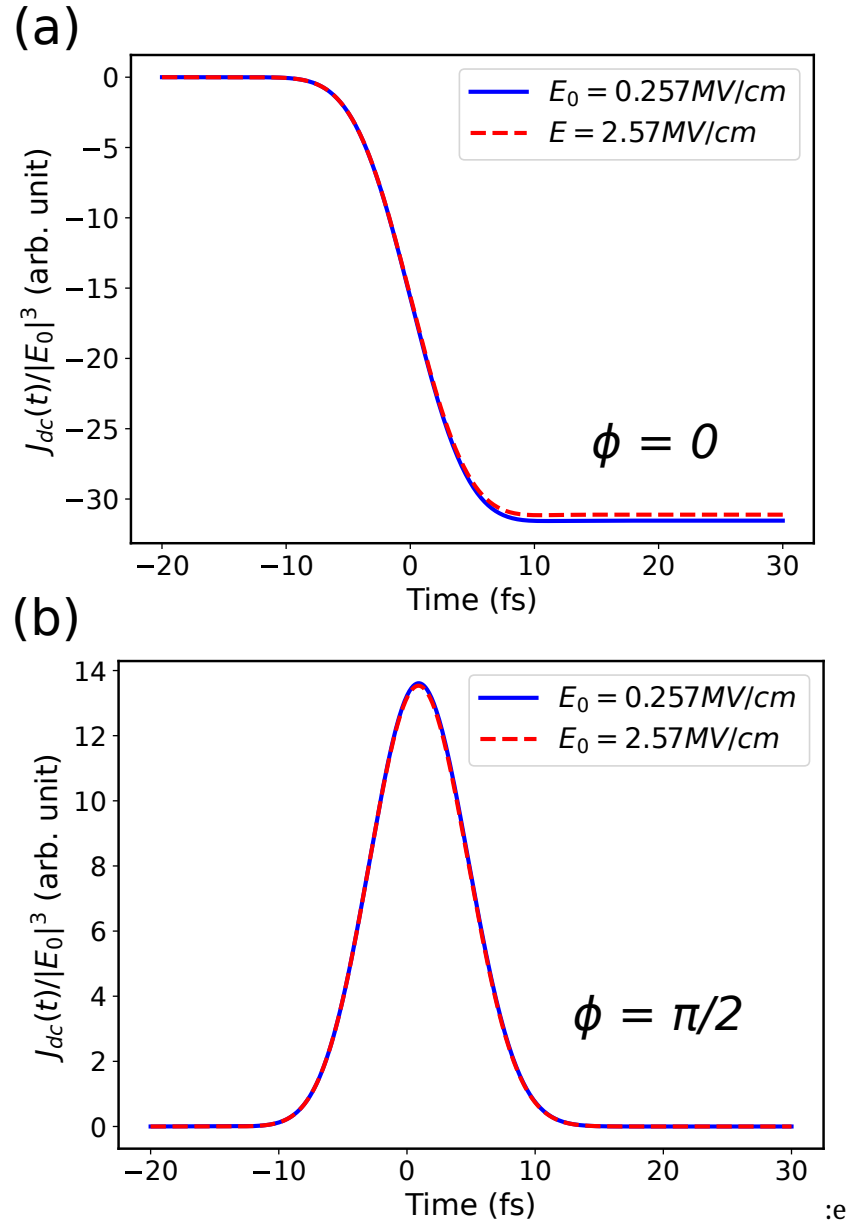


Figure 3.4: The dc components of the currents $J_{dc}(t)$ computed from Eq. (3.50) are shown as a function of time. The results using the relative phase of $\phi = 0$ are shown in panel (a), while those using $\phi = \pi/2$ are shown in (b)

In Figure 3.4(a), the calculated dc-current component of the scaled current, $J_{dc}(t)/E_0^3$, is presented for a relative phase of $\phi = 0$, including results for various field strengths, E_0 . Remarkably, the residual dc-current persists beyond the conclusion of the laser fields ($t > \tau/2$). Notably, the scaled quantity, $J_{dc}(t)/E_0^3$, maintains identical behavior across different field strengths. This consistency suggests that the dc component of the induced current can be interpreted as a third-order nonlinear optical effect. This interpretation aligns with the inherent nature of the 1+2 QuIC process, which involves interference between one- and two-photon absorption processes, classifying it as a third-order nonlinear optical phenomenon. The presented results shed light on the robust and field-independent nature of the observed third-order nonlinear optical effects in the system.

In Figure 3.4(b), the dc-current component of the scaled current, $J_{dc}(t)/E_0^3$, is illustrated with a relative phase of $\phi = \pi/2$. In stark contrast to the results with $\phi = 0$ shown in Fig.3.4(a), the currents in Fig.3.4 (b) do not show a persistent dc component after the conclusion of the laser irradiation. This outcome signifies that the applied field with a relative phase of $\phi = \pi/2$ does not disrupt time-reversal symmetry, and consequently, no population imbalance is induced, resulting in the absence of a sustained current. It is noteworthy that, even in the case of $\phi = \pi/2$, the dc-component of the current is induced solely during the laser irradiation, highlighting yet another instance of a third-order nonlinear optical process. This observation provides further insight into the nuanced interplay between field characteristics and the resulting dynamical responses in the system.

By manipulating the relative phase ϕ , one gains control over the extent of time-reversal symmetry breaking, thereby influencing the resulting population imbalance and dc- current injection [62]. For subsequent analysis, we systematically explore the persistent dc current by varying the relative phase ϕ . Figure 3.5 illustrates the dependence of the dc-current on the relative phase ϕ after laser irradiation, computed using a field with a strength of $E_0 = 2.57$ MV/cm. The amplitude of the induced dc current reaches its maximum when $\phi = 0$ and $\phi = \pi$, with opposite signs for these two phases. Moreover, the induced dc current exhibits continuous variation as the phase ϕ is manipulated, attaining zero when $\phi = \pi/2$ and $\phi = 3\pi/2$, corresponding to the points where the applied fields restore time-reversal symmetry. This straightforward phase dependence aligns with findings

from prior works[62, 64], providing further validation of the controllable nature of the induced dc current through manipulation of the relative phase.

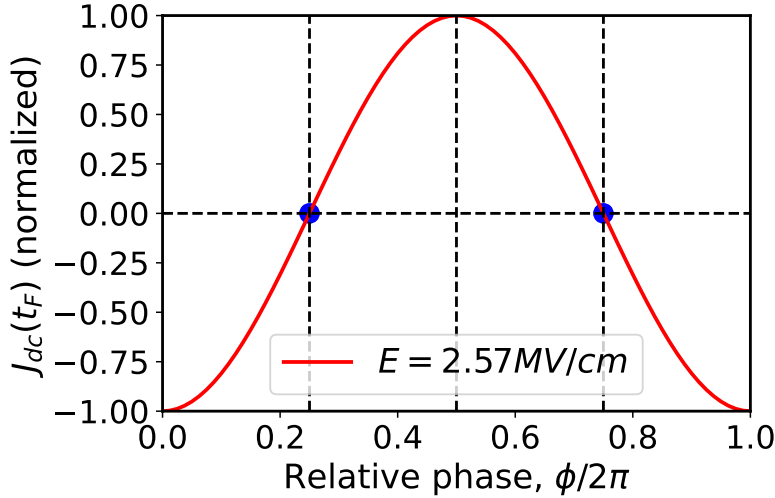


Figure 3.5: The persistent current $J_{dc}(t_f)$ as a function of the relative phase, ϕ . The results are computed by setting E_0 to 2.57 MV/cm and $\hbar\omega$ to 3 eV.

QuIC processes often exhibit resonance conditions at specific photon energies. By systematically investigating the photon-energy dependence, we can identify resonant regions where the interference effects are significantly enhanced. Investigating these dependencies helps identify the primary mechanisms. To investigate this phenomenon within our theoretical framework, we systematically evaluate the direct current (dc) after laser irradiation by varying the fundamental frequency ω in Eq.(3.49). Figure 3.6 (a) illustrates the resulting dc current following laser irradiation with a field strength of $E_0 = 1.03$ MV/m.

According to the expected behavior of the $1 + 2$ QuIC process, the dc-current decreases when the fundamental photon energy falls below half of the band gap, i.e., $\hbar\omega \leq E_g/2 = 2.95$ eV, since the fundamental photon energy $\hbar\omega$ must adhere to the condition $\hbar\omega \geq E_g/2$, where E_g signifies the band gap. In instances where the fundamental photon energy $\hbar\omega$ falls below the gap, both the $1 + 2$ QuIC process and the resultant dc-current vanish. This exploration not only clarifies the crucial role of photon energy in the indication of the $1 + 2$ QuIC process but also underscores the significance of satisfying specific conditions for its occurrence and subsequent dc-current induction. This behavior aligns with the expected characteristics of the $1 + 2$ QuIC process and provides valuable insights into the

influence of the fundamental frequency on the induced dc-current in our theoretical framework.

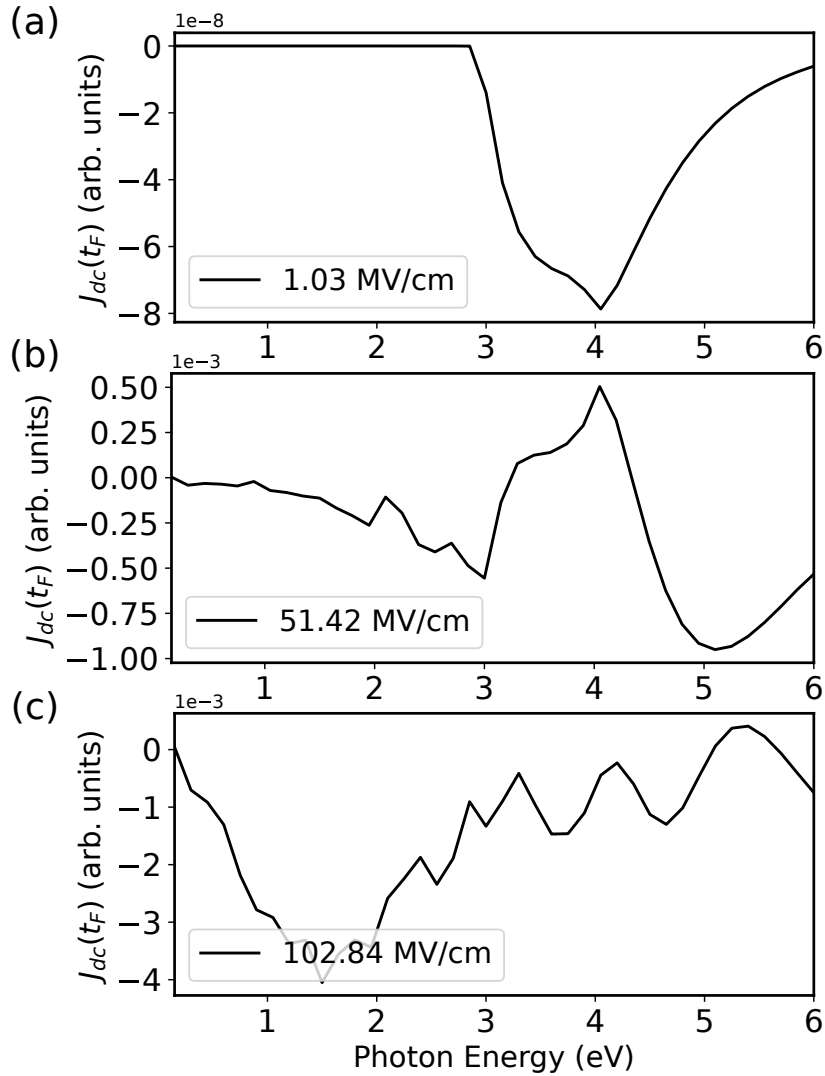


Figure 3.6: The current after the laser irradiation is shown as a function of the fundamental photon energy $\hbar\omega$. The results computed different field strengths: (a) $E_0 = 1.03 \text{ MV/cm}$, (b) 51.42 MV/cm , and (c) $E_0 = 102.84 \text{ MV/cm}$.

The investigation helps understand the complicated relationship between photon energy, field strength, and the nonlinear processes. It is essential to investigate the photon energy dependence of the dc-current after laser irradiation while varying the field strength, E_0 , to unravel the intricacies of this highly nonlinear optical phenomenon. Figures 3.6 (b) and (c) thoroughly illustrate the photon-energy dependence of the persistent current following laser irradiation, calculated for two distinct field strengths: (b) $E_0 = 51.42 \text{ MV/cm}$ and (c) $E_0 = 102.84 \text{ MV/cm}$.

In contrast to the weak field regime dominated by the $1 + 2$ QuIC, the direct current (dc) can be induced even under deeply off-resonant conditions, where the photon energy is smaller than half of the band gap ($\hbar\omega \leq E_g/2$), as evident in Figure 3.6(b). This convincing observation suggests that potent laser fields introduce additional pathways for electron excitation that extend beyond two-photon absorption. These additional processes, involving multiple photons, contribute to the creation of a population imbalance and a dc-current, even in the deeply off-resonant regime. The interaction between laser field strength and photon energy unveiled in these results provides invaluable insights into the complex dynamics governing persistent currents in strong-field regimes.

Illustrated in Figure 3.6 (c), a noteworthy observation emerges the magnitude of the direct current (dc) after laser irradiation in the deeply off-resonant regime ($\hbar\omega \leq E_g/2$) surpasses that in the $1 + 2$ QuIC regime ($\hbar\omega \geq E_g/2$) as the applied field strength reaches exceptionally large values. This interesting behavior finds its explanation in the ponderomotive energy, denoted as:

$$U_p = \frac{e^2 E_0^2}{4m\pi\omega_0^2} \quad (3.51)$$

The associated light-induced intraband transitions, both of which are more substantial for lower frequency driving[70]. Consequently, the ensuing nonlinear effects and the injection of dc current become more pronounced in the deeply off-resonant regime compared to the resonant condition. Our initial investigation focused on analyzing the electric current induced by these two-color laser fields within the weak field regime. We confirmed that the dc-component of the induced current persists even after laser irradiation when the fundamental photon energy $\hbar\omega$ exceeds the optical gap, $E_g/2$. This ballistic current phenomenon originates from a population imbalance in the Brillouin zone, arising from quantum interference between two distinct excitation paths: one involving one-photon absorption at the photon energy of $2\hbar\omega$, and the other involving a two-photon absorption path at the photon energy of $\hbar\omega$ [62–64].

This discovery sets the stage for a more in-depth exploration in the subsequent section, where we will study into the involvements of efficiently inducing dc-current through highly nonlinear optical processes in the deeply off-resonant

regime.

3.3 Deeply Off-resonant Highly-nonlinear Regime

Despite the significant interest in the nonlinear photovoltaic effect, there has been limited exploration of efficient current injection in the deeply off-resonant regime with multi-cycle light pulses, particularly using linearly polarized light. Subsequently, the scope of QuIC can be extended to involve general integer combinations, denoted as $M + N$ QuIC [66, 68]. In this extended scheme, two-color laser fields operating at frequencies ω and ω' induce M - and N -photon absorption processes, respectively. To investigate the mechanism of dc-current injection in the deeply off-resonant regime, as demonstrated in the previous section, we fix the fundamental photon energy $\hbar\omega$ in Eq. (3.49) at 1 eV. Notably, this value is much smaller than half of the band gap, $E_g/2 = 2.95$ eV, for this section.

Similarly, we start with evaluating the population imbalance induced by a strong field in the deeply off-resonant regime, we calculate the population distribution $n_{c\vec{k}}$ after irradiating the laser field with a strength of 100 MV/cm. A distinct pattern emerges in the excited carrier population distribution around the K and K' points. This pattern can be understood through the multi-photon absorption resonances of the light-induced Floquet states[71]. In Fig.3.7(a), we present the computed population distribution in the conduction band. As anticipated from the preceding discussion, the photo-carrier distribution reveals a significant population imbalance between \mathbf{k} and $-\mathbf{k}$ points. To enhance clarity in visualizing the population imbalance, we compute the population imbalance distribution $\Delta n_{c\mathbf{k}} = n_{c\mathbf{k}} - n_{c,-\mathbf{k}}$. Figure 3.7(b) displays the resulting population imbalance distribution $\Delta n_{c\mathbf{k}}$. Since $\Delta n_{c\mathbf{k}}$ is constrained by $-1 \leq \Delta n_{c\mathbf{k}} \leq 1$, the population imbalance between \mathbf{k} and $-\mathbf{k}$ is maximized when $|\Delta n_{c\mathbf{k}}| = 1$. As observed in Fig.3.7 (b), the population imbalance distribution takes significantly large values, comparable to the maximum values (± 1), across a wide range of the Brillouin zone.

We calculate the population imbalance ratio r_{im} defined as the maximum absolute value of the population imbalance distribution $\Delta n_{c\mathbf{k}}$ across the Brillouin

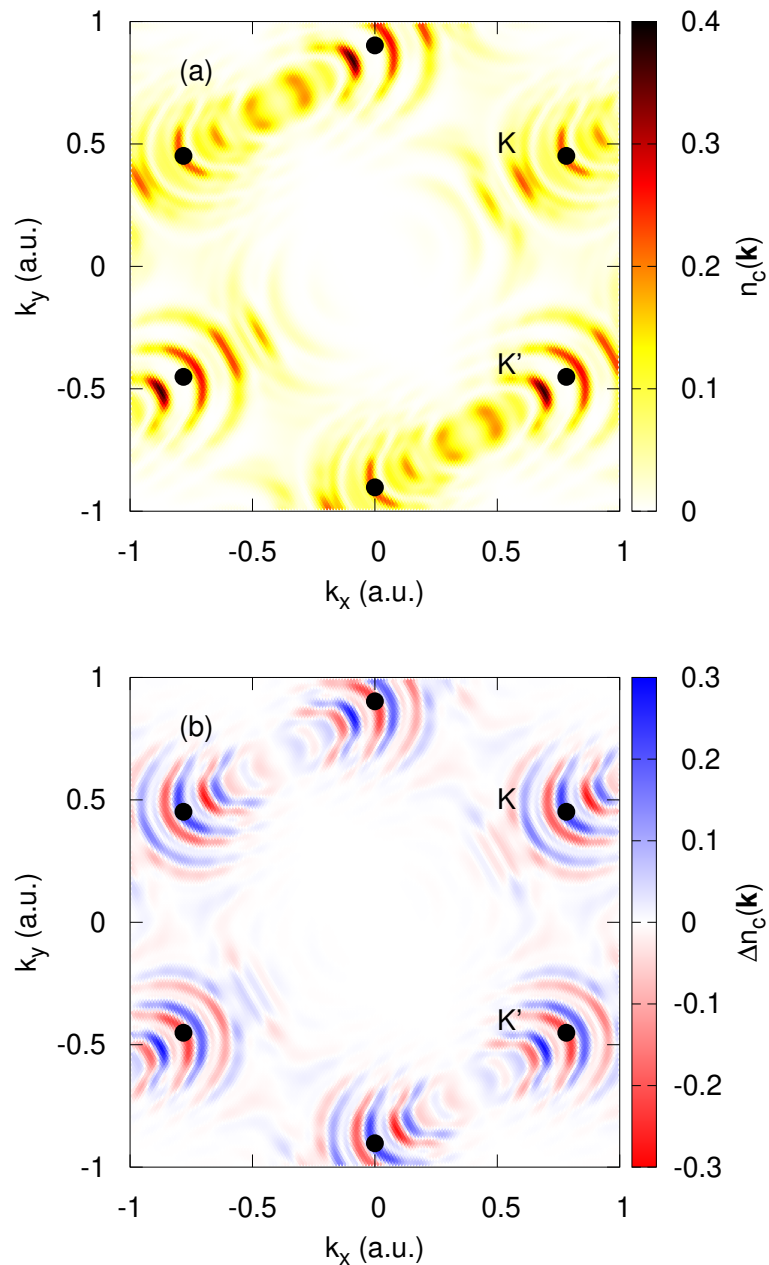


Figure 3.7: (a) The conduction population distribution $n_c(\mathbf{k})$ after the irradiation of the laser field, and (b) the population imbalance distribution $\Delta n_c(\mathbf{k})$ are shown. The results are computed by setting E_0 to 10^{10} V/m.

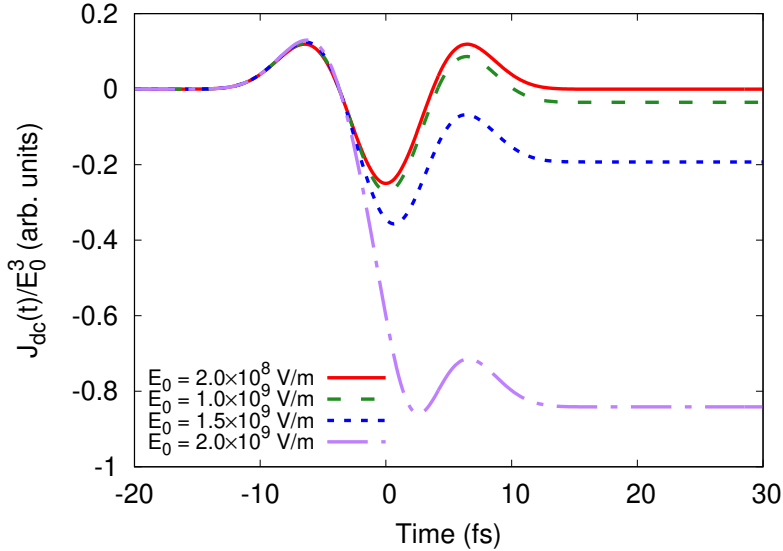


Figure 3.8: The dc components of the currents, $J_{dc}(t)$, are shown as a function of time. The results are computed with the deeply off-resonant condition, $\hbar\omega = 1.0$ eV.

zone for further qualification. Mathematically, it is expressed as:

$$r_{im} = \frac{\int_{BZ} d\mathbf{k} |\Delta n_{c\mathbf{k}}|}{\int_{BZ} d\mathbf{k} (n_{c\mathbf{k}} + n_{c,-\mathbf{k}})} = \frac{\int_{BZ} d\mathbf{k} |\Delta n_{c\mathbf{k}}|}{2 \int_{BZ} d\mathbf{k} n_{c\mathbf{k}}}. \quad (3.52)$$

The computed imbalance ratio, r_{im} , from Figs. 3.7(a) and (b) is about 0.307. Hence, more than 30% of the excited electrons contribute to the population imbalance. This implies the potential for realizing a significant population imbalance through the use of linearly polarized light alone.

In an earlier study [20], significant control over valley population was proposed using bi-circular fields with counter-rotating ω and 2ω two-color laser fields. In contrast, in this work, we demonstrate that significant valley population can be induced without relying on circular or elliptically polarized light; rather, bi-color linearly polarized light alone can break the time-reversal symmetry and cause such population control.

We commence our analysis of population imbalance by examining the light-induced current in the time domain within the deeply off-resonant regime. In Figure 3.8, we present the dc component of the current, $J_{dc}(t)/E_0^3$, computed with varying field strengths, E_0 . For this analysis, the relative phase ϕ is set to 0. Evidently, a third-order nonlinear response dominates the induced current in the

case of weak field strength. Given that the photon energy of the second harmonic is smaller than the band-gap ($2\hbar\omega < E_g$) and the QuIC process is forbidden, the third-order current returns to zero after the laser irradiation.

However, as the field strength becomes sufficiently strong, the dc-component remains finite even after laser irradiation, as illustrated in Fig. 3.8. This observation suggests that a higher-order nonlinear process contributes to the ballistic dc-current injection beyond the third-order nonlinear effect.

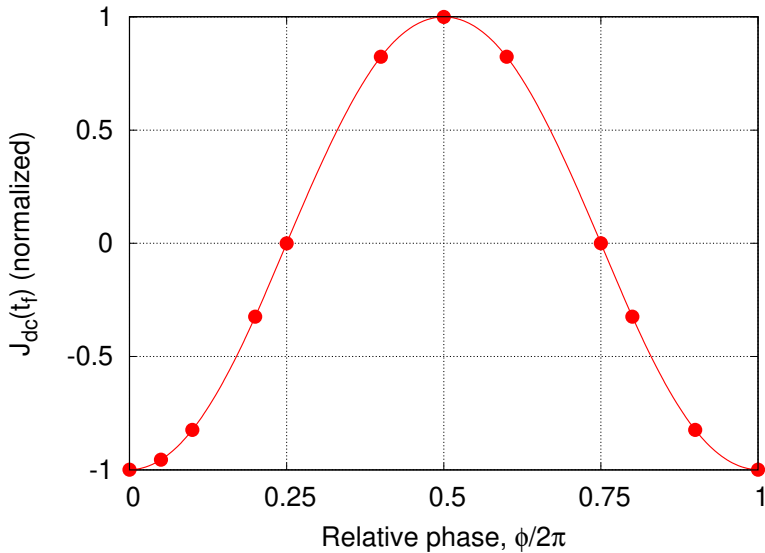


Figure 3.9: The persistent current, $J_{dc}(t_f)$, is shown as a function of the relative phase ϕ . The results are computed with the deeply off-resonant condition, $\hbar\omega = 1.0$ eV.

Next, we explore the dependence of the ballistic current induced by deeply off-resonant light on the relative phase, ϕ . Figure 3.9 illustrates the computed current as a function of the relative phase, ϕ , with calculations conducted at a field strength of $E_0 = 2 \times 10$ MV/cm. In accordance with the QuIC case shown in Fig. 3.5, the persistent current is maximized when the relative phase is $\phi = 0$ or $\phi = \pi$, and it vanishes when the applied field exhibits time-reversal symmetry ($\phi = \pi/2$ or $\phi = 3\pi/2$). Therefore, even in the deeply off-resonant regime, the direction and magnitude of the persistent current can be controlled by manipulating the relative phase ϕ between the two-color fields at frequencies ω and 2ω .

To gain a more detailed understanding of the complicated mechanism behind dc current injection in the deeply off-resonant regime, we study into an analysis

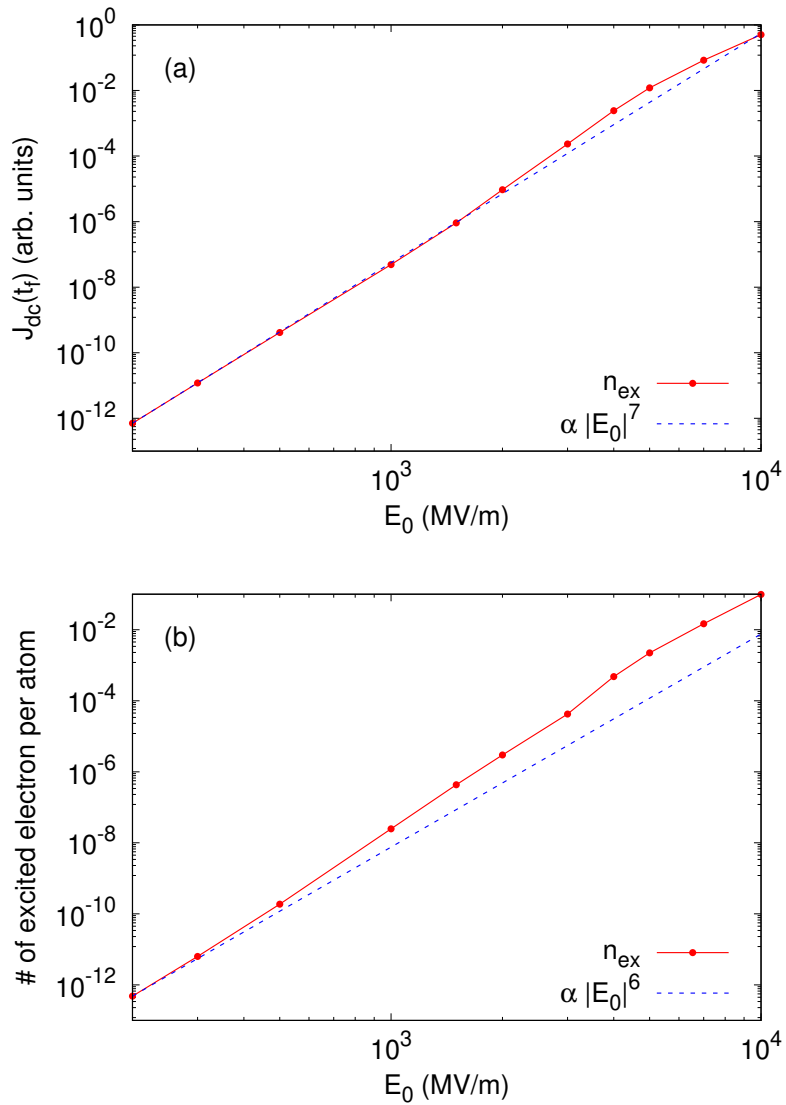


Figure 3.10: (a) The persistent current, $|J_{dc}(t_f)|$, is shown as a function of the field strength, E_0 . (b) The number of conduction population after the laser irradiation is shown as a function of the field strength E_0 .

3.3. DEEPLY OFF-RESONANT HIGHLY-NONLINEAR REGIME

of how the injected current scales with the applied field strength E_0 . As illustrated in Figure 3.10, the current amplitude after laser irradiation is plotted against the varying field strength. Notably, a reference line representing $|E_0|^7$ is included for comparison.

The compelling observation from the figure is that the induced current exhibits a clear proportionality to $|E_0|^7$ in the weak field regime. This insightful finding suggests that the seventh-order nonlinear process takes precedence in governing the dynamics of dc current injection under these conditions. This nuanced understanding provides a comprehensive insight into the complicated nonlinear optical processes that contribute to the observed dc current phenomena in the deeply off-resonant regime.

The observed scaling law of the induced dc current with the applied field strength might initially seem inconsistent with the expected behavior of a straightforward $M + N$ QuIC process. In the conventional $M + N$ QuIC situation, the M -photon absorption process is initiated by light with frequency ω , and the N -photon absorption process is generated by light with frequency 2ω , resulting in an overall $(M + N)$ -th order nonlinear process. For instance, if we consider a six-photon process for multi-photon absorption with light of frequency ω and a three-photon process for light of frequency 2ω , the anticipated simple $M + N$ QuIC process corresponds to the ninth-order nonlinear process ($M + N = 6 + 3 = 9$).

However, our experimental observations reveal a scaling that indicates seventh-order nonlinearity instead. This apparent discrepancy in the observed and expected nonlinearities of the injected dc current can be rationalized by the presence of an additional excitation channel involving a four-photon absorption process. In this situation, two photons at frequency ω and the other two photons at frequency 2ω combine to excite electrons. This additional four-photon excitation channel interferes with the three-photon absorption process at the photon energy of $2\hbar\omega$, resulting in seventh-order ($7 = 3 + 4$) nonlinear current injection.

To study deeper into the nonlinearity of the light-induced electron dynamics, we performed computations to determine the number of photo-excited carriers

after laser irradiation using the expression:

$$N_{ex} = \frac{2}{A_{BZ}} \int_{BZ} d\mathbf{k} n_{c,\mathbf{k}}, \quad (3.53)$$

where $A_{BZ} = \int_{BZ} d\mathbf{k}$ represents the area of the Brillouin zone.

Figure 3.10 (b) presents the number of excited electrons as a function of the field strength, E_0 , alongside a reference line proportional to $|E_0|^6$. In the weak field regime, the number of excited electrons exhibits proportionality to $|E_0|^6$, highlighting the dominance of the three-photon absorption process in the excitation mechanism. However, as the field strength increases, the deviation from the three-photon absorption line suggests the initiation of a nonperturbative mechanism in the excitation process.

In contrast to the $|E_0|^6$ -dependence of the number of photo-excited carriers in the weak field regime, the injected current and the corresponding population imbalance follow a $|E_0|^7$ scaling, as illustrated in Figure 3.10 (a). The difference in nonlinearities between the absolute photo-carrier population and the population imbalance implies that the population imbalance is negligible concerning the absolute photo-carrier population in the weak field regime. However, in a strong field regime, the relative significance of the population imbalance becomes substantial as it grows more rapidly than the absolute photo-carrier population. Therefore, the distinction in nonlinearities between the total photocarrier population and the population imbalance indicates the potential for large-amplitude valley carrier population control.

Expanding our analysis to the deeply off-resonant regime, where $\hbar\omega \ll E_g/2$, we observed an absence of population imbalance under weak applied field strength. However, as the field strength increased, a population imbalance in the Brillouin zone is formed, leading to the injection of the persistent dc-current after the laser irradiation. Scaling analysis of the ballistic current injection with respect to the applied field strength E_0 revealed that the population imbalance and the ballistic current result from an interference between three-photon absorption process with three photons of energy $2\hbar\omega$ and a four-photon absorption process with two photons of energy $2\hbar\omega$ and two photons of energy $\hbar\omega$. Consequently, we demonstrated that a multi-photon absorption process, incorporating photons with dif-

3.3. DEEPLY OFF-RESONANT HIGHLY-NONLINEAR REGIME

ferent energies, plays a pivotal role in addition to the multi-photon absorption process involving single-color photons.

In previous works [20–22], the formation of substantial population imbalance and valley-population control has been discussed in monolayer systems such as monolayer *h*-BN and graphene, using bi-circular laser fields with frequencies ω and 2ω . Recently, valley-population control with bi-circular fields has been extended to multi-layer and bulk systems [72] without relying on intrinsic inversion symmetry breaking and the Berry curvature at the valleys. In contrast to these works, our study demonstrates the induction of a large population imbalance and ballistic current injection without relying on the ellipticity of light. Instead, we rely on time-reversal symmetry breaking achieved through relative phase control between two-color linearly-polarized fields at frequencies ω and 2ω . Furthermore, similar to Ref. [72], the injection mechanism with bi-color linearly polarized light does not rely on intrinsic inversion symmetry breaking, indicating an efficient dc current injection and population control with the scheme using linearly-polarized light. The potential of population control and the photovoltaic effect with linearly polarized light, in addition to circularly/elliptically polarized light, unveils novel avenues for realizing ultrafast opto-electronics, marked by precise control of current and population dynamics on the femtosecond time scale.

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Chapter 4

THZ-INDUCED HHG AND NONLINEAR CHARGE TRANSPORT

Researchers have probed HHG in graphene particularly within the terahertz (THz) regime [73, 74]. THz-induced transparency of graphene, an intriguing nonlinear optical effect, has been explored [75–77], often analyzed through a thermodynamic model emphasizing the reduction of electric conductivity [49, 74]. Despite these advancements, a comprehensive understanding of the microscopic mechanisms governing these nonlinear effects remains elusive. The existing studies predominantly rely on thermodynamic models, lacking a thorough exploration of nonequilibrium quantum dynamics under dissipation. This research gap highlights the need for a deeper exploration of the underlying microscopic processes to unravel the intricacies of HHG and related nonlinear optical effects in graphene.

In this chapter, we delve into the intricate details of terahertz (THz)-induced high-order harmonic generation (HHG) and nonlinear electric transport in graphene. Our approach involves utilizing the quantum master equation with the relaxation time approximation to provide a comprehensive understanding of the underlying phenomena. To gain microscopic insights, we meticulously compare the outcomes of fully dynamic calculations with those obtained through a quasi-static approximation, wherein the electronic system is treated as a nonequilibrium steady state.

The key revelation from our investigation is that the THz-induced electron dynamics in graphene can be accurately represented by the nonequilibrium steady-state approach at each moment in time. Through a thorough population distribution analysis, we elucidate that THz-induced HHG in graphene stems from the reduction of effective conductivity, attributed to a significant displacement of electrons in the Brillouin zone.

To deepen our understanding, we draw comparisons between the nonequilibrium picture presented here and a thermodynamic perspective. This comparative analysis allows us to unravel the pivotal role of the nonequilibrium nature of electron dynamics in driving the extremely nonlinear optical and transport phenomena observed in graphene. Our study contributes valuable insights into the intricate interplay between THz fields, electron dynamics, and nonlinear behavior in graphene systems.

The comprehensive dynamical analysis derived from the quantum master equation provides a natural framework for understanding the intricate nonequilibrium features inherent in field-induced phenomena. Specifically, it allows for the exploration of phenomena characterized by symmetry breaking and delayed responses, unveiling the nuanced dynamics of the system under the influence of external fields. Ongoing theoretical investigations aim to delve deeper into these aspects, unraveling the subtleties of nonequilibrium behavior induced by light-matter interactions.

4.1 Fully Dynamical Simulations for THz Field

In this section, we first give a brief introduction of recent Experiments and Thermodynamic model on graphene. Next we delve into the microscopic intricacies governing the THz-induced high-order harmonic generation (HHG) in graphene, the electronic structure is described by the tight-binding model in Chapter 2. We initiate our investigation by conducting a detailed electron dynamics simulation using Eq. (2.21). The focus of this simulation is to analyze the high-order harmonic generation (HHG) in graphene under the influence of a linearly polarized

4.1. FULLY DYNAMICAL SIMULATIONS FOR THZ FIELD

laser pulse. To facilitate this analysis, we adopt a specific form for the applied vector potential, described by the equation:

$$A(t) = -\frac{E_0}{\omega_0} e_x \sin(\omega_0 t) \cos^4 \left(\frac{\pi}{T_{\text{full}}} t \right), \quad (4.1)$$

where the simulation is conducted in the domain $-T_{\text{full}}/2 < t < T_{\text{full}}/2$ and is zero outside this interval. Aligning with a previous experimental setup [73], we set specific parameters for the pulse: the peak field strength E_0 is chosen as 8.5 MV/m, the mean photon energy $\hbar\omega_0$ is set to 1.2407 meV, and the pulse duration T_{full} is established as 40 ps. Notably, the direction of the electric field e_x is defined along the Γ - M direction.

This detailed setup enables a thorough exploration of the electron dynamics under the influence of the specified laser pulse parameters, laying the groundwork for a comprehensive analysis of high-order harmonic generation in graphene.

Following the electron dynamics simulation governed by the field in Eq. (5.3), we proceed to compute the induced electric current, denoted as $J(t)$. To unveil the frequency content embedded within the current dynamics, we employ a Fourier transform applied to the current, yielding the high-order harmonics spectrum described by the expression:

$$I_{\text{HHG}}(\omega) \sim \omega^2 \left| \int_{-\infty}^{\infty} dt, J(t), e^{i\omega t} \right|^2. \quad (4.2)$$

Here, $I_{\text{HHG}}(\omega)$ encapsulates the contribution of high-order harmonics, and the spectrum is determined by the square of the magnitude of the Fourier transform of the induced current. This approach enables us to discern and analyze the harmonic content within the electric current, providing valuable insights into the high-order harmonic generation phenomenon induced by the specified laser pulse.

In Figure 4.1, we present the computed HHG spectra, denoted as $I_{\text{HHG}}(\omega)$, corresponding to different chemical potentials μ . Each chemical potential yields

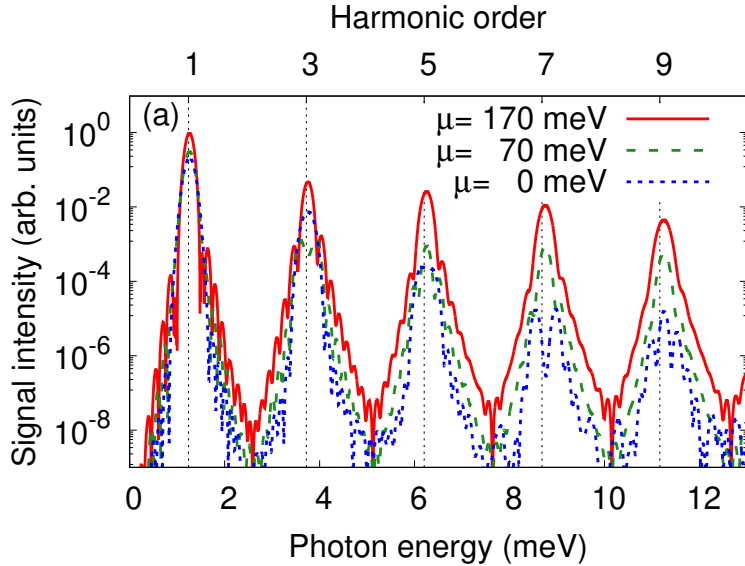


Figure 4.1: Computed harmonic spectra $I_{\text{HHG}}(\omega)$ with Eq. (4.2) for different chemical potentials, $\mu = 0, 70$ and 170 meV.

distinct harmonic peaks, and a noticeable trend emerges: the intensities of the emitted harmonics systematically increase with the rise in the chemical potential. This observation aligns with findings from a recent experiment [74], where an analogous increase in emitted harmonic intensity was noted with an elevation in gate voltage.

The results presented here echo the interpretations from prior research, such as [49], where THz-induced high-order harmonic generation in graphene was elucidated through a thermodynamic framework. In contrast, our current study aims to advance the understanding of these THz-induced nonlinear phenomena by incorporating a comprehensive microscopic perspective. Specifically, we delve into the nonequilibrium nature of electron dynamics to refine our description of light-matter interactions and provide a more nuanced interpretation of the observed trends in high-order harmonic spectra with varying chemical potentials.

Following the electron dynamics calculations under THz fields, our findings reveal that the emitted harmonics experience enhancement with an increase in chemical potential. This theoretical insight aligns with recent experimental observations, where high-order harmonic generation is similarly enhanced through the application of a gate bias voltage [74].

4.2 Quasi-static Approximation

Subsequently, we introduce a quasi-static approximation to dissect the THz-induced electron dynamics, facilitating a reexamination of the nonlinear electric transport and the field-induced transparency phenomena inherent to graphene [57]. In our analysis, we make the assumption that the variation of the THz field is sufficiently slow, allowing the electronic system to be effectively characterized by a nonequilibrium steady state at each point in time. This assumption holds true under the equilibrium established between the field-induced excitation and relaxation processes. Its accuracy is particularly pronounced when the mean frequency of the THz field is significantly smaller than the intrinsic relaxation rates, denoted as $1/T_1$ and $1/T_2$.

For practical considerations within the quasi-static approximation, we initiate our analysis by evaluating the electric current of a nonequilibrium steady state under a static electric field, represented as $E(t) = E_0 e_x$. The corresponding expression is given by:

$$J_S(E_0) = \lim_{t \rightarrow \infty} \frac{2}{(2\pi)^2} \int dk \text{Tr} [\hat{J}k(t)\rho k(t)] . \quad (4.3)$$

In this equation, the electron dynamics are computed under a static field, $A(t) = -E_0 e_x t$. Over time, the electronic system attains a nonequilibrium steady state as a result of the equilibrium between field-induced excitation and relaxation processes.

Within the quasi-static approximation, we substitute the instantaneous electric field in the induced current $J(t)$ with the steady current $J_S(E_0)$ from Eq. (4.3), resulting in the approximation:

$$J(t) \approx J_S(E(t)) . \quad (4.4)$$

To assess the validity of this approximation, we first evaluate the steady current in Eq.(4.3) for various field strengths. For practical computations, we analyze the electron dynamics under a static electric field, denoted as $E_0 = E_0 e_x$. Figure 4.2 depicts the computed current as a function of time under a static field. In this simulation, the chemical potential μ is set to 170 meV, and the field strength E_0 is established at 8.5 MV/m. The initial state at $t = 0$ corresponds to the thermal equilibrium state.

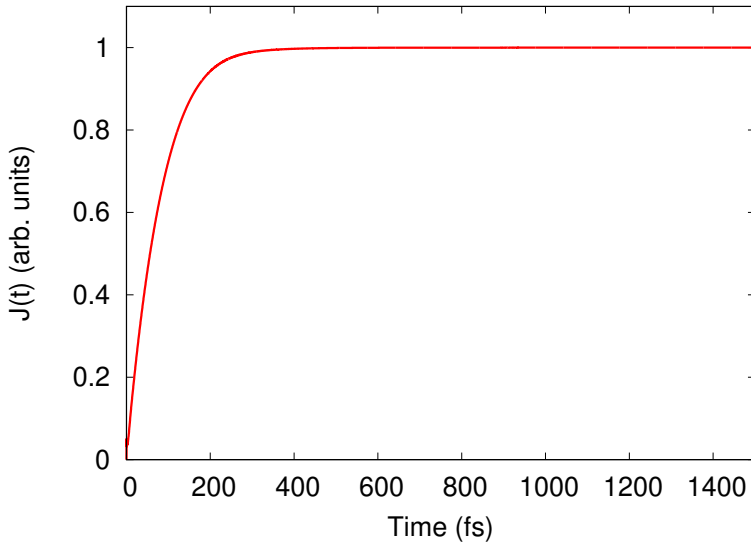


Figure 4.2: Electric current in graphene under a static electric field, $E_0 = 8.5$ MV/m.

As observed in Fig.4.2, the application of the electric field induces an electric current at $t = 0$, and it steadily approaches the steady-state value, denoted as $J_S(E_0)$. This observation confirms that the electronic system, evolving under Eq.(2.21) with a static electric field, eventually reaches a nonequilibrium steady state after a sufficiently extended period of time propagation. This validation supports the reliability of the quasi-static approximation in capturing the nonequilibrium dynamics induced by a slowly varying electric field.

In our iterative simulations, we systematically vary the field strength E_0 and assess the resulting values of the steady current. Denoting the k th set of employed field strength and evaluated current as E_k and J_k respectively, we represent the computed steady current J_k as red points in Figure 4.3 against the applied field strength E_k . To construct a continuous function $J_S(E_0)$ from the discrete data

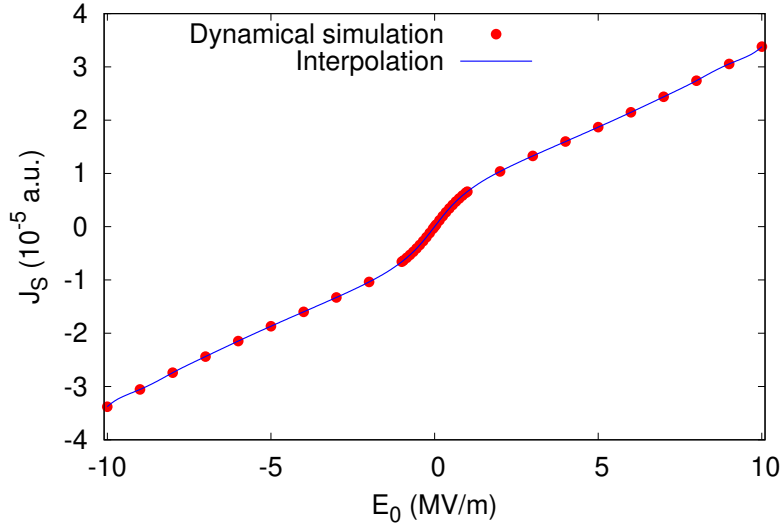


Figure 4.3: Steady current $J_S(E_0)$ as a function of field strength E_0 . The results of the fully dynamical calculation are shown as the red points, while the interpolated result is shown as the blue-solid line.

points E_k, J_k in Figure 4.3, we adopt a two-step interpolation procedure.

In the initial step of constructing the continuous function, we employ a polynomial regression with the following odd function:

$$J_{\text{polynomials}}(E_0) = \sum j = 0^4 e_x \alpha^{(2j+1)} E_0^{2j+1}, \quad (4.5)$$

where $\alpha^{(j)}$ represents optimization parameters. These parameters are fine-tuned to ensure that the polynomial function $J_{\text{polynomials}}(E_0)$ effectively reproduces the discrete points E_k, J_k in Figure 4.3.

Moving on to the second step, we aim to refine the discrepancy between the discrete points in Figure 4.3 and the polynomial function $J_{\text{polynomials}}(E_0)$. To achieve this, we define the residual error of the polynomial regression as:

$$\Delta J_k = J_k - J_{\text{polynomials}}(E_k). \quad (4.6)$$

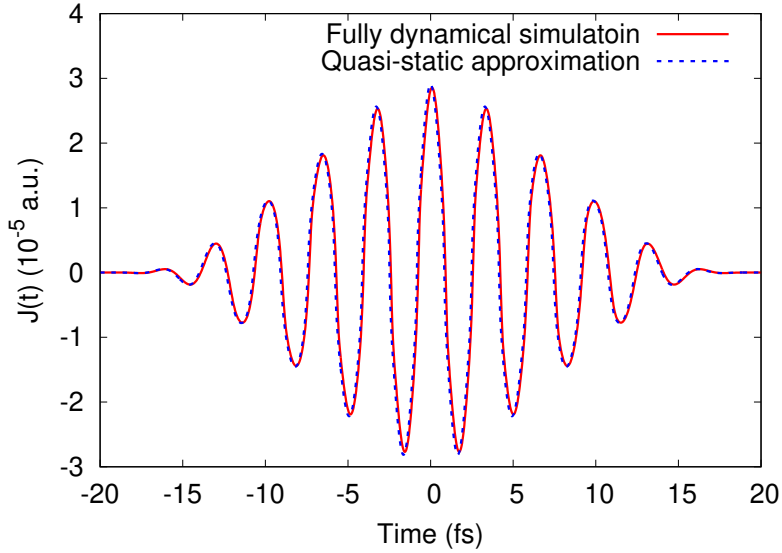


Figure 4.4: Comparison of the THz-induced current computed with the fully dynamical calculation and the quasi-static approximation.

Subsequently, we apply spline interpolation to the data points $E_k, \Delta J_k$, denoting the interpolated function as $\Delta J_{\text{spline}}(E_0)$. Finally, we approximate the continuous function, $J_S(E_0)$, as:

$$J_S(E_0) \approx J_{\text{polynomials}}(E_0) + \Delta J_{\text{spline}}(E_0). \quad (4.7)$$

Utilizing the approximated function in Eq.(4.7), we evaluate the THz-induced electric current with the quasi-static approximation, Eq.(4.4). The relationship between current and field as expressed in Eq. (4.3), we approximate the field-induced current $J(t)$ by the steady-state current, where the instantaneous electric field is denoted as $E(t)$, leading to the approximation $J(t) \approx J_S(E(t))$. Figure 5.2 showcases the computed current as a function of time with the quasi-static approximation. To facilitate comparison, the result of the fully dynamical calculation is also presented. Applying a Fourier transform to the obtained current in Figure 5.2 yields the high-order harmonic generation (HHG) spectra depicted in Figure 4.5.

To gauge the accuracy of the quasi-static approximation, we computed the high-order harmonic generation spectrum $I_{\text{HHG}}(\omega)$ using the approximated current, $J_S(E(t))$. Figure 4.5 illustrates the computed spectrum $I_{\text{HHG}}(\omega)$ with the

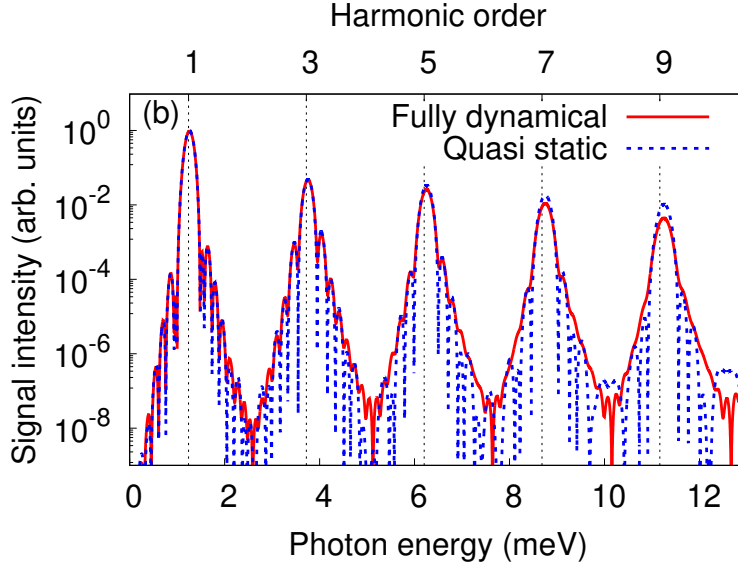


Figure 4.5: Comparison of the HHG spectra computed with the fully dynamical simulations in Sec. ?? and the quasi-static approximation Sec. ?? . Here, the chemical potential is set to $\mu = 170$ meV.

quasi-static approximation, with μ set to 170 meV. For reference, the corresponding result from the fully dynamical calculation is also presented. As evident from the figure, the quasi-static approximation faithfully reproduces the results obtained from the fully dynamical calculation. Thus, we confirm that the quasi-static approximation adeptly captures the electron dynamics in graphene under THz fields. This observation suggests that the microscopic mechanism of THz-induced high-order harmonic generation (HHG) in graphene can be effectively elucidated based on the nonequilibrium steady state, considering the delicate balance between field-induced excitation and relaxation. It is worth noting that the accuracy of the quasi-static approximation diminishes for higher-order harmonics due to the rapid dynamics that cannot be adequately captured by the quasi-static picture.

We have validated that the THz-induced high-order harmonic generation in graphene is effectively captured by the quasi-static approximation, highlighting the relevance of the nonequilibrium steady-state in portraying essential aspects of THz-induced electron dynamics. A detailed microscopic analysis has been conducted to examine the significance of the intraband current and the population distribution within the Brillouin zone in this steady-state.

4.3 Nonlinear Charge Transport in Nonequilibrium Steady-state

To provide a broader context, we juxtapose the nonequilibrium steady-state achieved within the quasi-static framework with insights garnered from a recently developed thermodynamic model [49]. This comparative analysis aims to elucidate the nonequilibrium mechanisms governing nonlinear optical and transport phenomena within graphene in the THz regime. By contrasting the quasi-static nonequilibrium picture with the thermodynamic model, we endeavor to unravel the underlying dynamics driving the intricate interplay between electron behavior and external THz fields in graphene systems. We then study the nonlinear electric conductivity in a static regime in order to develop microscopic insight into the THz-induced HHG. For this purpose, we first define the intraband component of the steady-state current in Eq. (4.3).

Next, we proceed to evaluate the effective conductivities using both the total steady current $J_S(E_0)$ and the intraband component $J_S^{\text{intra}}(E_0)$. The effective total and intraband conductivities are computed as follows:

$$\sigma(E_0) = \frac{e_x \cdot J_S(E_0)}{E_0} , \quad (4.8)$$

$$\sigma^{\text{intra}}(E_0) = \frac{e_x \cdot J_S^{\text{intra}}(E_0)}{E_0} . \quad (4.9)$$

Figure 4.6 illustrates the computed effective conductivities, $\sigma(E_0)$ and $\sigma^{\text{intra}}(E_0)$, as a function of the applied field strength E_0 for different chemical potentials μ . In this figure, the conductivities $\sigma(E_0)$ obtained from the total steady current $J_S(E_0)$ align well with those derived from the intraband current $J_S^{\text{intra}}(E_0)$ across all investigated field strengths E_0 and chemical potentials μ . This observation implies that the charge transport in graphene under static and THz fields is predominantly governed by the intraband current. The intraband current is described by the product of the band group velocity and the band population in the Brillouin zone.

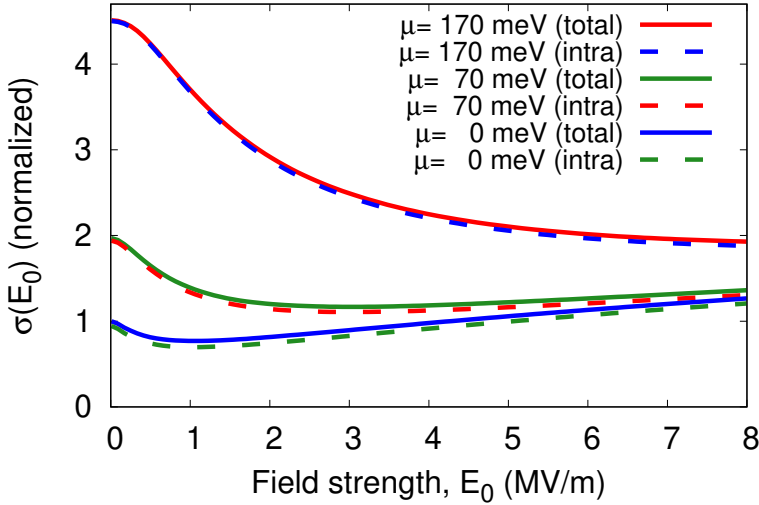


Figure 4.6: Nonlinear effective conductivities of graphene as a function of the static field strength E_0 evaluated with the total currents (solid lines) and intra-band currents (dashed lines) for different values of the chemical potential, $\mu = 0$, 70 and 170 meV.

In Fig. 4.6, the effective conductivities, $\sigma(E_0)$, exhibit an initial reduction across all investigated chemical potentials μ as the field strength increases from zero. This reduction in conductivity aligns with the field-induced transparency phenomenon observed in graphene [57], as the conductivity $\sigma(E_0)$ is directly related to photoabsorption via Joule heating, given by:

$$E_{\text{Joule}} = E_0 \cdot J_S(E_0) = \sigma(E_0) E_0^2 \quad (4.10)$$

As the field strength continues to increase, graphene with relatively small chemical potentials (e.g., $\mu = 0$ or 70 meV) exhibits an increase in conductivity, while graphene with a relatively large chemical potential (e.g., $\mu = 170$ meV) continues to show a reduction in conductivity. These trends are consistent with a previous theoretical study on nonlinear transport in graphene using the linear band approximation [57]:

$$H_k = v_F (\sigma_x k_x + \sigma_y k_y) \quad (4.11)$$

Given that the present work employs a more comprehensive electronic structure throughout the full Brillouin zone based on the tight-binding model, it serves as a

validation of the low-energy Hamiltonian approximation for the graphene band-structure used in the previous work. In the previous study [57], the decrease in effective conductivity was attributed to the dispersion of the population imbalance in the Brillouin zone, while the increase in conductivity was linked to additional carrier injection through the Zener tunneling mechanism. These interpretations naturally apply to the results obtained in the present study, further confirming the robustness and applicability of the previously proposed mechanisms.

Given the aptitude of the quasi-static approximation in accurately delineating THz-induced electron dynamics, the interpretation of THz-induced High-Order Harmonic Generation (HHG) finds its foundation in the effective conductivities, as portrayed in Fig. 4.6. Should the conductivity $\sigma(E_0)$ remain unswayed by the field strength E_0 , the induced current maintains linear proportionality to the field strength, thereby precluding the generation of harmonics. Consequently, within the quasi-static framework, the emergence of harmonics stems from the nonlinearity inherent in the current $J_S(E_0)$ and the field-strength-dependent conductivity $\sigma(E_0)$.

Examining Fig.4.6, it becomes evident that the conductivity exhibits heightened sensitivity to the field strength, particularly accentuated for larger chemical potentials. This accentuated dependence manifests as a pronounced reduction in conductivity with increasing field strength. This observation underpins the elucidation of the heightened HHG in Fig.4.1 accompanying a shift in chemical potential—attributed to the significant reduction in conductivity coincident with the amplified field strength.

In prior investigations [73, 74], the interpretation of THz-induced HHG in graphene was anchored in the reduction of conductivity, albeit within the framework of the thermodynamic model [49]. To unravel the influence of nonequilibrium dynamics in the steady state, we shall delve into the interrelation between the two models—the nonequilibrium steady-state model and the thermodynamic model—in the forthcoming section, Sec. 4.4.

Moving to the intraband current articulated in Eq.(2.29), it is crucial to recognize its composition—encompassing the product of band velocity and population. Given the inherent invariance of band velocity under the presence of

electric fields as an intrinsic material property, the crux of THz-induced current generation lies in the field-induced modulation of population. Furthermore, as expounded earlier, the THz-induced current is predominantly governed by the intraband component. For an intricate understanding of the THz-induced current at a microscopic level, we shall undertake an analysis of the population distribution within the Brillouin zone under the influence of the field. Figure 4.7 (a) delineates the equilibrium population distribution in the conduction band, denoted as $f^{FD}(\epsilon_{c,k})$, around a Dirac point (K point) in graphene, specifically at $k = \frac{2\pi}{\sqrt{3}a} \left(1, \frac{1}{\sqrt{3}}\right)$. Here, the chemical potential μ is established at 170 meV. The equilibrium population exhibits a circular symmetry around the Dirac point, reflecting the partial filling of the Dirac cone by doped electrons.

We define the field-induced alteration in conduction population within a nonequilibrium steady state as:

$$\Delta n_{c,k} = \left[n_{c,k'+eA(t)/\hbar}(t) - f^{FD}(\epsilon_{c,k'+eA(t)/\hbar}) \right]_{k'+eA(t)/\hbar=k} \quad (4.12)$$

Figures 4.7 (b-d) showcase the field-induced conduction population $\Delta n_{c,k}$ for varying field strengths: (b) 0.01 MV/m, (c) 3 MV/m, and (d) 10 MV/m.

As illustrated in Fig. 4.7(b), the field-induced population modulation emerges along the ring-shaped contour defined by the single-particle energy ϵ_{bk} and the Fermi energy $\epsilon_F = \mu|_{T_e=0}$ as $\epsilon_{bk} = \epsilon_F$, specifically where $\epsilon_{bk} = \epsilon_F$. The modulation occurs in proximity to the Fermi energy due to the mild field excitation, and the ring structure derives from the circular symmetry inherent in the Dirac cone. In the weak field regime, the augmentation and reduction in conduction population $\Delta n_{c,k}$ exhibit symmetric distribution along the field direction (x -axis). Conversely, in the strong-field regime, the distribution becomes non-symmetric, as evident in Figs. 4.7(c) and (d). Here, the red-colored region signifies an expanded range on the left side of the Dirac point, where population increase occurs, while the blue-colored region indicates a more confined area on the right side marked by population decrease. The pronounced elongation of the population increase along the field direction may be construed as a consequence of the field-induced intraband acceleration within the Brillouin zone. Simultaneously, the localized population decrease around the Dirac point, as observed in Fig. 4.7(a), can be at-

tributed to the field-induced displacement of initially localized electrons encircling the Dirac point.

In the preceding study [57], the decrease in conductivity was elucidated as the saturation of population imbalance surrounding the Dirac point. To scrutinize this interpretation, we present the conduction population distribution:

$$n_{c,k' + eA(t)/\hbar} \Big|_{k' + eA(t)/\hbar = k} \quad (4.13)$$

in lieu of the population change $\Delta n_{c,k}$ in Fig.4.7(e), with the field strength E_0 set to 10MV/m. It is noteworthy that the summation of the density in Fig.4.7(a) and the density change in Fig.4.7(d) corresponds to the density in Fig.4.7 (e).

Examining Fig.4.7(e), a discernible shift in the conduction population from the right to the left side of the Dirac cone is evident. This observation indicates that the population imbalance encircling the Dirac cone is already near its maximum saturation point, where no further population can be transferred from the right side to the left side. Consequently, the population imbalance, a pivotal determinant of the intraband current, reaches saturation in the strong-field regime, precluding any substantial increase. This saturation, in turn, leads to the saturation of the intraband current—predominant in the nonequilibrium steady state—ultimately culminating in the observed reduction in conductivity within the strong-field regime.

Our investigation reveals a substantial decrease in the effective conductivity of graphene in the strong-field regime, attributed to the saturation of population imbalance within the Brillouin zone. This reduction aligns with the observed THz-induced transparency in graphene, as reported experimentally [75–77] and theoretically investigated in prior studies [57]. Furthermore, we establish that this diminished conductivity leads to nonlinear current behavior in the strong-field regime, culminating in high-order harmonic generation in graphene. Thus, the origin of high-order harmonic generation can be comprehended through the lens of saturation of population displacement within the Brillouin zone in the context of nonequilibrium electron dynamics.

4.4 Comparison with Thermodynamic Model

Having elucidated the microscopic intricacies of THz-induced High-Order Harmonic Generation (HHG) in graphene within the framework of the nonequilibrium steady-state, our focus now shifts to investigating the distinctive role played by the nonequilibrium nature of THz-induced electron dynamics. To accomplish this, we undertake a comparative analysis with the previously formulated thermodynamic model [49]. In contrast to the present nonequilibrium model, the thermodynamic model relies on the utilization of the thermal Fermi–Dirac distribution to delineate laser-excited electronic systems. This model operates under the assumption that electrons swiftly undergo thermalization, allowing them to be effectively treated as an equilibrium state characterized by a notably high electron temperature T_e .

Within the thermodynamic model, equilibrium states find characterization through the electron temperature T_e , while in the developed nonequilibrium model, nonequilibrium steady-states are intrinsically defined by the applied field strength E_0 , devoid of any reliance on temperature considerations. To ensure a equitable comparison between the two models, it becomes imperative to establish a connection between the electron temperature T_e and the field strength E_0 . This connection is facilitated by the introduction of the field-induced excess energy for each model.

The total energy of the electronic system is formulated as:

$$E_{\text{tot}}(t) = \frac{2}{(2\pi)^2} \int dk \text{Tr} \left[H_{k+eA(t)/\hbar} \rho_k(t) \right]. \quad (4.14)$$

Subsequently, the field-induced excess energy of the nonequilibrium steady-state is defined as

$$\Delta E_{\text{excess}}^{\text{NEQ}}(E_0) = \lim_{t \rightarrow \infty} [E_{\text{tot}}(t) - E_{\text{tot}}(-t)], \quad (4.15)$$

Here, $\lim_{t \rightarrow \infty} E_{\text{tot}}(t)$ corresponds to the total energy in the nonequilibrium steady-state under the presence of the field E_0 , while $\lim_{t \rightarrow \infty} E_{\text{tot}}(-t)$ corresponds

to that of the equilibrium state without the field. Thus, the field-induced excess energy of the nonequilibrium model captures the energy difference between the nonequilibrium steady-state under an external field E_0 and the field-free equilibrium state.

In contrast, the field-induced excess energy of the thermodynamic model is defined as the energy difference between finite temperature states at T_e and 300K, the initial temperature of the present nonequilibrium model:

$$\Delta E_{\text{excess}}^{\text{TM}} = \sum_{b=v,c} \frac{2}{(2\pi)^2} \int dk \epsilon_{bk} \times [f^{\text{FD}}(\epsilon_{bk}, T_e, \mu) - f^{\text{FD}}(\epsilon_{bk}, T_e = 300 \text{ K}, \mu)]. \quad (4.16)$$

Therefore, $\Delta E_{\text{excess}}^{\text{TM}}$ is expressed as a function of the electron temperature T_e .

By employing Eq.(4.15) and Eq.(4.16), we establish a link between the applied field strength E_0 characterizing the nonequilibrium steady-state and the electron temperature T_e inherent to the thermodynamic model through the concept of excess energy. This connection enables a comparative analysis of the effective conductivity $\sigma(E_0)$ in the nonequilibrium steady-state and the linear conductivity of the thermodynamic model.

Figure 4.8 presents the conductivities of the nonequilibrium steady-state (depicted by the red-solid line) and the thermodynamic model (represented by the green-dashed line). The computations for the nonequilibrium steady-state consider a chemical potential μ set to 170 meV and an electron temperature T_e in the relaxation operator set to 300 K. In contrast, the linear conductivity of the thermodynamic model is evaluated under the influence of a weak field, ensuring the induced current aligns with a linear response. The results for the thermodynamic model involve varying the electron temperature T_e while maintaining the total population constant, as expressed by:

$$N_{\text{tot}} = \frac{2}{(2\pi)^2} \sum_{b=v,c} \int dk f^{\text{FD}}(\epsilon_{bk}, T_e, \mu), \quad (4.17)$$

to the value at $T_e = 300 \text{ K}$ and $\mu = 170 \text{ meV}$. Consequently, the chemical potential undergoes adjustments with changes in the electron temperature.

4.4. COMPARISON WITH THERMODYNAMIC MODEL

Figure 4.8 illustrates that the conductivity of the thermodynamic model (depicted by the green-dashed line) initially experiences a decline with an increase in the excess energy, followed by a notable upturn once the excess energy reaches a moderately large value. In contrast, the conductivity of the nonequilibrium steady-state (represented by the red-solid line) consistently diminishes with the escalation of the excess energy across the entire explored range. It's important to note that the conductivity of the nonequilibrium steady-state in Fig.4.8 aligns with that in Fig.4.6 when considering the converted x -axis. The fundamental contrast between the conductivities of the nonequilibrium steady-state and the thermodynamic model arises from the interband excitation influenced by temperature. In the thermodynamic model, thermal excitation propels electrons from the valence band to the conduction band, augmenting the effective carrier population and subsequently enhancing conductivity with elevated electron temperatures. In contrast, the nonequilibrium steady-state experiences a significant suppression of field-induced interband excitation due to Pauli blocking, mitigating the artificial rise in effective carrier population and the resultant boost in conductivity.

In their previous study [49], the authors delved into the microscopic intricacies of THz-induced high-order harmonic generation and field-induced transparency in graphene using the thermodynamic model. Notably, the investigation adopted a single-band approximation in which only the conduction band was considered, while the valence band remained frozen. Surprisingly, this single-band approximation yielded better agreement with experimental results than the two-band approximation, where both valence and conduction bands were considered [74]. Despite the intuitive expectation that the two-band approximation would offer greater accuracy, the results indicated that the single-band approximation provided a more accurate depiction within the thermodynamic model.

To elucidate the role of the single-band approximation in the thermodynamic model, we extended our comparison between the thermodynamic model and the nonequilibrium steady-state by incorporating the single-band approximation into our analysis. In this adaptation, we phenomenologically constrained the population in the valence band, while maintaining the use of the Fermi–Dirac

distribution for the conduction band. This modification involved transforming the Fermi–Dirac distribution as follows:

$$\tilde{f}^{\text{MFD}}(\epsilon, T_e, \mu) = f^{\text{FD}}(\epsilon, T_e, \mu)\Theta(\epsilon) + \Theta(-\epsilon), \quad (4.18)$$

where $\Theta(\epsilon)$ represents the Heaviside step function. By replacing the original Fermi–Dirac distribution (Eq.(2.26)) with the modified version (Eq.(4.18)), we conducted a comparative analysis of conductivity with the thermodynamic model. The results of the thermodynamic model incorporating the single-band approximation are depicted by the blue-dotted line in Fig. 4.8. Remarkably, this modified thermodynamic model effectively reproduces the conductivity trend observed in the nonequilibrium steady-state, showcasing a consistent monotonic decrease with an increase in excess energy.

This intriguing outcome suggests that the phenomenological freezing of the valence band in the single-band approximation curtails the artificial interband excitation in the thermodynamic model, leading to a more accurate portrayal of conductivity. In contrast, the nonequilibrium steady-state, based on a fully dynamical model, naturally captures the suppression of interband excitation, providing an accurate representation of electron dynamics in graphene under THz fields without resorting to the phenomenological freezing of the valence band.

Consequently, our findings indicate that the thermodynamic model exhibits an artificial augmentation of electric conductivity when subjected to intense THz fields, stemming from pronounced interband transitions between the valence and conduction bands. In alignment with prior work [74], we introduced a single-band approximation, freezing the valence band to mitigate this spurious interband excitation. Correspondingly, the computed conductivity in the thermodynamic framework, employing this single-band approximation, accurately reflects the anticipated decreasing trend under field irradiation. This trend aligns with experimental observations of the field-induced transparency of graphene [75–77]. In contrast, the nonequilibrium model developed in this study effectively captures the diminishing trend in conductivity under field irradiation without the need for artificially freezing the valence band. This underscores the essential role of the nonequilibrium nature of electron dynamics in describing conductiv-

4.4. COMPARISON WITH THERMODYNAMIC MODEL

ity reduction under field irradiation and preventing spurious interband excitation, as evidenced in the comparison with the thermodynamic model.

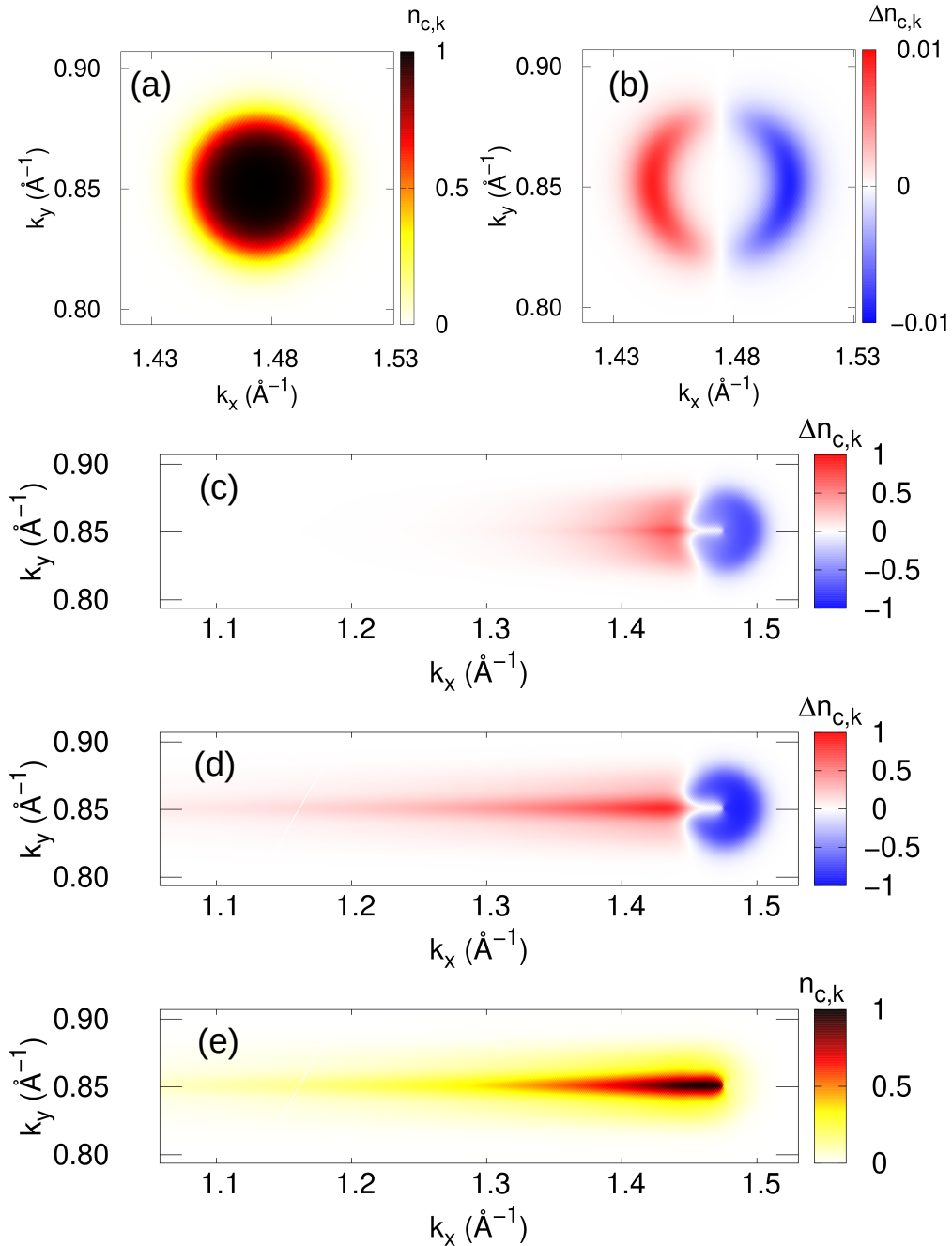


Figure 4.7: (a) The equilibrium population distribution in the conduction band $f^{\text{FD}}(\epsilon_{c,k})$. (b-d) The field induced conduction population change for different field strengths, (b) 0.01 MV/m, (c) 3 MV/m, and (d) 10 MV/m. (e) The population distribution in the conduction band in the nonequilibrium steady-state under a static field, $E_0 = 10 \text{ MV/m}$.

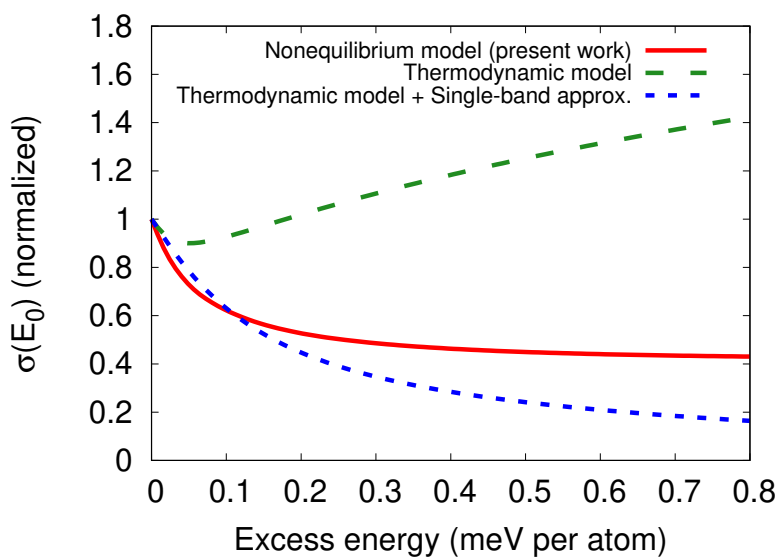


Figure 4.8: Computed effective conductivities are shown as a function of the excess energy. The results for the nonequilibrium steady-state (red-solid), the thermodynamic model (green-dashed), and the thermodynamic model plus the single-band approximation (blue-dotted) are shown.

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Chapter 5

ENHANCEMENT OF MIR-INDUCED HHG BY COHERENT COUPLING WITH THZ FIELD

Following a detailed exploration of the phenomenon of DC-current injection and the generation of population imbalance through the application of two-color linearly polarized laser fields in Chapter 2, it is also imperative to enhance the efficiency of solid-state High Harmonic Generation (HHG) for the development of innovative HHG-based light sources and spectroscopies. Recent investigations have indicated the potential enhancement of HHG from graphene using two-color laser fields, leveraging various mechanisms [21, 78, 79].

In the work by Ref.[78], the concept of two-color HHG is proposed, involving the interplay of electron-hole pair creation induced by high-frequency pump light and the subsequent acceleration of these created pairs by low-frequency light. Mrudul *et al.* delved into HHG from graphene under bicircular fields, showcasing the ability to control valley polarization[21]. Additionally, Avetissian *et al.* explored HHG from graphene under linearly polarized light and its second harmonics. They demonstrated that, when the two-color fields are perpendicularly polarized, stronger harmonics can be emitted compared to parallel polarization [79]. Recently, High Harmonic Generation (HHG) from graphene has garnered experimental attention in the mid-infrared (MIR)[80, 81] and terahertz

(THz)[82, 83] regimes, revealing distinctive ellipticity dependence and remarkable efficiency. Similar to our previous exploration of HHG from graphene in the THz regimes in Chapter 4, based on a quantum master equation, this theoretical methodology has been adeptly applied in the MIR region [84] for the elucidation of experimental results [80, 81].

Furthermore, in the MIR regime, the coupling between field-induced intra-band and interband transitions unfolds crucial channels for HHG, leading to enhanced HHG with finite ellipticity [84]. Real-time electron dynamics simulations in the THz regime have underscored the significance of considering the nonequilibrium steady-state, resulting from the delicate balance between field-driving and relaxation. This approach transcends the confines of the equilibrium thermodynamic framework and provides a more comprehensive understanding of HHG from graphene [85].

In this chapter, we delve into the prospect of leveraging a terahertz (THz) field to augment mid-infrared (MIR)-induced High Harmonic Generation (HHG) in graphene, drawing insights from our cumulative knowledge. Initially, we employ a quantum master equation to scrutinize electron dynamics under both MIR and THz fields, subsequently assessing the emitted harmonic spectra. The outcomes from fully dynamical calculations are juxtaposed against a thermodynamic model that incorporates the equilibrium Fermi–Dirac distribution. Additionally, a nonequilibrium population model is considered, accounting for a population distribution in a nonequilibrium steady-state. Through our analysis, we unveil the pivotal role played by coupling induced coherence via THz and MIR fields in enhancing MIR-induced HHG. This elucidation underscores the significance of field-induced coherence, extending beyond mere population effects.

5.1 MIR-induced HHG in Graphene under THz Fields

In our analysis of High Harmonic Generation (HHG) induced by a Mid-Infrared (MIR) laser pulse in the presence of Terahertz (THz) fields, we adopt a practical

form for the MIR pulse, expressed as follows:

$$A_{MIR}(t) = -\frac{E_{MIR}}{\omega_{MIR}} e_{MIR} \sin(\omega_{MIR} t) \cos^4\left(\frac{\pi}{T_{MIR}} t\right) \quad (5.1)$$

This pulse is defined in the domain $-T_{MIR}/2 < t < T_{MIR}/2$ and is zero outside this range. Here, E_{MIR} represents the peak strength of the MIR field, ω_{MIR} is the mean frequency, e_{MIR} is a unit vector indicating the polarization direction of light, and T_{MIR} is the pulse duration. Specifically, we set the pulse duration T_{MIR} to 0.4 ps and the mean frequency ω_{MIR} to 0.35424 eV/ \hbar for this study, while other parameters are varied in our computation.

First, we investigate HHG in graphene only with the MIR fields. For practical analysis, the direction of the angle 0° is fixed to the Γ - M axis (the x -axis in our setup), and the peak field strength of the MIR field E_{MIR} is fixed at 6.5 MV/cm. The emitted harmonics are investigated by manipulating the polarization direction of the MIR field, e_{MIR} .

To analyze the HHG efficiency, we compute the signal intensity of the emitted harmonics from Eq (4.2) at each order by integrating the power spectrum within a finite energy range as the integrated intensity of the emitted n th harmonic $I_{\text{total}}^{n\text{th}}$:

$$I_{\text{total}}^{n\text{th}} = \int_{(n-\frac{1}{2})\omega_{MIR}}^{(n+\frac{1}{2})\omega_{MIR}} d\omega I_{\text{HHG}}(\omega). \quad (5.2)$$

In our evaluation of the angular dependence of the emitted harmonic yield $I_{\text{total}}^{n\text{th}}$ without Terahertz (THz) fields, where we analyze the intrinsic symmetry of graphene, Figures 5.1 illustrate the computed angular dependence of the emitted harmonic yield $I_{\text{total}}^{n\text{th}}$ using only the Mid-Infrared (MIR) field. The emitted harmonics exhibit a six-fold symmetry, mirroring the hexagonal lattice symmetry of graphene. As depicted in Fig. 5.1, the lower-order harmonics display an almost circular angular dependence, indicative of the circular symmetry inherent in Dirac cones. In contrast, the higher-order harmonics demonstrate a more intricate six-fold symmetry in their angular dependence, owing to the deviation of the electronic structure of graphene from a simple Dirac cone when a single-particle energy is distant from the Dirac point. Similarly, we adopt the subsequent ex-

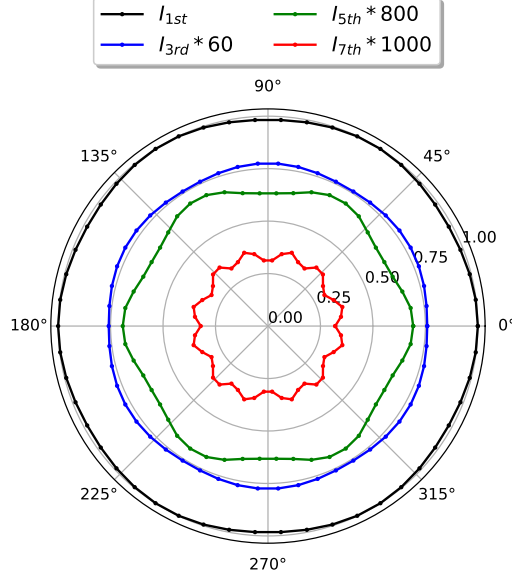


Figure 5.1: The angular dependence of the harmonic yield obtained from the electron dynamics calculations in the presence of the MIR field. The third, fifth, and seventh harmonic yields are scaled by factors of 60, 800, and 1000, respectively.

pression for the Terahertz (THz) pulse:

$$A_{THz}(t) = -\frac{E_{THz}}{\omega_{THz}} e_{THz} \sin(\omega_{THz} t) \cos^4\left(\frac{\pi}{T_{THz}} t\right) \quad (5.3)$$

within the interval $-T_{THz}/2 < t < T_{THz}/2$, and zero outside this range. Here, E_{THz} denotes the peak strength of the THz field, ω_{THz} is the mean frequency, e_{THz} represents a unit vector along the polarization direction, and T_{THz} stands for the pulse duration. In our investigation, the pulse duration T_{THz} is fixed at 40 ps, and the mean frequency ω_{THz} is set to 1.2407 meV/h. The time profile of the applied THz electric field is depicted in the inset of Fig.5.2(a).

To illuminate the intricacies of Terahertz (THz)-assisted Mid-Infrared (MIR)-induced High Harmonic Generation (HHG) in graphene, we conduct an electron dynamics calculation in the presence of both THz and MIR fields, denoted as $E_{THz}(t) + E_{MIR}(t)$. Here, we set E_{MIR} to 6.5 MV/cm and E_{THz} to 0.5 MV/cm. It is pertinent to note that experimentally available intense THz pulses can exhibit amplitudes exceeding 1 MV/cm [86]. The polarization direction of the THz field e_{THz} is aligned with the Γ - M direction (the x -direction in our setup), while the polarization direction of the MIR field e_{MIR} is considered as a tunable pa-

5.1. MIR-INDUCED HHG IN GRAPHENE UNDER THZ FIELDS

parameter. Figures 5.2(a) and (b) depict the computed current $J(t)$ induced by $E_{THz}(t) + E_{MIR}(t)$ as a function of time. The result for the parallel configuration ($e_{MIR} = e_x = e_{THz}$) is presented in Fig.5.2(a), while the result for the perpendicular configuration ($e_{MIR} = e_y \perp e_{THz}$) is shown in Fig.5.2(b). The x and y components are represented by blue and red lines, respectively. Evidently, Figs.5.2 (a) and (b) illustrate that the THz field induces a current on the picosecond time scale, whereas the MIR field induces a current on a much shorter time scale.

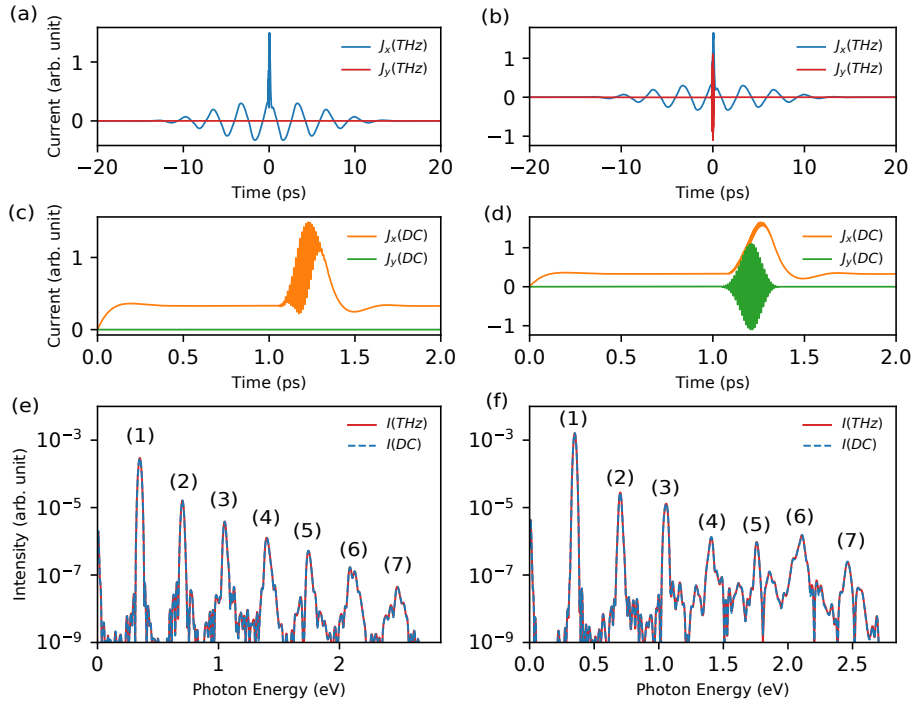


Figure 5.2: (a, b) The current $J(t)$ induced by THz and MIR fields, $E_{THz}(t) + E_{MIR}(t)$. The x -component of the current is shown as the blue line, whereas the y -component is shown as the red line. The inset is the panel (a) shows the time profile of the applied THz field. (c, d) The current $J(t)$ induced by the static and MIR fields, $E_{dc}(t) + E_{MIR}(t - \tau_{MIR})$. The x component of the current is shown as the orange line, whereas the y -component is shown as the green line. In the panels (a) and (c), the polarization of all the fields is parallel to the Γ - M direction (the x -direction in the present setup) as $e_{THz} = e_{dc} = e_{MIR} = e_x$. In the panels (b) and (d), the polarization of THz and static fields is parallel to the x -direction as $e_{THz} = e_{dc} = e_{MIR} = e_x$, while that of the MIR field is perpendicular as $e_{MIR} = e_y$. (e) The power spectra $I_{HHG}(\omega)$ computed using the current in (a) and (c). (f) The power spectra $I_{HHG}(\omega)$ computed using the current in (b) and (d).

To delve into Mid-Infrared (MIR)-induced High Harmonic Generation (HHG) amidst the presence of Terahertz (THz) fields, we scrutinize the current induced by the MIR field under the influence of the THz field. For this analysis, we delin-

eate two types of currents. Firstly, we denote the current induced by both the THz and MIR fields as $J^{THz+MIR}(t)$. Secondly, we denote the current induced solely by the THz field as $J^{THz}(t)$. Defining the current induced by the MIR field in the presence of the THz field as $J^{eff}(t) = J^{THz+MIR}(t) - J^{THz}(t)$, we subject it to Fourier transformation, followed by computation of the power spectrum of the emitted harmonics using Eq.(4.2). The solid line in Fig.5.2(e) showcases the power spectrum computed utilizing the current $J(t)$ depicted in Fig.5.2(a), wherein the polarization directions of the THz and MIR fields are parallel. Conversely, the solid line in Fig.5.2(f) exhibits the power spectrum computed employing the current $J(t)$ illustrated in Fig.5.2(b), where the polarization directions of the two fields are perpendicular. Notably, Fig.5.2(e) elucidates the generation of second and higher even-order harmonics alongside odd-order harmonics, attributed to the local breakdown of the system's inversion symmetry induced by the THz field. This phenomenon, known as electric-field-induced second-harmonic generation (EFISH) or THz-induced second-harmonic generation (TFISH), has been extensively documented[87–90]. Similarly, even-order harmonics are generated in the perpendicular configuration ($e_{MIR} \perp e_{THz}$), as depicted in Fig.5.2(f).

Incorporating the THz pulse explicitly in the electron dynamics computation extends the propagation time (42 ps in the current scenario), as illustrated in Figs.5.2(a) and (b). Consequently, performing electron dynamics calculations with the explicit inclusion of THz pulses entails a substantial computational burden. To alleviate the computational overhead associated with modeling Mid-Infrared (MIR)-induced High Harmonic Generation (HHG) in graphene under a THz field, we adopt a static field approximation based on the quasistatic approximation [85].

For practical analysis, we conduct two simulations. In the first simulation, electron dynamics are computed under a static field $E_{dc}(t) = e_{dc}E_{dc}\Theta(t)$, abruptly initiated at $t = 0$. Here, e_{dc} represents the unit vector along the polarization direction of the static field, and E_{dc} denotes the field strength. Upon the abrupt activation of the static field, the induced electron dynamics prompt a current. Following a sufficiently prolonged time of propagation, the driven system attains a steady state, and the current stabilizes over time. We designate the current induced solely by $E_{dc}(t)$ as $J^{dc}(t)$.

In the second simulation, electron dynamics are computed under both the MIR

and static fields, $E_{dc}(t) + E_{MIR}(t - \tau_{MIR})$, where the pulse center of the MIR field is shifted by τ_{MIR} . We denote the current induced by $E_{dc}(t) + E_{MIR}(t - \tau_{MIR})$ as $J^{dc+MIR}(t)$. The shift τ_{MIR} can be made sufficiently large to investigate the MIR-induced electron dynamics for a full nonequilibrium steady state realized by the static field $E_{dc}(t)$. Subsequently, the MIR-induced current can be extracted as $J^{eff}(t) = J^{dc+MIR}(t) - J^{dc}(t)$ to analyze MIR-induced HHG in the presence of the static field.

Figures 5.2(c) and (d) depict the current $J^{dc+MIR}(t)$ induced by both the static and MIR fields. The orange and green lines represent the x and y components of the current, respectively. Here, the static field aligns with the Γ - M direction (the x -direction in our setup), and its strength E_{dc} matches the peak strength of the THz field, $E_{dc} = E_{THz} = 0.5\text{MV/cm}$. In Fig.5.2(c), the MIR field aligns parallel to the static field, while in Fig.5.2(d), it aligns perpendicular to the static field. To incorporate the MIR field into the nonequilibrium steady-state under the static field, we set the time delay τ_{MIR} of the MIR field to 1 ps, exceeding the relaxation time scales of the quantum master equation, T_1 and T_2 .

To investigate HHG in the presence of the static field $E_{dc}(t)$, we extract the current $J^{eff}(t)$ induced by the MIR field in the presence of the static field by subtracting $J^{dc}(t)$ from $J^{dc+MIR}(t)$: $J^{eff}(t) = J^{dc+MIR}(t) - J^{dc}(t)$. The dashed lines in Figs 5.2(e) and (f) represent the HHG spectra computed using the current shown in Figs 5.2(c) and (d), respectively. Remarkably, the results obtained through the quasistatic approximation with a static field align perfectly with those computed by explicitly including the THz pulse. This consistency underscores the validity of the quasistatic approximation for analyzing HHG under MIR and THz fields. Additionally, we verified the consistency of the quasistatic approximation across various static field strengths (refer to Figure 5.1). Henceforth, we employ the static field within the quasistatic approximation rather than explicitly incorporating the THz pulse.

We further investigate the influence of relaxation times, T_1 and T_2 , on HHG in the presence of THz and MIR fields. Employing the methodologies outlined in Section 2.3.2, we compute the yields of third- and fifth-order harmonics under varying relaxation times. The results, depicted in Fig.5.3, exhibit consistent qualitative trends in HHG enhancement with THz field irradiation across different

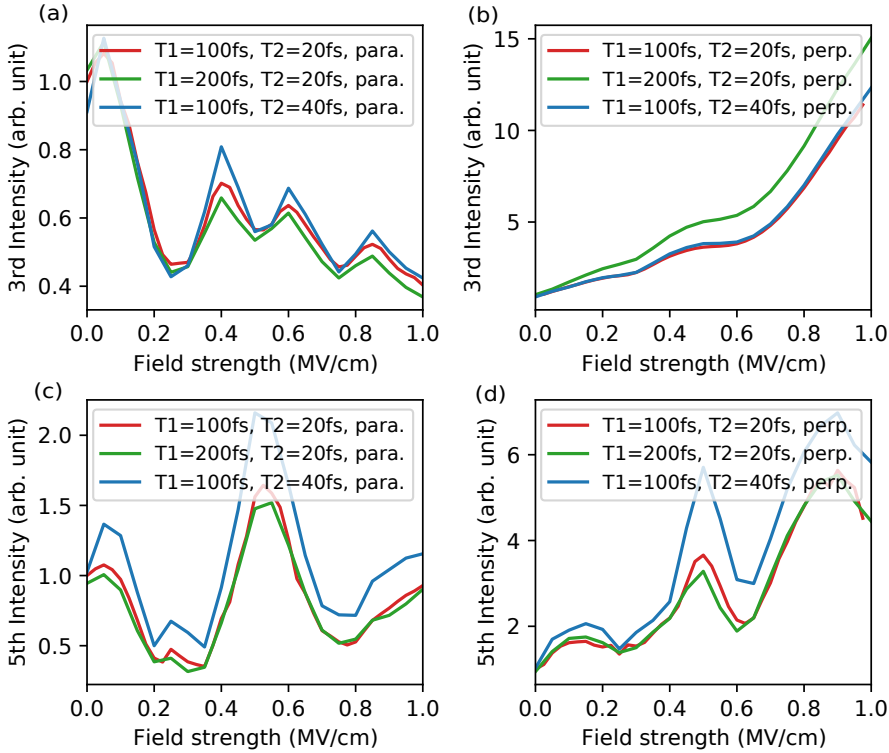


Figure 5.3: The harmonic yields are shown as a function of the static field strength E_{dc} . In each panel, the results obtained using the different relaxation times, T_1 and T_2 , are compared. The results of the third harmonics are shown in the panels (a) and (b), whereas those of the fifth harmonics are shown in the panels (c) and (d). The results obtained using the parallel configuration ($e_{MIR} = e_x = e_{THz}$) are shown in the panels (a) and (c), whereas those obtained using the perpendicular configuration ($e_{MIR} = e_y \perp e_{THz}$) are shown in the panels (b) and (d).

relaxation times. Thus, the specific choice of relaxation times does not substantially alter the enhancement phenomenon.

Relaxation times are determined by diverse scattering mechanisms, including electron-electron, electron-phonon, and electron-impurity interactions. Consequently, the actual relaxation times in practical settings depend on experimental conditions. Nonetheless, the findings presented in Fig. 5.3 suggest that HHG enhancement via THz field irradiation can manifest as a robust phenomenon across a broad spectrum of experimental scenarios.

we extend our inquiry to validate the quasi-static approximation across varying strengths of the THz field. We repeat the analyses presented in Figs. 5.2(e) and (f) while altering the THz field strength. Results obtained under a weak THz

5.1. MIR-INDUCED HHG IN GRAPHENE UNDER THZ FIELDS

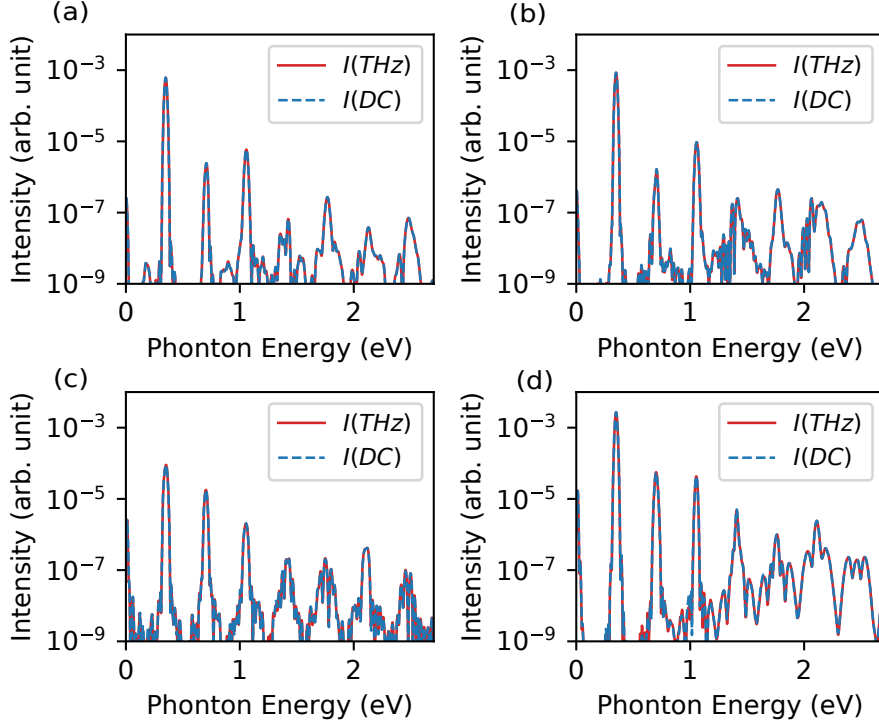


Figure 5.4: The power spectra of emitted harmonics, $I_{HHG}(\omega)$, are shown. The results obtained using a weak THz field ($E_{THz} = 0.1$ MV/cm) are shown in the panels (a) and (b), while those obtained using a strong THz field ($E_{THz} = 1.0$ MV/cm) are shown in the panels (c) and (d). The results obtained using the parallel configuration ($e_{MIR} = e_x = e_{THz}$) are shown in the panels (a) and (c), whereas those obtained using the perpendicular configuration ($e_{MIR} = e_y \perp e_{THz}$) are shown in the panels (b) and (d).

field ($E_{THz} = 0.1$ MV/cm) are depicted in Figs. 5.4(a) and (b), while those under a strong THz field ($E_{THz} = 1.0$ MV/cm) are shown in Figs. 5.4(c) and (d). As observed from the figures, the outcomes of the quasi-static approximation closely mirror those obtained from calculations with THz laser pulses across all investigated field strengths and polarization directions. Hence, we affirm the efficacy of the quasi-static approximation in accurately describing electron dynamics in graphene under THz and MIR fields, encompassing both weak and strong field regimes. This agreement between the quasistatic approximation and the explicit inclusion of the THz pulse underscores the pivotal role played by the nonequilibrium steady state under the static field in MIR-induced HHG in graphene in the presence of a THz field.

5.2 Orientational Dependence of HHG

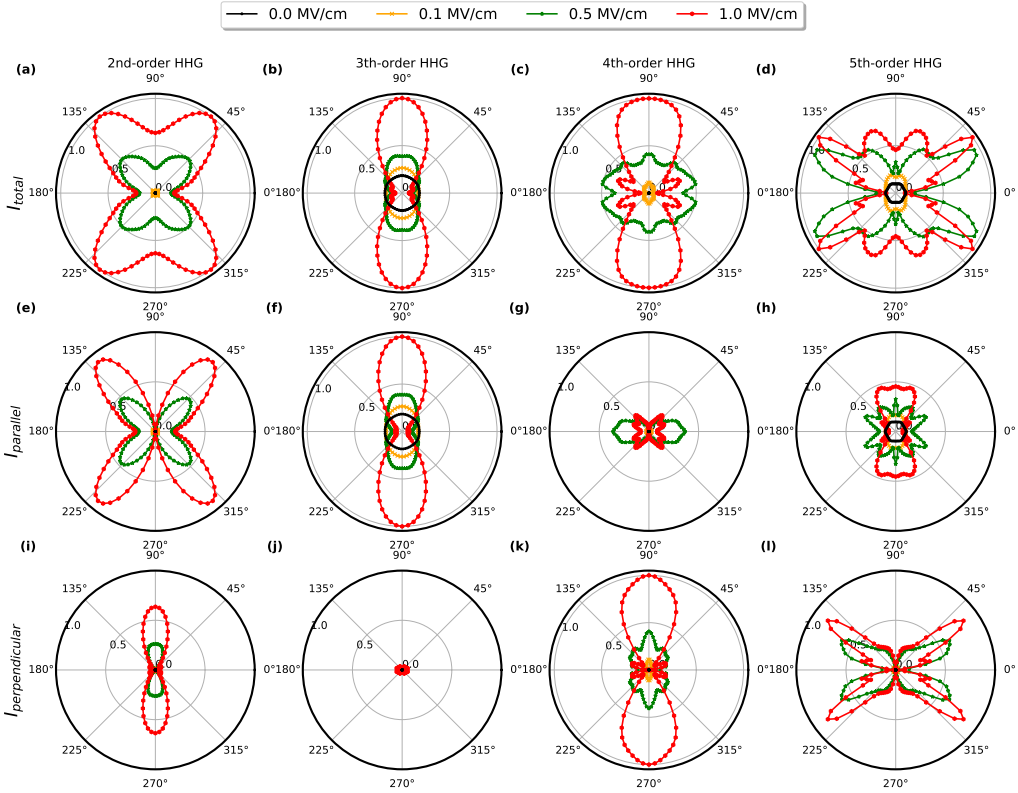


Figure 5.5: The angular dependence of the harmonic yield in the nonequilibrium steady-states under a static field along the Γ - M direction is shown for different static field strengths, E_{dc} . The angle θ denotes the relative angle between the static field and the *MIR* field. (a-d) The total intensity $I_{\text{total}}^{\text{n}^{\text{th}}}$ is shown for the second, third, fourth, and fifth harmonics. (e-h) The component of the intensity parallel to e_{MIR} is shown for each harmonic. (i-l) The component of the intensity perpendicular to e_{MIR} is shown for each harmonic. The results are normalized by the maximum total intensity $I_{\text{total}}^{\text{n}^{\text{th}}}$ for each harmonic.

We explore high-harmonic generation (HHG) in graphene within the quasi-static approximation, varying the relative angle between the static and mid-infrared (MIR) fields. For our analysis, we maintain the direction of the static field e_{dc} along the Γ - M axis (the x -axis in our setup) and set the peak field strength of the MIR field E_{MIR} to 6.5 MV/cm. We investigate the emitted harmonics by manipulating the polarization direction of the MIR field, e_{MIR} , and adjusting the strength of the static field, E_{dc} .

Figures 5.5(a-d) illustrate the angular dependence of the emitted harmonic yield $I^{\text{n}^{\text{th}}}$ for various harmonic orders. Here, θ represents the relative angle

between the MIR and static fields. In Fig.5.5(a), absence of a static field results in no second harmonic generation, given the intrinsic inversion symmetry of graphene. However, with the introduction of a static field, second harmonics emerge due to the breakdown of this symmetry. Notably, for a static field strength of 0.5MV/cm, the emitted second-harmonic intensity peaks at approximately 45° relative angle.

In Fig.5.5(b), the third-harmonic yield appears nearly isotropic (depicted by the black line) in the absence of a static field, reflecting the rotational symmetry of the Dirac cone (refer also to Figure 5.1). Conversely, under the influence of a strong static field ($E_{dc} = 1.0\text{MV/cm}$), the third-harmonic intensity exhibits significant angular dependence: it is notably enhanced when the static and MIR fields are perpendicular to each other, whereas it is suppressed for parallel field orientations. This enhancement for the perpendicular configuration can be attributed to the coupling between the intraband transition induced by the static field and the interband transition induced by the MIR field, as previously suggested[84].

The angular dependence of higher-order harmonics becomes more intricate under a static field, as depicted in Figs.5.5(c) and (d). Notably, the fifth-order harmonic emission shows significant enhancement in the presence of either static or THz fields (Fig.5.5(d)). For instance, applying a static field of 0.5MV/cm boosts the fifth-order harmonic intensity by more than tenfold compared to that induced solely by the MIR field (indicated by the green line in Fig.5.5(d)). This enhancement ratio surpasses that observed for the third-order harmonic, suggesting a greater potential for field-induced enhancement in higher-order harmonics. Indeed, the seventh-order harmonic exhibits a 25-fold enhancement with a static field strength of 0.5MV/cm (refer to Figure 5.1).

To further elucidate the angular dependence of HHG in graphene, we decompose the harmonic intensity $I_{\text{HHG}}(\omega)$ into parallel and perpendicular components with respect to the polarization of the driving MIR field. The parallel component of the HHG intensity is defined as

$$I_{\text{HHG}}^{\text{para}}(\omega) \sim \omega^2 \left| \int_{-\infty}^{\infty} dt e_{MIR} \cdot J(t) e^{i\omega t} \right|^2, \quad (5.4)$$

where e_{MIR} is the unit vector along the polarization direction of the MIR field. Likewise, the perpendicular component is defined as

$$I_{\text{HHG}}^{\text{perp}}(\omega) \sim \omega^2 \left| \int_{-\infty}^{\infty} dt \bar{e}_{MIR} \cdot J(t) e^{i\omega t} \right|^2, \quad (5.5)$$

where \bar{e}_{MIR} is a unit vector perpendicular to e_{MIR} , i.e., $\bar{e}_{MIR} \cdot e_{MIR} = 0$. The total intensity I_{HHG} in Eq. (4.2) is reproduced by the sum of $I_{\text{HHG}}^{\text{para}}(\omega)$ and $I_{\text{HHG}}^{\text{perp}}(\omega)$ as $I_{\text{HHG}}(\omega) = I_{\text{HHG}}^{\text{para}}(\omega) + I_{\text{HHG}}^{\text{perp}}(\omega)$.

Equations (5.4) and (5.5) are utilized to dissect the emitted harmonic intensity into parallel and perpendicular components. Figures 5.5 (e–h) and (i–l) elucidate the angular dependence of the parallel and perpendicular components of the harmonic intensity, respectively, for different orders.

In Figs.5.5(a), (e), and (i), the parallel component of the second harmonic under the static field peaks around 45° , constituting the dominant contribution to the total second-harmonic intensity at this orientation. Conversely, the maximum perpendicular component is consistently achieved when the MIR and static fields are orthogonal to each other. In Figs.5.5(b), (f), and (j), the third harmonic is predominantly governed by its parallel component across all angles and static field strengths examined. Notably, for both second- and third-harmonic generation, the parallel components prevail when the emitted harmonic intensity is maximized.

Qualitative distinctions emerge between the lower-order harmonics (second and third) and the higher-order ones (fourth and fifth). In Fig.5.5(c), the fourth harmonic yield peaks at an angle θ of 90° under the strongest applied static field, $E_{dc} = 1.0\text{MV/cm}$. A comparison of Figs.5.5(g) and (k) reveals the predominance of the perpendicular component in the emitted harmonic intensity under these conditions. Conversely, as depicted in Figs.5.5(d), (h), and (l), the emitted fifth harmonic at the most efficient angle is primarily governed by the perpendicular component, despite the dominance of the parallel component at all angles in the absence of a static field. Hence, the emission pathways associated with the perpendicular components are anticipated to play a crucial role in enhancing MIR-induced HHG by a THz field. This trend persists for higher-order harmonics as well (see Figure5.1).

5.3 Comparison of Nonequilibrium Steady State and Thermodynamic Model

In this investigation, we delve into the role of nonequilibrium steady states in high harmonic generation (HHG) by juxtaposing the outcomes of the quasistatic approximation with those derived from the thermodynamic model [49], a framework previously utilized in studying HHG in graphene under THz fields [82, 83]. The quasistatic approximation replaces the THz pulse with a static field to describe the electronic system's behavior under THz irradiation, whereas the thermodynamic model approximates the system's response to a THz pulse as a high-temperature thermal state [49]. The discrepancy between these models elucidates the influence of nonequilibrium distributions, delineating their significance in HHG.

The quasistatic approximation is characterized by the static field strength, E_{dc} , while the thermodynamic model relies on the electron temperature T_e . To enable a direct comparison between these models, we introduce the concept of excess energy [85] as a common metric of excitation intensity. The excess energy under the quasistatic approximation, denoted as $\Delta E^{\text{non-eq excess}}(E_{dc})$, quantifies the change in total energy due to the static field $E_{dc}(t)$. Conversely, the excess energy within the thermodynamic model, $\Delta F_{\text{excess}}^{\text{thermo}}(T_e)$, measures the energy change arising from an increase in electron temperature from room temperature ($T_e = 300$ K). This common excess energy facilitates an objective and quantitative comparison between the two models [85].

Figure 5.6 illustrates the comparison between the results obtained from the quasistatic approximation and the thermodynamic model. Setting the MIR field strength to 6.5 MV/cm and its polarization direction to the Γ - M direction (the x -axis), we observe distinct behaviors in odd-order harmonics between the two models. Figure 5.6(a) showcases the substantial enhancement and suppression of the MIR-induced third harmonic under the quasistatic approximation for parallel and perpendicular configurations, respectively. In contrast, the thermodynamic model yields nearly constant results. Figures 5.6 (b) and (c) further reveal significant enhancements in the fifth- and seventh-harmonic yields under a static

field within the quasistatic approximation, while the thermodynamic model exhibits marginal variations in harmonic yields with increasing electron temperature. Consequently, the observed HHG enhancement cannot be solely attributed to the simple heating of electronic systems within the thermodynamic model, underscoring the crucial role of non-equilibrium electronic dynamics induced by the field. The minimal alterations in harmonic yields within the thermodynamic model relative to those predicted by the nonequilibrium steady-state model suggest that modifications in the population distribution around the Fermi level exert negligible influence on HHG spectra.

5.4 Contribution of Nonequilibrium Population

In our exploration of the coherent coupling between the MIR and THz fields, beyond the population contribution induced by the THz field, we introduce a nonequilibrium population distribution model as an extension of the thermodynamic model.

Within the thermodynamic model, the THz field's contribution is represented by adjusting the population distribution via an increased electronic temperature in the reference Fermi–Dirac distribution. This model captures only the population contribution, corresponding to the diagonal elements of the density matrix, based on the thermal distribution.

To delve into the coherent coupling contribution, we extend the thermodynamic model by substituting the reference Fermi–Dirac distribution in the relaxation operator (Eq. (2.25)) with the population distribution of the nonequilibrium steady state under a static field. This extension incorporates the population contribution, signified by diagonal elements of the density matrix, while omitting THz-induced coherence, represented by the off-diagonal elements of the density matrix.

By comparing the nonequilibrium population model with the fully dynamical model, which encompasses both population and coherence effects, we can

discern the role of coherent coupling between the THz and MIR fields. This comparative analysis helps elucidate the distinct contributions of population and coherence effects to HHG, providing valuable insights into the underlying mechanisms governing this phenomenon. To formulate the nonequilibrium population model, we first analyze the population distribution in the nonequilibrium steady state under a static field. The population distribution in the Brillouin zone can be expressed as

$$\begin{aligned} n_{bk}(t) &= \int dk' \delta(k - K'(t)) \text{Tr} \left[|u_{bk'}^H(t)\rangle\langle u_{bk'}^H(t)| \rho_{k'}(t) \right] \\ &= \langle u_{b,k-eA(t)}^H(t) | \rho_{k-eA(t)}(t) | u_{b,k-eA(t)}^H(t) \rangle, \end{aligned} \quad (5.6)$$

where $K'(t)$ is the accelerated wavevector in accordance with the acceleration theorem, $K'(t) = k' + eA(t)$. The population distribution in the nonequilibrium steady state can be evaluated in the long-time propagation limit under a static field $A(t) = -E_{dc}t$,

$$n_{bk}^{\text{neq-steady}} = \lim_{t \rightarrow \infty} n_{bk}(t). \quad (5.7)$$

In Fig.5.7(a), we illustrate the population distribution in the conduction band for the nonequilibrium steady-state under a static field with a strength of $E_{dc} = 0.5 \text{ MV/cm}$. The static field is oriented along the Γ - M direction (x -axis), and the blue circle marks the Dirac point (K point).

In this depiction, the region to the left of the Dirac point is predominantly occupied by the field-induced population in the nonequilibrium steady-state, while the region to the right of the Dirac point appears nearly empty. This asymmetry disrupts the inversion symmetry of the system. We utilize this nonequilibrium population distribution as the reference distribution of the relaxation operator in Eq. (2.25) instead of the Fermi–Dirac distribution to establish the nonequilibrium population model.

In Fig.5.7(b), we present the angular dependence of the second-harmonic yield under a static field with a strength of $E_{dc} = 0.5 \text{ MV/cm}$. The corresponding angular dependences of the third, fourth, and fifth harmonics are depicted in Figs.5.7(c–e), respectively. Each panel displays results obtained using the nonequilibrium

population model as the blue solid line, juxtaposed with results derived from the quasi-static approximation, depicted as the green solid line, which matches the result shown in Fig.5.5.

Figs 5.7 (b) and (d) highlight that even-order harmonics computed with the nonequilibrium population model are notably weaker compared to those calculated using the fully dynamical approach based on the quasi-static approximation. This discrepancy indicates that under the charge-neutral condition ($\mu = 0$) examined here, the resonant effects of the MIR field at two- and four-photon resonances are significantly distant from the Fermi level. Consequently, modifications to the population near the Fermi surface yield minor contributions to even-order harmonic generation. In contrast, the fully dynamical calculation reveals that the THz field can coherently couple with the MIR field via off-diagonal elements of the density matrix, enabling coherent coupling not only around the Fermi level but also across the Brillouin zone wherever dipole transitions are permitted. Thus, the coherent coupling component may strengthen even-order harmonic generation by enhancing contributions from resonant quantum pathways.

Fig.5.7(c) demonstrates that the third-harmonic yield computed using the fully dynamical model is 1.57 times stronger than that obtained using the nonequilibrium population model when the fields are perpendicular. This result suggests that the THz field enhances third-harmonic generation for the perpendicular configuration, with both coherent coupling and incoherent population playing crucial roles in this THz-assisted enhancement mechanism. Conversely, when the fields are parallel, the third-harmonic yield calculated using the fully dynamical approach is 0.57 times weaker than that computed using the nonequilibrium population model. This finding indicates that contributions from coherent coupling and incoherent population counteract each other, diminishing the overall signal. Thus, both coherent coupling and incoherent population influence third-harmonic generation depending on the relative angle θ between the THz and MIR fields.

In Fig.5.7(e), we observe that the fifth-order harmonic yield computed using the fully dynamical model is significantly higher than that obtained using the

5.4. CONTRIBUTION OF NONEQUILIBRIUM POPULATION

nonequilibrium population model, except when the MIR and THz fields are parallel. This observation suggests that coherent coupling predominantly contributes to the enhancement of fifth-harmonic generation for most angles, although both coherent coupling and incoherent population effects are relevant when the fields are parallel. These consistent results are similarly observed for higher-order harmonics (see Appendix ??).

In summary, when only a MIR field is applied to graphene, the induced HHG is attributed to interference between multiple excitation pathways involving nonlinear coupling between MIR-induced intraband and interband transitions. Conversely, the substantial enhancement of HHG observed in the presence of THz fields indicates the activation of an additional nonlinear coupling mechanism. This mechanism arises from coherent coupling between MIR- and THz-induced transitions, suggesting its predominance over other processes in contributing to overall harmonic yield.

The comparison between the results obtained using the fully dynamical calculation and the nonequilibrium population distribution model has provided valuable insights into the roles of coherent coupling between the MIR and THz fields. The dominance of coherent coupling in generating THz-induced even-order harmonics and enhancing high-order harmonics suggests its crucial role in driving nonlinear optical processes in solids. Conversely, the enhancement of third harmonics under a THz field is influenced by both coherent coupling and the nonequilibrium population. Furthermore, coherent coupling appears to predominantly contribute to the enhancement of higher-order harmonics.

Importantly, these enhancement mechanisms are not confined to specific laser parameters but can be realized under more general conditions. Therefore, effective control of both coherent coupling and population dynamics becomes essential for boosting HHG from solids using multicolor laser fields.

Furthermore, the significance of coherent coupling extends across various orders of harmonic generation, as evidenced by the coherent coupling mechanism's influence on low-order harmonic phenomena (see Fig. 5.7). This underscores the indispensability of field-induced coherence in nonlinear optical effects more

broadly. Consequently, these findings hint at the potential for efficiently controlling electron and spin dynamics through coherent coupling, leveraging multi-color lasers. Such capabilities would transcend mere frequency conversion of light, opening avenues for the advancement of ultrafast optoelectronics and optospintronics.

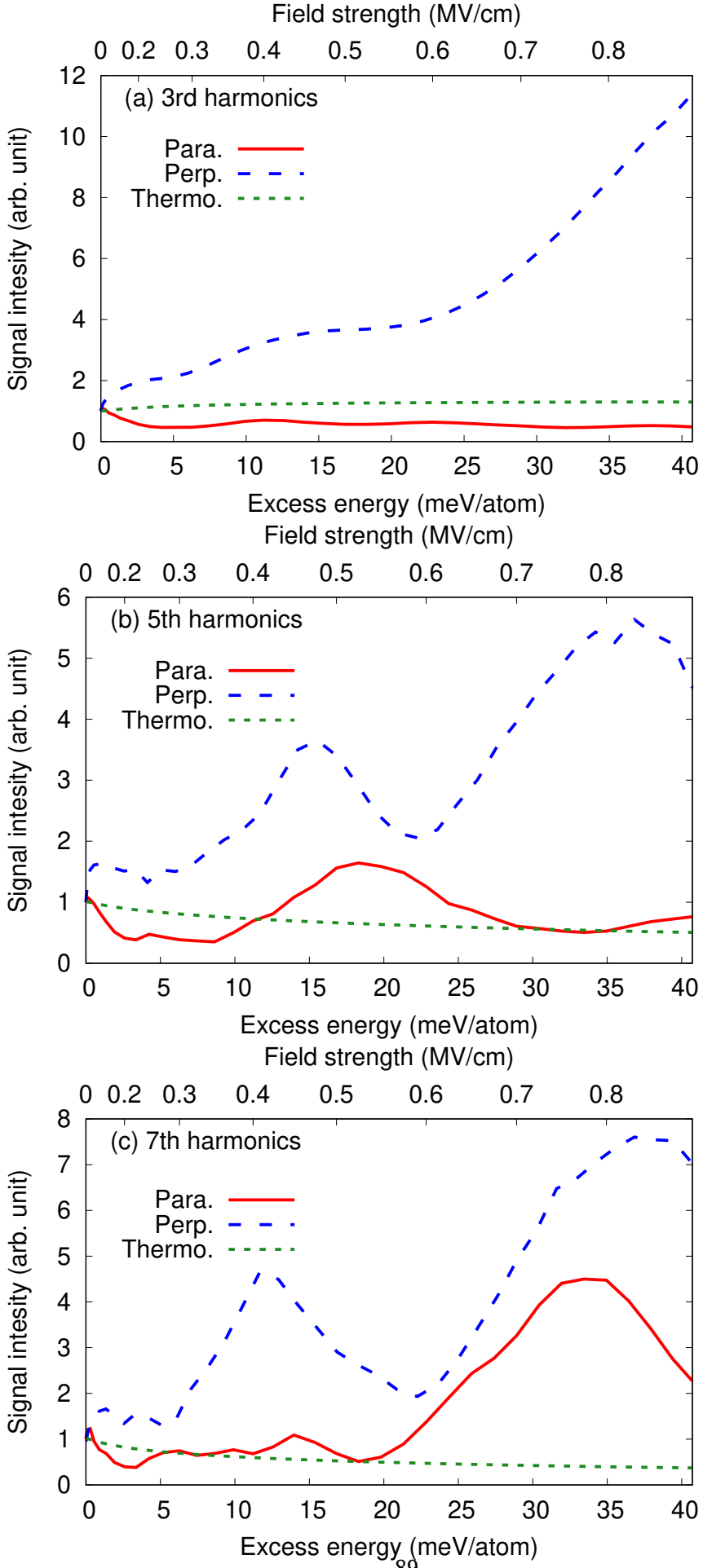


Figure 5.6: The emitted light intensity, I^{nth} , is shown as a function of the excess energy for (a) third (b) fifth, and (c) seventh harmonics. The results for the

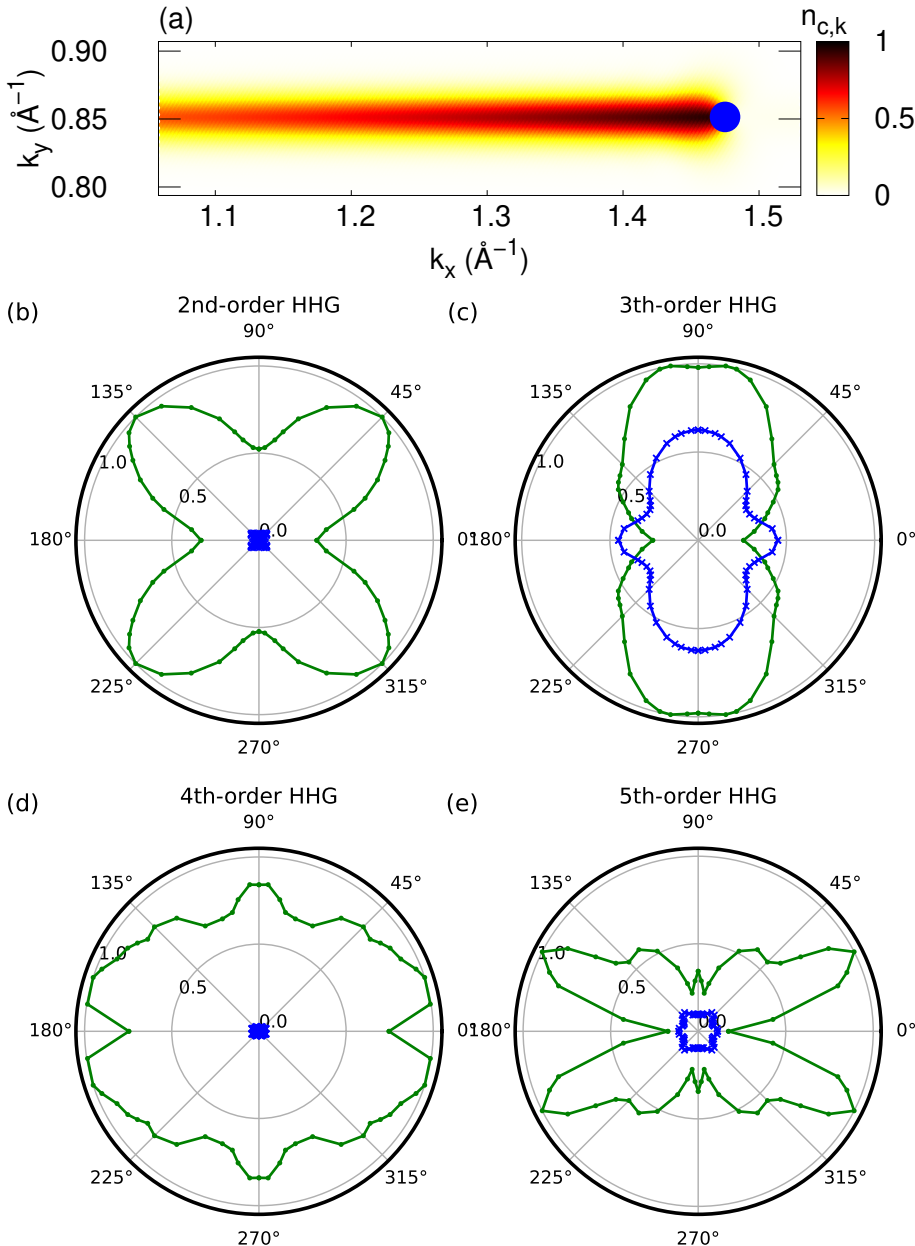


Figure 5.7: (a) The calculated conduction population distribution, $n_{ck}^{\text{neq-steady}}$ for the nonequilibrium steady-state is shown. Here, the Dirac point is indicated by the blue circle. (b–e) The angular dependence of the emitted harmonic intensity is shown for the (b) second, (c) third, (d) fourth, and (e) fifth harmonics. The results obtained using the nonequilibrium population model and the nonequilibrium steady-state are shown by the blue and green solid lines, respectively.

Chapter **6**

CONCLUSION AND OUTLOOK

CHAPTER 6. CONCLUSION AND OUTLOOK

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Appendix A

ADIABATIC BASIS REPRESENTATION

The adiabatic approximation is used almost all the time in solving time-propagation, which will be explained in more detail here. To analytically investigate nonlinear photocarrier injection in solids, we first introduce the equation of motion in the adiabatic basis representation. In this representation, we can naturally separate the interband transitions, the dynamical phase factor, and the geometric phase factor. To introduce the representation, we consider the following one-body Schrödinger equation for a \mathbf{k} -point,

$$i \frac{d}{dt} |\psi_{\mathbf{k}}(t)\rangle = H [\mathbf{k} + \mathbf{A}(t)] |\psi_{\mathbf{k}}(t)\rangle, \quad (\text{A.1})$$

where $\mathbf{A}(t)$ is an external vector potential, which is related to the external electric field as $\mathbf{E}(t) = -d\mathbf{A}(t)/dt$. In this note, we assume that the vector potential is zero for the negative time; $\mathbf{A}(t \leq 0) = 0$.

To introduce the adiabatic basis representation, we introduce the instantaneous eigenstates of the Hamiltonian as

$$H [\mathbf{k} + \mathbf{A}(t)] |u_{b,\mathbf{k}+\mathbf{A}(t)}\rangle = \epsilon_{b,\mathbf{k}+\mathbf{A}(t)} |u_{b,\mathbf{k}+\mathbf{A}(t)}\rangle, \quad (\text{A.2})$$

where b is the band index. Hereafter, we assume the two-band system, which has the valence band ($b = v$) and the conduction band ($b = c$). However, we can straightforwardly extend it to general systems.

APPENDIX A. ADIABATIC BASIS REPRESENTATION

On the basis of the instantaneous eigenstates defined by Eq. (A.2), we consider the following expansion of the wavefunction

$$|\psi_{\mathbf{k}}(t)\rangle = c_{v,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{v,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{v,\mathbf{k}}^g(t)}} |u_{v,\mathbf{k}+\mathbf{A}(t)}\rangle + c_{c,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{c,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{c,\mathbf{k}}^g(t)}} |u_{c,\mathbf{k}+\mathbf{A}(t)}\rangle, \quad (\text{A.3})$$

where $c_{b,\mathbf{k}}(t)$ are the expansion coefficients. In the expansion, we explicitly include the dynamical phase factor, $e^{-i\int_0^t dt' \epsilon_{v,\mathbf{k}+\mathbf{A}(t')}}$, and the additional phase factor, $e^{i\phi_{b,\mathbf{k}}^g(t)}$. The latter one will be defined later as the geometric phase factor.

Inserting Eq. (A.3) into Eq. (??), one obtains

$$\begin{aligned} \left[i\frac{d}{dt} - H[\mathbf{k} + \mathbf{A}(t)] \right] |\psi_{\mathbf{k}}(t)\rangle &= i\dot{c}_{v,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{v,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{v,\mathbf{k}}^g(t)}} |u_{v,\mathbf{k}+\mathbf{A}(t)}\rangle \\ &\quad + i\dot{c}_{c,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{c,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{c,\mathbf{k}}^g(t)}} |u_{c,\mathbf{k}+\mathbf{A}(t)}\rangle \\ &\quad - \dot{\phi}_{v,\mathbf{k}}^g(t)c_{v,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{v,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{v,\mathbf{k}}^g(t)}} |u_{v,\mathbf{k}+\mathbf{A}(t)}\rangle \\ &\quad - \dot{\phi}_{c,\mathbf{k}}^g(t)c_{c,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{c,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{c,\mathbf{k}}^g(t)}} |u_{c,\mathbf{k}+\mathbf{A}(t)}\rangle \\ &\quad - ic_{v,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{v,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{v,\mathbf{k}}^g(t)}} \mathbf{E}(t) \cdot \frac{\partial|u_{v,\mathbf{k}+\mathbf{A}(t)}\rangle}{\partial\mathbf{k}} \\ &\quad - ic_{c,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{c,\mathbf{k}+\mathbf{A}(t')} e^{i\phi_{c,\mathbf{k}}^g(t)}} \mathbf{E}(t) \cdot \frac{\partial|u_{c,\mathbf{k}+\mathbf{A}(t)}\rangle}{\partial\mathbf{k}} = 0. \end{aligned} \quad (\text{A.4})$$

By multiplying $e^{+i\int_0^t dt' \epsilon_{v,\mathbf{k}+\mathbf{A}(t')} e^{-i\phi_{v,\mathbf{k}}^g(t)}} \langle u_{v,\mathbf{k}+\mathbf{A}(t)}|$ to Eq. (A.4), one obtains

$$\begin{aligned} i\dot{c}_{v,\mathbf{k}}(t) - \dot{\phi}_{v,\mathbf{k}}^g(t)c_{v,\mathbf{k}}(t) - ic_{v,\mathbf{k}}(t)\mathbf{E}(t) \cdot \left\langle u_{v,\mathbf{k}+\mathbf{A}(t)} \left| \frac{\partial u_{v,\mathbf{k}+\mathbf{A}(t)}}{\partial\mathbf{k}} \right. \right\rangle \\ - ic_{c,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{c,\mathbf{k}+\mathbf{A}(t')} - \epsilon_{v,\mathbf{k}+\mathbf{A}(t')}} e^{i(\phi_{c,\mathbf{k}}^g(t) - \phi_{v,\mathbf{k}}^g(t))} \mathbf{E}(t) \cdot \left\langle u_{v,\mathbf{k}+\mathbf{A}(t)} \left| \frac{\partial u_{c,\mathbf{k}+\mathbf{A}(t)}}{\partial\mathbf{k}} \right. \right\rangle \quad (\text{A.5}) \\ = 0. \quad (\text{A.6}) \end{aligned}$$

Likewise, by multiplying $e^{+i\int_0^t dt' \epsilon_{c,\mathbf{k}+\mathbf{A}(t')} e^{-i\phi_{c,\mathbf{k}}^g(t)}} \langle u_{c,\mathbf{k}+\mathbf{A}(t)}|$ to Eq. (A.4), one ob-

tains

$$\begin{aligned} i\dot{c}_{c,\mathbf{k}}(t) - \dot{\phi}_{c,\mathbf{k}}^g(t)c_{c,\mathbf{k}}(t) - ic_{c,\mathbf{k}}(t)\mathbf{E}(t) \cdot \left\langle u_{c,\mathbf{k}+\mathbf{A}(t)} \middle| \frac{\partial u_{c,\mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right\rangle \\ - ic_{v,\mathbf{k}}(t)e^{-i\int_0^t dt' \epsilon_{v,\mathbf{k}+\mathbf{A}(t')} - \epsilon_{c,\mathbf{k}+\mathbf{A}(t')}} e^{i(\phi_{v,\mathbf{k}}^g(t) - \phi_{c,\mathbf{k}}^g(t))} \mathbf{E}(t) \cdot \left\langle u_{c,\mathbf{k}+\mathbf{A}(t)} \middle| \frac{\partial u_{v,\mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right\rangle = 0. \end{aligned} \quad (\text{A.7})$$

Combining Eq. (A.6) and Eq. (A.7), one can obtain the following matrix form,

$$i\frac{d}{dt}\mathbf{c}_{\mathbf{k}}(t) = \begin{pmatrix} \dot{\phi}_{v,\mathbf{k}}^g(t) & 0 \\ 0 & \dot{\phi}_{c,\mathbf{k}}^g(t) \end{pmatrix} \mathbf{c}_{\mathbf{k}}(t) + i\mathbf{E}(t) \cdot \begin{pmatrix} M_{11} & M_{12} \\ M_{21} & M_{22} \end{pmatrix} \mathbf{c}_{\mathbf{k}}(t), \quad (\text{A.8})$$

$$M_{11} = \left\langle u_{v,\mathbf{k}+\mathbf{A}(t)} \middle| \frac{\partial u_{v,\mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right\rangle \quad (\text{A.9})$$

$$M_{12} = e^{-i\int_0^t dt' \Delta\epsilon_{cv,\mathbf{k}+\mathbf{A}(t')} + i\Delta\phi_{cv,\mathbf{k}}^g(t)} \left\langle u_{v,\mathbf{k}+\mathbf{A}(t)} \middle| \frac{\partial u_{c,\mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right\rangle \quad (\text{A.10})$$

$$M_{21} = e^{-i\int_0^t dt' \Delta\epsilon_{vc,\mathbf{k}+\mathbf{A}(t')} + i\Delta\phi_{vc,\mathbf{k}}^g(t)} \left\langle u_{c,\mathbf{k}+\mathbf{A}(t)} \middle| \frac{\partial u_{v,\mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right\rangle \quad (\text{A.11})$$

$$M_{22} = \left\langle u_{v,\mathbf{k}+\mathbf{A}(t)} \middle| \frac{\partial u_{v,\mathbf{k}+\mathbf{A}(t)}}{\partial \mathbf{k}} \right\rangle \quad (\text{A.12})$$

where $\Delta\epsilon_{bb',\mathbf{k}+\mathbf{A}(t)}$ is defined by the difference of the single particle energies as $\epsilon_{b,\mathbf{k}+\mathbf{A}(t)} - \epsilon_{b',\mathbf{k}+\mathbf{A}(t)}$, and $\Delta\phi_{bb',\mathbf{k}}^g(t)$ is defined by the difference of the geometric phases as $\phi_{b,\mathbf{k}}^g(t) - \phi_{b',\mathbf{k}}^g(t)$. Here, the coefficient vector was introduced as

$$\mathbf{c}_{\mathbf{k}}(t) = \begin{pmatrix} c_{v,\mathbf{k}}(t) \\ c_{c,\mathbf{k}}(t) \end{pmatrix}. \quad (\text{A.13})$$

Here, we define the geometric phases as

$$\phi_{b,\mathbf{k}}^g(t) = -i \int_0^t dt' \mathbf{E}(t') \cdot \left\langle u_{b,\mathbf{k}+\mathbf{A}(t')} \middle| \frac{\partial u_{b,\mathbf{k}+\mathbf{A}(t')}}{\partial \mathbf{k}} \right\rangle \quad (\text{A.14})$$

$$= i \int_0^t dt' \frac{dA(t')}{dt'} \cdot \left\langle u_{b,\mathbf{k}+\mathbf{A}(t')} \middle| \frac{\partial u_{b,\mathbf{k}+\mathbf{A}(t')}}{\partial \mathbf{k}} \right\rangle \quad (\text{A.15})$$

$$= i \oint_{\mathbf{A}(0)}^{\mathbf{A}(t)} d\mathbf{A} \cdot \left\langle u_{b,\mathbf{k}+\mathbf{A}} \middle| \frac{\partial u_{b,\mathbf{k}+\mathbf{A}}}{\partial \mathbf{k}} \right\rangle.$$

APPENDIX A. ADIABATIC BASIS REPRESENTATION

As seen from the last expression in Eq. (A.15), the phase $\phi_{b,\mathbf{k}}^g$ depends only on the geometry of the path of the integral.

With the expression of the geometric phases in Eq. (A.15), one can rewrite the equation of motion for the coefficient vector as

$$i \frac{d}{dt} \mathbf{c}_\mathbf{k}(t) = \mathcal{H}(t) \mathbf{c}_\mathbf{k}(t). \quad (\text{A.16})$$

$$\mathcal{H}(t) = i \mathbf{E}(t) \cdot \begin{pmatrix} 0 & M_{12} \\ M_{21} & 0 \end{pmatrix} \quad (\text{A.17})$$

This is the time-dependent Schrödinger equation on the adiabatic basis, and it is closely related to the Houston basis expansion [59, 60].

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