The objective of this thesis is to elucidate the femtosecond dynamics of coupled low-energy excitations in both strongly correlated materials and artificially engineered quantum structures. By means of near-infrared pump/multi-THz probe spectroscopy and a series of technological innovations (a novel collinear four-pass Ti:Sapphire amplifier and shot-noise reduced electro-optic sampling), fundamentally new insights into the many-body physics of two representative strongly correlated materials are obtained:

- It is clarified which microscopic mechanisms underlie the formation of the charge density wave in the transition-metal dichalcogenide titanium diselenide (1T-TiSe₂).
- A study of the high-temperature superconductor YBCO reveals that there exists no temporal hierarchy between electron-electron and electron-phonon scattering processes in this system.

Furthermore, non-adiabatic activation of ultrastrong light-matter interaction between a tailor-cut photonic resonance and an electronic excitation is realized and studied on a sub-cycle timescale. This further paves the way towards the observation of novel quantum-electrodynamical phenomena.

Cover: An ultrashort light pulse (flash) transiently separates two constituent orders of the charge density wave in 1T-TiSe₂: While the exciton-like periodic arrangement of electrons (blue clouds) gets optically quenched, the periodic deformation of the crystal lattice (highlighted by transparent surface, lower half) persists in an oscillating state (motion blur).
Ultrafast low-energy dynamics of strongly correlated systems

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Chapter 1

Introduction

Electromagnetism is one of the four fundamental interactions in nature and determines the structure of atoms, molecules and solids. On the quantum level, electromagnetic interactions between charged particles occur via the exchange of photons. Quantum-electrodynamics provides a mathematical description of this process and has proven to be one of the most accurate theories in physics. In condensed matter systems, where many charged particles (on the order of $10^{23}/\text{cm}^3$) are correlated via the infinitely ranged Coulomb potential, our understanding of the resulting many-body phenomena can be considered to be very basic, at best. Contemporary models allow one to explain many fundamental structural, magnetic and electronic properties of insulators, some semiconductors and simple metals by assuming non-interacting electrons [Kit04]. Even novel materials like graphene can be understood reasonably well from the single-particle point of view [Mor10]. The situation becomes more complicated when many-body interactions play the central role in determining the electronic, magnetic, optical, and sometimes even mechanical properties of a material [Dag05, Mor10]. In the simplest case, one particular interaction mechanism correlates electrons with each other. Already this is sufficient to yield fascinating phenomena such as BCS superconductivity [Onn11] or the spin-density-wave in elemental chromium [Faw88]. These effects are now well understood and can be modeled by incorporating the relevant electronic interaction mechanism into established single-particle models [Bar57, Faw88]. For strongly correlated materials, the situation is far less clear-cut. In these systems, the intricate low-energy electromagnetic interplay between many interacting electrons and a variety of elementary excitations, such as magnons, phonons, or excitons, often results in competing ground states with different symmetries, different macroscopic properties and different low-energy excitations. A phase transition between these ground states is often induced by slight changes to a tuning parameter,
such as temperature, pressure, or doping. Many of these phases and phase transitions bear great technological potential. Colossal magnetoresistance, an orders of magnitude change in resistivity associated with a magnetic-field-induced metal-to-insulator transition [Ram97], and high-temperature superconductivity are just two famous examples. Nonetheless, in order to exploit these phenomena for future technology, science first needs to obtain a microscopic understanding of the underlying mechanisms and to find out why it is often sufficient to slightly tune a parameter to tip the balance to favor one ground state over another. With sufficient knowledge, it may ultimately be possible to engineer systems with tailor-cut properties, e.g. an increased critical temperature for superconductivity.

These challenges call for experiments that are capable of directly accessing the underlying microscopic degrees of freedom in order to study their role in many-body interactions. Typically, the relevant elementary excitations in strongly correlated materials, such as phonons [Kli07], magnons [Hut72], intraexcitonic transitions [Lei08], and collective plasma excitations [Hub01] are resonant to far- to mid-infrared radiation with photon frequencies in the range of approximately 0.1 THz to 100 THz (1 THz = 10^{12} Hz). Other elementary excitations in the terahertz window of the electromagnetic spectrum such as internal transitions of artificial quasiparticles [Por12, Mén14] and vibrational transitions in large biomolecules [Ton07] are pivotal for many other scientific fields. In recent years, this has reliably spurred significant advances in the technology employed to study the terahertz spectral region [Lee00, Heb04, Gan05, Ton07, Kam13, Sch14, Eis14]. Fourier transform infrared spectroscopy and ellipsometry [Tom05] are widely employed time integrated techniques to study the spectral fingerprints of THz optical resonances in thermal equilibrium. However, time integrated spectroscopic approaches fail to capture the very essence of the microscopic physics in strongly correlated systems: The delicate and highly dynamical interplay of different types of elementary excitations that, in fact, occurs on femtosecond timescales (1 fs = 10^{-15} s) [Ulb11]. The invention of mode-locked Ti:sapphire lasers [Spe91] together with the development of increasingly efficient emitters and detectors for THz radiation [Fer02, Ton07] paved the way for femtosecond optical-pump/multi-THz-probe spectroscopy. During the last decade, this technology has become a well established tool to study the ultrafast dynamics of low-energy elementary excitations [Hub01, Sch04, Ulb11, Kim12]. In these experiments, a femtosecond optical pump pulse prepares a non-equilibrium state of the system under study by either exciting a specific type of the correlated elementary excitations or by melting a correlation induced phase itself. A THz probe pulse that is offset by a given pump-probe delay time subsequently captures the momentary complex-valued dielectric response of the sample. By repeating this experiment as a function of the pump-probe delay time and recording the electric

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1The term multi-THz usually refers to photon frequencies above 10 THz.
field time-trace of the THz probe for each delay time, one obtains a femtosecond slow-
motion picture of the ultrafast photoinduced dynamics of the full complex THz dielectric
response [Hub01]. Femtosecond THz spectroscopy often allows one to separate spectrally
overlapping resonances via their individual femtosecond dynamics and, most importantly,
to resonantly monitor the femtosecond live interplay of low-energy excitations in strongly
correlated materials after a controlled perturbation of the system.

In this work, optical pump/multi-THz-probe spectroscopy (introduced in Chapter 2) is
technologically advanced [Por14b] and employed to elucidate the fundamental mechanisms
in systems with strong coupling between their individual low-energy excitations. Crucial
new insights are gained for two representative strongly correlated materials:

- It is clarified [Por14a] which microscopic mechanisms underly the formation of
  the charge density wave in the transition-metal dichalcogenide titanium diselenide
  (1T-TiSe2). This study, presented in Chapter 3, is the main focus of this thesis.
- The investigation of phonon and quasiparticle dynamics in the high-temperature
cuprate superconductor YBa2Cu3O7−δ, documented in Chapter 4, answers the ques-
tion of whether there exists a temporal hierarchy between electron-electron and
electron-phonon scattering processes in this system [Pas10].

A third experiment, discussed in Chapter 5, targets the ultrafast dynamics of low-energy
elementary excitations in the regime of ultrastrong coupling:

- Non-adiabatic activation of ultrastrong light-matter interaction between a tailor-cut
  photonic resonance and an electronic excitation is implemented and studied on a
  sub-cycle timescale [Por12]. This part of the work further paves the way towards
  the observation of novel phenomena in non-adiabatic quantum-electrodynamics.

Separating structural and electronic order of the charge density wave in 1T-TiSe2

1T-TiSe2 is a particularly interesting and intensely studied strongly correlated model
material. Despite its relatively simple quasi-two-dimensional structure with a unit cell
consisting of only three atoms, several many-body effects play important roles in defining
its macroscopic properties. The most prominent phenomenon is the occurrence of a charge
density wave. For more than 40 years, scientists have aimed to identify the microscopic
origin of this phase transition. The solution of this enigma may not lead to immediate
advances in our technology, yet it stands to reason that a comprehensive understanding
of relatively simple model systems is an essential prerequisite on the route to understand
and disentangle all the intertwined coupling mechanisms in more complex many-body
systems, such as high-temperature superconductors.
Chapter 1. Introduction

Figure 1.1: Schematic mechanism of a Peierls transition in a 1D metal. (a) Situation in the normal metallic phase: The free electrons (light blue) are homogeneously distributed throughout the crystal lattice (red dots) of period $a$. The half-filled conduction band (red line) is populated with electrons up to the Fermi energy $E_F$ and the Fermi wavevector $k_F = \pi/2a$. In the charge ordered phase (b), the electron density is spatially modulated with a period of $\lambda_{\text{CDW}} = 2a$. The new Brillouin zone boundary at $k = k_F$, induced by the additional periodicity, yields the formation of a band gap $2\Delta_{\text{CDW}}$ and thereby energetically lowers the populated electronic states.

The basic concept of a Peierls-type charge density wave phase transition does not hold for the case of 1T-TiSe$_2$, yet it is well suited for introducing the fundamental features of the phenomenon: In 1930, Peierls proposed that a one-dimensional metal formed by an equally spaced chain with one electron per ion is unstable [Pei55]. For the normal metallic phase, the model system implies a half-filled conduction band with a Fermi vector of $k_F = \pi/2a$, where $a$ is the lattice constant (see Figure 1.1(a)). In this case, the conduction electrons are free and delocalized throughout the crystal lattice. The instability comes from the fact that a periodic modulation of the crystal lattice with a wave vector of $k_F$ (i.e. with a periodicity of $2a$) can lower the total energy of occupied electronic states by causing the formation of a band gap at the Fermi energy $E_F$ (see Figure 1.1(b)). When the electronic energy gain outweighs the elastic energy associated with the periodic lattice distortion, the distorted phase represents the ground state of the system. The formation of a band gap at $E_F$ makes the Peierls transition a metal-to-insulator transition. In real space, the renormalization of the electronic bands implies the localization of electrons (i.e. the formation of a charge density wave) with a periodicity of $2a$. 

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For $1T$-TiSe$_2$, many different mechanisms have been proposed to explain the origin of the commensurate\textsuperscript{2} charge density wave that emerges below a critical temperature of $T_c \approx 200$ K. Previous experimental results were mainly interpreted in favor of either a scenario that the charge density wave is a result of electron-phonon coupling analogous to a Peierls-type transition (e.g. [Hol01, Ros02b]) or that exciton-like coupling of free electrons and holes drives the phase transition (e.g. [Cer07, Mon10]). While low-energy excitations such as phonons, plasmons, charge density wave collective modes, and energy gaps provide unique THz fingerprints of lattice [Hol77, Sno03, MV11] and electronic orders [Li07b], their ultrafast dynamics have not been directly and simultaneously resolved so far. In this work, these degrees of freedom are monitored on the femtosecond scale after near-infrared (NIR) photoexcitation of the system. The novel experimental insights gained this way provide strong evidence that the concerted action of excitonic electron-hole correlations and a structural Jahn-Teller-like effect creates the charge density wave in $1T$-TiSe$_2$. Furthermore, it is found that in the highly non-thermal transient state after photoexcitation, the structural component of the charge density wave can persist even when the excitonic effects are quenched. A quantum-mechanical theory based on the work from van Wezel et al. [Wez10b] corroborates the above conclusions.

**Phonon and quasiparticle dynamics in superconducting YBa$_2$Cu$_3$O$_{7-\delta}$**

Unconventional superconductors are materials in which the mechanism that stabilizes the Cooper-pairs is not exclusively based on BCS-like electron-phonon interactions. These synthetic materials have a rather complex crystal structure and bear a plethora of correlation induced phenomena. For the class of copper based “cuprate” high-temperature\textsuperscript{3} superconductors, these include commensurate and incommensurate spin and charge density waves, stripe-ordering, antiferromagnetic states and a peculiar pseudo-gap phase, that either stand in direct competition with the superconducting phase [Fau11, Cha12, Has14] or coexist with it [Hau10, Hay14, He14b]. Among the multitude of elementary low-energy excitations and their coupling mechanisms in these systems, the key ingredient for unconventional superconductivity has not yet been identified. Electron-phonon interactions, however, appear to be a pivotal factor. Although any type of purely phonon-based electron pairing mechanism fails to explain the defining property of high-temperature superconductors, evidence of significant electron-phonon contributions in cuprates has been found

\textsuperscript{2}A charge or spin density wave is termed *commensurate* if its periodicity is a rational multiple of the lattice constant and is termed *incommensurate* if not.

\textsuperscript{3}Systems that feature unconventional superconductivity with a critical temperature of $T_c \gtrsim 30$ K are usually referred to as *high-temperature* superconductors.
via various experimental techniques \cite{Bat87b, Ope99, Rez06, Iwa08}. Pump-probe studies have been utilized with the goal of establishing a temporal hierarchy of microscopic interaction processes in order to qualitatively elucidate their potential relevance to the pairing mechanism. Near-infrared \cite{Dem99, Kus08}, mid-infrared \cite{Kai00} and THz probe pulses \cite{Ave01} as well as Raman scattering \cite{Sai09} have allowed researchers to study the recombination of photogenerated quasiparticles and the recovery of the superconducting condensate following a strong optical perturbation. In a time- and angle-resolved photocollection spectroscopic (tr-ARPES) study of the system Bi-Sr-Ca-Cu-O the relaxation of the quasi-equilibrium electronic temperature was analyzed assuming a selective electron-phonon coupling \cite{Per07}. However, since these experiments do not directly monitor the lattice degrees of freedom themselves, a more detailed picture of the role of the various phonon modes during an initial non-thermal regime has been beyond reach. Femtosecond electron diffraction has the potential to follow the evolution of the lattice directly \cite{Ged07}, but current sources are not yet sufficiently advanced to observe the key lattice modes that possibly assist the correlation of electrons in high-temperature superconductors \cite{Sci11}.

In this work, NIR-pump/multi-THz-probe spectroscopy is employed to address the question of whether there exists a temporal hierarchy between electron-electron and specific electron-phonon scattering processes in the prototypical high-temperature superconductor YBa$_2$Cu$_3$O$_{6.93}$ ($T_c = 92$ K). In order to study the ultrafast interplay between the electrons and lattice, femtosecond multi-THz probe pulses resonantly trace specific phonon modes and quasiparticles during a photoinduced transition to the normal state. The results demonstrate that in the system Y-Ba-Cu-O electron-phonon scattering occurs on a time scale comparable to that of the electron-electron scattering.

Femtosecond optical control and ultrafast dynamics of low-energy light-matter mixed states in the ultrastrong coupling regime

The detailed and proven understanding of fundamental electromagnetic interaction processes on the quantum level has led theoreticians to predict fascinating, fundamentally new quantum-electrodynamical phenomena (e.g. \cite{Moo70, Ful76, Yab89, Lib07, Nat12, Wes14}). Many of these theories still await the development of novel technology that facilitates their experimental confirmation. One of the most prominent examples of the predicted phenomena is the dynamical Casimir effect: It describes the conversion of electromagnetic quantum fluctuations into real excitations when the vacuum ground state of a quantum system is modulated non-adiabatically (i.e. faster than a single oscillation of its fundamental electromagnetic eigenmodes). The first theoretical description of this process \cite{Moo70} is based on the textbook scenario of the static Casimir effect. One considers two parallel,
perfectly reflecting uncharged metallic plates that are placed in vacuum. If the distance between the plates is sufficiently small, vacuum fluctuations yield a net “Casimir force” that measurably [Bre02] pushes the plates together. In this situation, the release of virtual electromagnetic excitations out of the quantum vacuum, as described by the dynamical Casimir effect, is predicted to occur when one of the mirrors undergoes a sufficiently accelerated motion at relativistic velocities [Moo70, Ful76]. Obviously, this intuitive model scenario cannot be realized in a laboratory. Fortunately, there are analogous systems that are technologically more feasible [Lib07, Wil11] to test this idea. Very promising in this respect are light-matter coupled intersubband cavity polariton systems in the regime of so-called ultrastrong coupling [Din03, Ciu05, Gün09]. Upon a sufficiently strong, non-adiabatic modulation of the light-matter coupling strength, virtual excitations of the THz electromagnetic eigenmodes are predicted to emit as Casimir photons into the external space [Ciu05, Lib07]. In a seminal work, Günter et al. [Gün09] demonstrated sub-cycle control of ultrastrong light-matter coupling in these systems. Yet, access to the relevant optical eigenmodes was rather limited and required highly complex experimental techniques. Since the virtual excitations are predicted to emerge as real excitations of the polaritonic eigenmodes [Lib07], full access to these modes constitutes an essential prerequisite for the detection of Casimir photons.

This work introduces a novel device that enables non-adiabatic optical activation of ultrastrong light-matter coupling between an eigenmode of a one-dimensional photonic crystal and an electronic intersubband resonance while providing straightforward access to all relevant eigenmodes. Femtosecond multi-THz spectroscopy is employed to demonstrate full access to the photonic band structure and to trace its non-adiabatic switching dynamics. Finite difference frequency domain (FDFD) simulations are in quantitative agreement with the experimental data and, additionally, allow for a microscopic understanding of the electromagnetic eigenmodes of the system.
Chapter 2

Ultrafast multi-terahertz spectroscopy

This chapter introduces the femtosecond multi-THz experimental techniques employed in this work. First, operational principles and modifications of the laser source, a home-built Ti:sapphire amplifier system, are discussed. Next, generation and detection of phase-stable THz pulses are explained along with a discussion on methods of two-time THz spectroscopy. Particular focus of this chapter is set on newly developed technology: A novel, high-gain collinear four-pass Ti:sapphire amplifier and a scheme to enhance the sensitivity of shot-noise limited field-resolved detection of THz radiation (published in Optics Letters [Por14b]).

2.1. The Ti:sapphire amplifier system

The accessible parameter space and noise characteristics of femtosecond multi-THz studies critically depend on the driving laser source. As will be seen in section 2.2.1, generation and field resolved detection of THz radiation require a pulse duration of the driving laser shorter than a half-cycle of the THz wave. Since detection of THz frequency components that approach the near infrared [Küb04] is an essential prerequisite for this work, the laser source must provide pulses with a duration of \( \lesssim 15 \) fs. Within the class of table-top lasers, this criterion can be met by setups based on mode-locked Ti:sapphire oscillators [Bal97, Ell01, Hub03, Sch03, Sud05] or fiber laser systems [Sel09, Kra10]. Another stringent requirement to the laser source is that it delivers a pulse energy high enough to optically manipulate correlation effects (e.g. to drive a phase transition). For most experimental scenarios, this excludes both fiber and Ti:sapphire oscillators with pulse energies typically far below \( \sim 0.1 \) µJ. While current Ti:sapphire regenerative amplifier systems easily provide pulse energies up to several mJ, their noise characteristics renders them unsuitable for
Chapter 2. Ultrafast multi-terahertz spectroscopy

Figure 2.1: Layout and specifications of the Ti:sapphire amplifier system. (a) Cavity-dumped Ti:sapphire oscillator. BP1, BP2: Brewster-prisms. AOM: Acousto-optic modulator. (b) Novel collinear four-pass Ti:sapphire amplifier. PBS1, PBS2: Polarizing beam splitter cubes. FR: Faraday rotator. (c) Adaptive pulse recompression stage. BPP1, BPP2: Brewster-prism pairs. DM: Deformable mirror.

The studies performed here. Besides that, their comparably low repetition rate of a few kHz implies unfavorable measurement statistics.

To meet all experimental requirements for this work, a unique home-built Ti:sapphire amplifier system [Hub03] is employed. The system does not set a record with any individual feature, but closes the gap between Ti:sapphire oscillators and regenerative amplifier systems: It combines a high repetition rate of up to \( f_{\text{rep}} = 4 \text{ MHz} \) with a pulse energy of up to \( E_{\text{max}} = 1.3 \mu\text{J} \) (see section 2.1.2) and a minimum pulse duration of \( t_{\text{min}} = 12 \text{ fs} \). The central wavelength \( \lambda_c \) is tunable between 740 nm and 850 nm and a spectral bandwidth \( \delta \lambda \) of greater than 100 nm (full width at half maximum, “FWHM”) can be achieved. Excellent noise characteristics [Hub03] are maintained by employing a low noise Nd:YVO\(_4\) continuous wave (CW) pump laser. Figure 2.1 shows the layout of the system: A cavity-dumped Ti:sapphire oscillator (a) generates seed pulses for a novel Ti:sapphire four-pass collinear amplifier (b). Dispersion of the amplified pulses is compensated in a pulse compression stage (c). Figure 2.2 depicts a typical spectrum and an interferometric autocorrelation of the pulses delivered by the system. The following sections elaborate on the components of the laser system in more detail.
2.1. The Ti:sapphire amplifier system

A cavity-dumped, passively Kerr-lens mode-locked Ti:sapphire oscillator (figure 2.1(a)) based on the design by Pshenichnikov et al. [Psh94] forms the first element of the amplifier system.

Ti$^{3+}$-doped sapphire ($\text{Al}_2\text{O}_3$) has become one of the most important gain media for femtosecond solid state lasers due to its outstanding optical [Mou86, Bra00] and mechanical properties. Since ultrashort laser pulses of duration $\Delta t$ (standard deviation of the intensity envelope) comprise a large bandwidth of photon frequencies $\Delta \omega$ as demanded by energy-time uncertainty ($\Delta \omega \Delta t \geq 1/2$), the enormous fluorescence bandwidth of Ti:sapphire is key to the generation of femtosecond pulses: It allows for efficient amplification of longitudinal cavity modes with wavelengths ranging from 0.65 $\mu$m to 1 $\mu$m. The formation of femtosecond pulses requires a fixed phase-relation between the amplified modes. This is commonly achieved by exploiting the nonlinear Kerr effect [Mil10] in the Ti:sapphire gain medium itself: Intensity peaks resulting from constructive interference of the longitudinal modes are self-focused via a Kerr-lens (i.e. an intensity dependent refractive index profile with intensity dependent focal length) such that the spatial overlap of an intensity spike with the population inverted region in the gain medium is enhanced as compared to the situation in CW operation. Kerr-lens mode-locked Ti:sapphire lasers allow for the formation of near-infrared (NIR) laser pulses with a duration as short as 5 fs [Ell01].

Here, a highly doped (absorption coefficient $\alpha(532 \text{ nm}) = 6.27 \text{ cm}^{-1}$) Brewster angle cut Ti:sapphire crystal is pumped with a CW power of 4 W from a Nd:YVO$_4$ laser at a wavelength of 532 nm. The output power of the latest generation commercial pump source

![Figure 2.2: (a) Intensity spectrum and (b) interferometric autocorrelation of the pulses delivered from the Ti:sapphire amplifier system. (b) is reproduced from [Hub03]. A Gaussian fit to the autocorrelation (dashed line) reveals a pulse duration of 12 fs.](image-url)
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is actively stabilized for frequencies up to 1 MHz (integrated rms-noise: 0.02%). This is tailor-cut for the operation with Ti:sapphire lasers, since pump noise at frequencies below \(\sim 1 \text{ MHz}\) directly transfers to the output power of Ti:sapphire lasers [Mul08]. The lifetime of \(\sim 4 \mu\text{s}\) of the upper laser levels in Ti:sapphire [Mou86] accounts for this fact. An acousto-optic modulator (AOM) in the short arm of the laser cavity serves as pulse picker: An acoustic wave of nanosecond duration in a Brewster cut quartz crystal diffracts the intra-cavity laser pulse in double-pass geometry to the amplifier stage (figure 2.1(b)) with an efficiency of up to 70%. The radio frequency driver unit of the AOM is synchronized with the laser oscillator repetition rate \(\nu_{\text{osc}} = 80 \text{ MHz}\). Laser pulses can be coupled out at an integer fraction of \(\nu_{\text{osc}}\). In addition, the AOM is used to initiate mode-locked operation. Astigmatism introduced by focal points of the cavity mode inside the Brewster cut optical elements is compensated with tilted folding mirrors [Kog72]. A fused-silica prism compressor (BP1, BP2) compensates intra-cavity group velocity dispersion. For further details on the oscillator and advantages of Ti:sapphire technology, see e.g. [Bra00, Hub03, Hub04, Küb07, Mil10].

### 2.1.2. A novel collinear four-pass Ti:sapphire amplifier

The energy of the pulses dumped from the oscillator via the AOM amounts to up to \(\sim 35 \text{ nJ}\). To enable the class of experiments performed in this work (see chapters 3-5) a Ti:sapphire amplifier stage is employed to increase the pulse energy. Figure 2.1(b) shows the amplifier stage integrated into the laser system while Figure 2.3 depicts a detailed close-up view. In the first generation of the amplifier setup, a double pass geometry allowed for a maximum amplification factor of 16 for low average seed power [Hub03, Küb07]. A maximum pulse energy of 210 nJ was reached in this scheme. Nevertheless, a yet higher pulse energy extends the experimental parameter space (e.g. by allowing for an increased photoexcitation intensity to melt more robust correlation induced phases) and improves the general flexibility of the system. For example, the output of the novel amplifier setup presented below would support the operation of up to six parallel beam lines in the formerly used configuration.

The main goal in the design of laser amplifiers is to achieve an efficient transfer of energy from the pump to the seed beam. Since the power of the seed light is usually limited, the common strategy underlying all amplifier designs is to maximize the spatial and temporal overlap of the seed photons with the population inversion in the gain medium. For efficient absorption of the pump light and a high spatial density of the population inversion, a highly doped gain medium is essential. At present, three different amplifier designs implemented with Ti:sapphire technology are common. When highest pulse energy
in the multi-mJ regime is of demand, it comes at the cost of a reduction in repetition rate to typically several kHz. Amplification in this regime is either accomplished with regenerative [Kme91] or non-collinear multi-pass [Bac95] chirped-pulse [Str85] amplifiers. In both cases, the effective spatial interaction volume for pump and seed beams is increased by sending the seed pulse multiple times through the population inverted region of the gain medium. In the case of regenerative amplifiers, an active element, such as a Pockels-cell, traps a seed pulse inside a laser cavity for several round-trips. In non-collinear multi-pass amplifiers the multi-pass beam path is set by geometrical means. Due to the lifetime of the upper laser levels of $4 \mu s$ (for $T < 200 K$ [Mou86]), pulsed pump lasers are necessary to maintain temporal overlap between population inversion and seed pulses at these repetition rates. When the repetition rate approaches the MHz regime, low-noise CW pumping of the gain medium becomes possible in combination with collinear amplification schemes [Hub03, Dan07, Oza10].

In this work, a novel collinear four-pass Ti:sapphire amplifier is developed. Apparently, this concept has never been applied routinely for the amplification of laser pulses, although a slightly different amplifier scheme is documented in the literature [And94]. Even though patented [McI93], no commercial Ti:sapphire laser system seems to make use of this approach.

For four-pass collinear amplification (see Figure 2.3), the horizontally polarized seed pulse is first transmitted through a Faraday isolator consisting of a polarizing beam splitter
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(PBS1), a Faraday rotator (FR) and a half wave plate. In forward direction, the seed pulse experiences no change in polarization after passing these elements. After transmission through another polarizing beam splitter (PBS2), the first amplification pass through the CW-pumped Ti:sapphire amplifier crystal follows. Next, the combination of a quarter wave plate and a folding mirror (M2) rotates the seed polarization by 90° to vertical and focuses the beam back into the amplifier crystal for the second amplification pass. Thereafter, PBS2 and the end mirror M1 come into play by reflecting the vertically polarized seed pulse back into the amplifier crystal for the third pass. M2 and the quarter wave plate then rotate the polarization back to horizontal and send the seed pulse through the Ti:sapphire crystal for the fourth time. Now, the horizontally polarized amplified pulse again transmits through PBS2. In backwards propagation direction, the Faraday rotator and the quarter plate wave turn the polarization of the pulse to vertical and PBS1 finally separates the beam paths of incoming and outgoing pulses. In pass one and four, the seed pulse is polarized along the c-axis of the Ti:sapphire crystal, resulting in a higher gain in these passes due to a maximized gain cross section [Mou86].

The highly doped ($\alpha$(532 nm) = 6.4 cm$^{-1}$) Ti:sapphire amplifier crystal is pumped with a power of 12 W from the same pump laser as the oscillator and is mounted in an evacuated housing equipped with anti-reflection coated fused silica windows. The crystal is cooled to a temperature of 224 K via a cascade of Peltier elements to increase the lifetime of the upper laser levels and to reduce thermal lensing effects [Hub04]. An anti-reflection coated fused silica lens focuses the collinearly superimposed pump and seed beams to a focal beam waist of 20 µm (FWHM) inside the amplifier crystal. A custom made high power anti-reflection coating on both sides of the Ti:sapphire crystal allows for normal incidence of the beams without significant reflection losses. About 80% of the pump power of 12 W is absorbed in the 3 mm long crystal. This length is chosen to approximately match the two-fold Rayleigh length of the pump beam [Oza10]. Theoretically, even tighter focusing should increase the gain per pass [Oza10], but strong thermal lensing effects would be the consequence [Hub04] and impair the beam quality.

Figure 2.4 shows the amplification factor (blue dots) and average output power (red squares) of the laser system measured after the prism compressor stage as a function of the average seed power measured after the AOM. The low power amplification factor is close to 100. This represents a more than six-fold improvement compared to the performance of the previous double-pass design. With increasing seed power, the amplification factor decreases due to depletion of population inversion. The steep drop observed at comparably low seed power below 20 mW demonstrates an already very efficient depletion of the population inversion by stimulated emission. Consequently, the output power of the system scales strongly sub-linearly with the seed power. As an advantageous side effect,
2.1. The Ti:sapphire amplifier system

Figure 2.4: Net amplification factor (blue dots) and average output power (red squares) of the Ti:sapphire four-pass amplifier in combination with the pulse recompression stage as a function of the average incident NIR seed power at a repetition rate of \( f_{\text{rep}} = 4 \text{ MHz} \). The Ti:sapphire amplifier crystal is pumped with a CW power of 12 W.

The sub-linear scaling reduces the influence of seed power fluctuations on the output power. At the maximum cavity-dumping level at a repetition rate of 4 MHz, the average output power is as high as 1.6 W. A quantum efficiency of 23% for conversion of absorbed pump photons is reached in the amplifier in this case. With that, the presented four-pass amplifier operates at the same level of performance as state-of-the art regenerative amplifiers [Zha12]. A very interesting reference is given by the commercially available Coherent RegA 9000 series: Its amplifier stage generates a maximum average output power of 1.6 W with a pump power of 12 W [Coh14], but only allows for amplification of pulses with a duration of \( \approx 40 \text{ fs} \) at much lower repetition rates.

An outstandingly simple alignment protocol is another advantage of the collinear four-pass amplifier design. Slight detuning of the quarter wave plate results in the formation of a laser cavity between M1 and M2 for a vertically polarized mode. This cavity can now be optimized by maximizing the laser power coupled out through PBS2. The diameter and the direction of the beam after PBS2 reveal the extra-cavity profile of the mode with maximum amplification and thus define the optimum beam parameters for the seed beam.

In summary, for repetition rates typically used for experiments of 400 kHz, 800 kHz and 4 MHz, the maximum energy of the pulses delivered by the system is increased to 1.3 \( \mu \text{J} \), 1.0 \( \mu \text{J} \), and 0.4 \( \mu \text{J} \), respectively. This represents an up to six-fold increase compared to the double pass design [Hub03]. The other specifications of the system (see page 10) are defined by the Ti:sapphire oscillator and thus remain unchanged.
2.1.3. Dispersion management

For amplification, ultrashort pulses unavoidably pass through dispersive optical elements. Here, especially the two-fold transmission through the 2 cm long terbium-gallium-garnet \cite{Sch94} Faraday rotator and multiple passes through the 5 mm thick polarizing beam splitters fabricated of SF2 glass induce optical chirp on the NIR pulses. This stretches the pulses to a calculated duration of 2.9 ps (total group velocity dispersion: $11.2 \times 10^3 \text{fs}^2$, total third order dispersion: $5.8 \times 10^3 \text{fs}^3$). Compared to the double pass setup \cite{Hub03}, the overall chirp in the new amplifier design is increased by only $\sim 20\%$.

For recompression of the pulses (figure 2.1(c)), the group velocity dispersion is compensated by two pairs of Brewster prisms (BPP1, BPP2) \cite{Pro92}. A deformable mirror (DM) in the Fourier plane of the prism compressor allows for correction of remaining higher order chirp \cite{Zee99}. A genetic optimization algorithm is employed to determine the surface profile of the deformable mirror which is controlled by 19 electro-static actuators. Feedback for the algorithm is given either by the intensity of optical second harmonic generation from the compressed pulses or the maximum electro-optically detected electric field amplitude of a generated THz electric field transient (see section 2.2.1).

2.2. Femtosecond multi-THz spectroscopy

The NIR femtosecond pulses delivered by the laser system described above are employed to operate the setup for femtosecond NIR-pump/multi-THz-probe spectroscopy. This technique is tailor-cut to study the live interplay of strongly correlated low-energy elementary excitations. In the experiment, a NIR pump pulse first induces a non-equilibrium state of the system under study. Then, a subsequent THz probe pulse captures the momentary dielectric response at a given delay time after photoexcitation. With field-resolved detection of the THz probe transients and variation of the pump-probe delay time, one obtains 2D-maps of the femtosecond evolution of the full complex THz dielectric response without the need for a Kramers-Kronig transformation. These datasets allow for investigation of the dynamic interplay of low-energy excitations via their transient THz spectral fingerprints.

In the following section, generation of phase-stable THz probe pulses from NIR femtosecond pulses via optical rectification as well as conventional electro-optic sampling (EOS) are summarized. The subsequent section elucidates the principles of two-time THz-spectroscopy. The experimental setup employed for this work is sketched and explained in Figure 2.5.
2.2. Femtosecond multi-THz spectroscopy

Figure 2.5: NIR-pump/multi-THz-probe setup. The 12-fs Ti:sapphire amplifier system (see section 2.1) serves as laser source. A first fraction of an incident NIR pulse photoexcites the sample (beam path (i)). A mechanical beam chopper (MC) switches the pump beam at a frequency of up to 1 kHz to facilitate lock-in detection techniques. A second portion of the NIR pulse (beam path (ii)) generates the THz probe (blue beam) via optical rectification (see section 2.2.1) in a nonlinear emitter crystal (EX). The polarization of the generation beam is periodically tuned with a photoelastic modulator (PEM) to modulate the emitted THz field amplitude at a frequency of $\nu = 84$ kHz for lock-in detection. A first set of parabolic mirrors focuses the THz probe beam onto the sample (orange box), which optionally can be cooled down to a temperature $T > 4$ K with a continuous flow helium cryostat. After interaction with the photoexcited sample (depicted for transmission geometry), the THz probe is focused on the nonlinear detector crystal (DX) via a second pair of parabolic mirrors. Here, the THz probe is spatiotemporally overlapped with the third fraction of the NIR pulse, the optical gate pulse (beam path (iii)) for (shot-noise reduced) EOS (see sections 2.2.1 and 2.3). Optical delay lines in branch (i) and (ii) set the pump-probe delay time $t_D$ and the EOS delay time $t$, respectively.
Chapter 2. Ultrafast multi-terahertz spectroscopy

2.2.1. Generation and detection of phase-stable THz radiation

THz generation via optical rectification of femtosecond near-infrared pulses

In this work, tunable generation of ultrabroadband phase-stable THz pulses is realized by optical rectification (OR) of the 12-fs NIR laser pulses in a nonlinear THz emitter crystal (Figures 2.5 and 2.6). OR is a nonlinear optical $\chi^{(2)}$-process that describes difference frequency generation (DFG) between frequency components contained in the spectrum of the driving laser pulse [Bon95]. Due to the large bandwidth of the 12-fs NIR laser pulses, difference frequencies as high as several ten THz can be generated this way. OR is most efficiently driven when the phase relation between the nonlinear difference frequency polarization and the intensity envelope of the NIR pulse is constant throughout the interaction length in the emitter crystal [Bon95]. This phase-matching condition can be naturally fulfilled in birefringent crystals when the polarization states of the generation and THz photons are suitably chosen with respect to the optical axis of the crystal [Hub00, Por14b]. A typical phase-matching configuration employed in this work is shown in Figure 2.6. Due to dispersion of the ordinary and extraordinary refractive indices, phase-matching cannot be fulfilled equally well for all THz frequencies. In a very inefficient case, the phase mismatch between the THz wave and the NIR pulse envelope accumulated at the exit surface of the crystal amounts to $2\pi$ and the THz wave generated in the first half of the crystal destructively interferes with the wave generated in the second half. Sufficiently thin crystals can minimize this effect and consequently permit the generation of THz pulses with a more broadband frequency spectrum.
2.2. Femtosecond multi-THz spectroscopy

Figure 2.7: Geometry for type II phase-matched EOS of phase stable THz radiation (blue waveform) using a GaSe electro-optic sensor. For abbreviations see captions of Figures 2.6 and 2.5.

Since DFG in OR occurs between frequency components of a single NIR generation pulse, the non-stable NIR carrier-envelope offset phase [Mil10] cancels in the DFG process and the THz pulses generated via OR are inherently phase-stable, i.e. the transient electric field profile is identical in every THz pulse.

**Electro-optic sampling of phase-stable THz radiation**

One of the most important achievements of terahertz photonics is the possibility to detect free-space THz radiation with respect to the absolute phase and amplitude of its oscillating carrier wave [Fer02, Ton07]. In comparison to other technological implementations of field-sensitive detection, i.e. photoconductive antennas [Smi88] and plasma based methods [Dai06], electro-optic sampling [Wu95, Liu04, Küh04] stands out due to its excellent sensitivity. Besides its application in THz time-domain spectroscopy, EOS is widely used for sensing and imaging applications [Ton07, Moo12, Bla13]. Steady technological progress has pushed the frequency bandwidth accessible with EOS close to the near-infrared [Sel08b, Mat12]. It has been demonstrated, that for sufficiently strong THz fields, EOS readily allows for recording a full THz waveform with a single laser shot [Sha00].

For EOS, a phase-stable THz transient (blue waveform in Figure 2.7) is focused into a nonlinear optical detector crystal, where it co-propagates with an ultrashort NIR gate pulse (red waveform) [Gal99]. In a simplified picture, the THz electric field induces a quasi-instantaneous birefringence in the electro-optic sensor via the Pockels effect, leading to a phase retardation $\Delta \varphi$ between the linear polarization components of the gate pulse. This mechanism requires the gate pulse to be shorter than a half-cycle of the THz electric field. Phase matching can be employed to maximize the interaction length between the THz field and the gate pulse [Liu04, Küh04]. $\Delta \varphi$ is read out with an ellipsometer consisting...
Figure 2.8: Single-cycle multi-THz field transient generated by phase-matched OR in a 15 µm thick GaSe crystal (type II phase-matching, \( \vartheta' = 55^\circ \)) and electro-optically sampled employing another GaSe crystal with the same thickness and orientation. The amplitude spectrum of the transient is shown in Figure 2.9. The shaded curve shows the intensity envelope with a duration (FWHM) of 35 fs.

of a quarter wave plate and a Wollaston prism (WP) (see Figure 2.7). The polarization optics split the gate power equally between two identical photodiodes (PD) as long as no THz field is applied, while a THz-induced phase retardation \( \Delta \varphi \) causes an imbalance \( S = I_a - I_b \) of the photocurrents \( (I_a, I_b) \) measured in the diode pair. Repeating this procedure as a function of the delay time \( t \) between the THz and the gate pulse results in a differential signal \( S(t) \) that is directly proportional to the time trace of the THz electric field \( E_{\text{THz}}(t) \).

Spectral coverage

In this work, NIR-pump/multi-THz-probe spectroscopy is accomplished in a frequency window ranging from 2 to 56 THz. For full coverage of this more than four optical octaves spanning frequency region, three different materials are employed as emitter and detector crystals. (i) Gallium phosphide (GaP) is suited for quasi-phase-matched OR and EOS for frequencies up to 7.5 THz. Transparency in the corresponding THz frequency window is the main advantage of the material and comes from the relatively high frequency phonon reststrahlen band beginning at \( \sim 9 \) THz [Bar68]. (ii) Gallium selenide (GaSe) is employed to cover frequencies from 8 to \( \sim 40 \) THz. Birefringent GaSe has become a standard material for multi-THz generation due to its wide phase-matching tuning range and its high nonlinearity [Liu04, Küb04]. (iii) Birefringent silver thiogallate (AgGaS\(_2\), AGS) is introduced with this work to cover frequencies above \( \sim 30 \) THz.
2.2. Femtosecond multi-THz spectroscopy

Figure 2.9: Linear amplitude spectra of THz transients recorded with EOS. The figure provides an overview of the spectral coverage of the experimental setup. Orange filled curve: 250 µm thick GaP crystals as emitter (EX) and detector (DX). Red filled curve: 15 µm thick GaSe as EX and DX, $\vartheta' = 55^\circ$ (spectrum of the single cycle transient shown in Figure 2.8). Yellow filled curves: AGS as EX and DX. Blue curve, yellow filling: 50 µm thick AGS crystals as EX and DX, $\vartheta = 53^\circ$, $\varphi = 45^\circ$. Green curve, yellow filling: 200 µm thick EX, 100 µm thick DX, $\vartheta = 60^\circ$, $\varphi = 45^\circ$. Horizontal arrows indicate the spectral range where OR and EOS is most efficient with the stated nonlinear crystal.

Figure 2.8 shows a typical multi-THz transient generated with phase-matched OR and recorded with phase-matched EOS. The duration of the waveform is practically bandwidth-limited and the electric field performs only one single optical cycle. The temporal intensity envelope (red shaded area) is as short as 35 fs (FWHM). GaSe is used as emitter and detector crystal. Phase-matching for OR and EOS is implemented in type II configuration (see Figures 2.6 and 2.7) with an external phase-matching angle of $\vartheta' = 55^\circ$. The ultrabroadband amplitude spectrum of the single-cycle THz transient is depicted in Figure 2.9 (red shaded curve). It contains frequency components from 8 to more than 40 THz. The type II phase-matching configuration together with the large external phase-matching angle close to the Brewster angle minimizes internal reflections in the $z$-cut GaSe crystals and thus suppresses undesired replicas of the main pulse [Küb07]. To fulfill phase-matching sufficiently for all THz frequency components contained in the spectrum, extremely thin GaSe crystals (thickness: 15 µm) are employed. These single-cycle transients are used to probe the electronic degree of freedom in $1T$-TiSe$_2$. For further details on phase-matching geometries in GaSe for OR and EOS see e.g. [Hub04, Küb04, Lin04, Küb05, Küb07, Rei07].

The orange shaded curve in Figure 2.9 shows the amplitude spectrum of THz transients generated and recorded with 250 µm thick GaP as emitter and detector crystals. This
configuration covers the spectral range from 2 to 7.5 THz and is employed to study the lattice degree of freedom in $1T$-TiSe$_2$.

THz transients with the amplitude spectra shown as yellow shaded curves in Figure 2.9 are generated and recorded using AGS as emitter and detector crystals. While the second order nonlinearity in AGS is estimated to be slightly lower than that of GaSe [Rot00, K"ub04], its tuning range for phase-matching is even wider: The phase velocity of frequencies up to 90 THz can be matched with the group velocity of NIR femtosecond pulses. Despite this fact, application of AGS for OR and EOS has never been reported. Simulation of phase-matched OR [Bon95] in AGS based on a Sellmeier parametrization of its refractive indices [Fan84] reveals most broadband phase-matching in type I configuration (i.e. extraordinary and ordinary polarized NIR generation light, ordinary polarized THz radiation). This requires a crystal orientation of $\phi = 45^\circ$ [Rot00]. In contrast to GaSe, which can be cleaved only along the $z$-plane, single-crystals of AGS can be cut along any arbitrary plane and, hence, any internal phase-matching angle $\vartheta$ (angle between NIR/THz wavevector and optical axis) can be realized. In GaSe, $\vartheta$ is practically limited to $18^\circ$, resulting in a highest possible phase-matchable frequency of $\sim 40$ THz. By using AGS crystals with a thickness of 50 $\mu$m in the orientation described above and by setting $\vartheta = 53^\circ$, THz transients with an amplitude spectrum shown by the yellow shaded blue curve are obtained. More stringent phase-matching in thicker crystals (emitter: 200 $\mu$m, detector: 100 $\mu$m) tuned for even higher THz frequencies by increasing $\vartheta$ to $60^\circ$ yields THz pulses with a center frequency as high as 52 THz. The yellow filled green curve shows the corresponding amplitude spectrum. In summary, by using AGS crystals for OR and EOS, the spectral range accessible for femtosecond spectroscopy with the present setup could be extended to cover photon frequencies of up to 56 THz.

2.2.2. Two-time THz spectroscopy for pump-probe studies

Probing the dielectric response of rapidly evolving non-equilibrium systems is a non-trivial task (for comprehensive reviews see e.g. [Kin99, N"em02, N"em05]). Conventional intensity based pump-probe experiments often suffer from limited time resolution and do not allow to directly access real and imaginary parts of the optical response functions simultaneously. In contrast, the unique possibility of two-time THz spectroscopy to resolve both the absolute phase and amplitude of a THz probe pulse that has interacted with a photoexcited sample at a given pump-probe delay time provides an ideal solution.

In an equilibrium system, the time-dependent polarization $P(t)$ induced by an electromagnetic waveform $E(t')$ is given by the convolution of the electric probe field with the
2.2. Femtosecond multi-THz spectroscopy

Figure 2.10: (a) In thermal equilibrium, a THz probe transient (depicted as a delta-like maximum, blue spike), incident at time $t'$, induces a polarization response, recorded electro-optically at time $t$ with absolute amplitude and phase (green wave). (b) The non-equilibrium response following optical excitation with a pump pulse (red spike) at time $t''$ is probed by a THz field (blue spike) at time $t'$, and the subsequent polarization (green wave) is recorded electro-optically at time $t$.

time-domain dielectric response function $\varepsilon(t)$:

$$P(t) = \varepsilon_0 \int_{-\infty}^{t} E(t') \left[ \varepsilon(t - t') - 1 \right] dt'$$

(2.1)

Causality requires that the integral extends over all times $t'$ prior to the time of observation $t$. Figure 2.10 (a) illustrates the situation for the case of a delta-like probe pulse. When the waveform of the incident probe field $E(t')$ is known, it is sufficient to measure the polarization $P(t)$. After Fourier transformation, the convolution decomposes into a product and the dielectric function $\varepsilon(\nu)$ is directly proportional to $P(\nu)/E(\nu)$. Both the probe $E(t')$ and the polarization trace $P(t)$ are captured with complete amplitude and phase information via EOS. This way, the full complex dielectric function is accessible without the need for a Kramers-Kronig transformation.

If photoexcitation induces a non-equilibrium state of the system, the response function itself may rapidly change while the induced polarization is recorded. This scenario is schematically depicted in Figure 2.10 (b): A pump pulse incident at time $t''$ drives the system out of equilibrium. Subsequently, an electric probe field (shown as a blue delta peak) at time $t'$ induces a polarization response. Electro-optic sampling permits to capture this response with absolute amplitude and phase as a function of the EOS delay time $t - t'$. 
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It is important to note that the dielectric response of a non-equilibrium system cannot be described by a one-time-dependent dielectric function since the system evolves in time while its polarization is recorded. For a proper description, \( \varepsilon \) has to be represented by a function of two independent time intervals, namely \( t - t'' \) and \( t - t' \), such that the polarization response of the photoexcited system reads \cite{Kin99}

\[
P(t, t'') = \varepsilon_0 \int_{-\infty}^{t} E(t') \left[ \varepsilon(t - t', t - t'') - 1 \right] dt'
\] (2.2)

It is not straightforward to retrieve the two-time response function from the above equation. In particular, it is not possible to deconvolve equation (2.2) by recording \( P(t, t'') \) for a fixed pump time \( t'' \) and subsequently performing a Fourier transformation since both arguments of \( \varepsilon \) depend explicitly on \( t \). It has been shown \cite{Kin99}, however, that this problem can be resolved when, instead of a single polarization transient \( P(t, t'') \) with a set pump time \( t'' \), one considers a set of data points taken from different pulses, measured with a fixed time delay \( t_D = t - t'' \) with respect to the pump pulse. Experimentally, acquisition of data in this way is achieved by keeping the relative arrival time between the pump and the electro-optic gate pulse fixed while the THz probe transient is continuously shifted in time. This is implemented in the setup (see Figure 2.5) by moving the delay line in branch (ii) and keeping the delay in branch (i) fixed. Ultimately, this approach allows one to retrieve

\[
P(t, t - t_D) = \varepsilon_0 \int_{-\infty}^{t} E(t') \left[ \varepsilon(t - t', t_D) - 1 \right] dt'
\] (2.3)

Since \( t_D \) is kept constant, \( P(t, t - t_D) \) can be deconvolved from \( E(t') \) to yield \( \varepsilon(t - t', t_D) \) or, after Fourier transformation, \( \varepsilon(\nu, t_D) \). It is to note that the sampling delay time \( t - t' \) and the pump-probe delay time \( t_D \) are not connected with each other by an uncertainty product. The maximum experimental time resolution is thus not limited by the duration of the probe pulse, but essentially set by the response time of the electro-optic detector \cite{Kin99}. With a 15 \( \mu \)m thin GaSe sensor, the dynamics of \( \varepsilon(\nu, t_D) \) can be retrieved with a time resolution of better than 40 fs.

2.2.3. Data acquisition and analysis

A THz probe transient acquired in the way described above directly encodes the polarization response of a photoexcited sample \( P(t, t - t_D) \). In principle, by comparing this transient with a reference waveform recorded without the sample in place, the momentary dielectric response of the sample \( \varepsilon(\nu, t_D) \) can be determined.
Another, experimentally more robust approach to measure $\varepsilon(\nu, t_D)$ is usually chosen when the dielectric response of the sample in thermal equilibrium $\varepsilon(\nu)$ is known. It is then sufficient to record the electric field that has interacted with the sample in thermal equilibrium $E_{\text{eq}}$ and the pump-induced changes $\Delta E$. With serial lock-in detection one can record both waveforms simultaneously [Küb07]. This approach is implemented as follows: A photoelastic modulator (PEM in Figure 2.5) modulates the THz pulse train with a frequency of 84 kHz. A first lock-in amplifier synchronously demodulates the current imbalance at the photodiodes and reads out the electro-optic signal. At the same time, the power of the NIR pump is modulated with a mechanical chopper (MC) at a much lower frequency of up to 1 kHz. Thus, for a sufficiently low integration time constant, the first lock-in amplifier measures alternately the electric field $E_{\text{eq}}$ of a waveform that has interacted with the equilibrium sample and the electric field $E_{\text{xc}} = E_{\text{eq}} + \Delta E$ of a waveform that has interacted with the excited sample. While integration of the lock-in amplifier output yields the average reference field $E_{\text{avg}} = \frac{1}{2}(E_{\text{eq}} + E_{\text{xc}})$, demodulation of its output with a second lock-in amplifier at the frequency of the MC in the pump beam yields $\Delta E$. From these quantities, $E_{\text{eq}}$ and $E_{\text{xc}}$ are directly calculated.

After recording the full waveforms, Fourier transformation along the time axis $t$ for a fixed $t_D$ yields $E_{\text{eq}}(\nu)$ and $E_{\text{xc}}(\nu, t_D)$. Since these quantities encode the polarization response of the entire probed volume, the exact geometry of the probe interaction of the THz pulse with the entire sample structure has to be taken into account for extraction of $\varepsilon(\nu, t_D)$. The optical transfer-matrix formalism is a convenient way to calculate the complex valued field transfer coefficient (i.e. the complex field reflection or transmission coefficient) of an arbitrary layered structure as a function of its refractive indices [Bor99]. It is employed here to express the field transfer coefficient of the excited sample

$$t_E(\nu, t_D) = \frac{E_{\text{xc}}(\nu, t_D)}{E_{\text{ref}}(\nu)}$$

(2.4)

as a function of $\varepsilon(\nu, t_D)$. In equation (2.4), $E_{\text{ref}}(\nu)$ corresponds to the reference electric field, that would be recorded without the sample in place. Since this quantity is usually not captured in the experiment, one may consider $t_E$ factorized as

$$t_E(\nu, t_D) = t_{E,\text{pp}}(\nu, t_D) \cdot t_{E,\text{eq}}(\nu) = \frac{E_{\text{xc}}(\nu, t_D)}{E_{\text{eq}}(\nu)} \cdot \frac{E_{\text{eq}}(\nu)}{E_{\text{ref}}(\nu)}.$$  

(2.5)

Here, the equilibrium complex field transmission coefficient $t_{E,\text{eq}}(\nu)$ can be expressed via the dielectric response of the sample in thermal equilibrium, eliminating $E_{\text{ref}}$. With the pump-induced change $t_{E,\text{pp}}(\nu, t_D)$ given by the experiment, the full field transfer coefficient of the photoexcited sample $t_E(\nu, t_D)$ is known and can be equated with the transfer-matrix expression. Numerical inversion of the transfer-matrix formalism then allows for extraction of $\varepsilon(\nu, t_D)$. 

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2.3. Shot-noise reduced electro-optic sampling

Currently, the ultimate limit of detector sensitivity in EOS is set by the quantum granularity, i.e. the shot-noise, of the gate laser pulses [Küb04, Mil10]. An overall increase of the sensitivity in EOS techniques may ultimately enable novel types of experiments in THz quantum optics including single-shot sampling of few-photon squeezed THz pulses or detection of THz photon bunches emitted from the quantum vacuum [Ciu05, Gün09, Por12]. Recent efforts to optimize the detector performance have aimed for improved detection electronics [Dar11], but have not diminished the shot-noise itself. This work presents a method that lowers the shot-noise in EOS and allows for reduction of the EOS acquisition time by more than one order of magnitude.

Balanced differential detection largely eliminates technical noise, such as excess power fluctuations of the gate pulses, since it typically affects the currents in both photodiodes equally. In state-of-the-art EOS, technical noise can be routinely suppressed to a level that the shot-noise of the gate pulses remains as the dominant noise source. Shot-noise arises from the quantization of the light field: The number of photons incident on each of the photodiodes follows a non-deterministic quantum distribution and can, hence, not be fully balanced. For a train of coherent gate pulses with an average power $P_g$, Poissonian statistics generates shot-noise $\Delta_S$ that scales with $\sqrt{P_g}$ [Mil10]. Since the electro-optic signal $S$ itself is directly proportional to $P_g$ [Gal99], the maximum signal-to-noise ratio (SNR) should, in principle, rise with $\sqrt{P_g}$. In practice, $P_g$ is often limited by available laser power or undesired higher-order nonlinear processes in the DX (e.g. two-photon absorption). Furthermore, eliminating the excess noise sufficiently to reach the shot-noise level is more challenging for high $P_g$.

![Schematic spectra and polarization directions](image)

**Figure 2.11:** Schematic spectra and polarization directions of the incident gate pulse (red) (centered at $\nu_g$) and of the sum frequency photons (blue) if the THz frequency $\nu_{\text{THz}}$ is small (a) and comparable (b) to the bandwidth of the gate spectrum. The electro-optic signal is located at the overlap of both spectra (gray shaded area).
2.3. Shot-noise reduced electro-optic sampling

Figure 2.12: Setup for spectrally filtered EOS consisting of a nonlinear crystal (NLC) and an ellipsometer (λ/4 plate; Wollaston prism (WP); photodiodes (PD)). Black waveform: Schematic THz field transient. The nonlinear optical interaction of the linearly polarized incident gate (red pulse) with the THz wave generates new frequency components at perpendicular polarization (blue pulse). For spectrally post-filtered EOS, an optical spectral filter (SF) is inserted behind the NLC.

In the following it will be shown, that even for shot-noise limited detection the SNR can be further enhanced in commonly used EOS configurations. In order to explain the underlying idea, EOS is described in the more rigorous picture of frequency mixing between the gate and the THz photons [Gal99] (i.e. as the inverse quantum effect of optical rectification). Depending on the interaction geometry, the $\chi^{(2)}$ susceptibility of the DX can give rise to either sum or difference frequency generation between the gate and the THz pulse. These nonlinear interactions create a phase-coherent replica of the broadband gate spectrum which is up- or down-shifted by the THz frequency (see Figure 2.11). The newly generated photons contain polarization components perpendicular to the incident gate light. Interference of converted and fundamental photons alters the polarization state in the frequency region where both spectra overlap. When the resulting polarization is analyzed in a base rotated by $\pi/4$ with respect to the incident gate polarization, the phase difference between the orthogonal polarization components ($\Delta \varphi$) scales linearly and sign-sensitively with the THz electric field. Note that the electro-optic signal is solely generated by photons from the spectral overlap region. When the frequency of the THz photons $\nu_{\text{THz}}$ is small compared to the bandwidth $\delta_g$ of the gate spectrum, the interference occurs practically throughout the gate spectrum and the Pockels effect picture remains valid (see Figure 2.11(a)). If $\nu_{\text{THz}}$ is comparable to $\delta_g$, the overlap region containing the electro-optic signal is located only at the wings of the gate spectrum and the electro-optic signal is reduced (see Figure 2.11(b)). In this situation, still all gate photons contribute equally to the shot-noise. Thus, a spectral filter (SF) inserted into the gate beam after the NLC (see Figure 2.12) can be used to select photons that contribute to the electro-optic signal and block those which only generate noise.
To quantify the influence of spectral filtering on the SNR, the following analysis assumes perfect phase-matching for sum frequency generation, a frequency independent nonlinear susceptibility $\chi^{(2)}$ and bandwidth limited pulse durations. Figure 2.13(a) schematically depicts typical spectra (solid lines) commonly seen in multi-THz spectroscopy based on Ti:sapphire lasers [Küb04, Liu04]. The red line shows the field amplitude spectrum of the incident gate pulses $E(\nu)$ centered at a frequency of $\nu = \nu_g$. The sum frequency spectrum $A(\nu - \nu_{THz})$ and the THz spectrum $A_{THz}(\nu)$ are shown in blue and black, respectively. The sum frequency spectrum results from the gate spectrum and a given THz component $\nu_{THz}$. Under the above assumptions, the spectral density of the electro-optic signal for a given frequency $\nu_{THz}$ (gray shaded area) is proportional to $\nu A(\nu) A(\nu - \nu_{THz})$ [Gal99]. Integrating over $\nu$ yields the electro-optic response as a function of $\nu_{THz}$. In a conventional EOS setup, the photodiodes account for the spectral integration. If the gate pulses are spectrally filtered with a high-pass (HP) optical filter, the lower integration boundary is set by the cut-on frequency $\nu_{HP}$. In summary, the spectral signal amplitude $S$ then depends on the interacting pulses as follows:
2.3. Shot-noise reduced electro-optic sampling

\[ S(\nu_{THz}) \propto A_{THz}(\nu_{THz}) \int_{\nu_{HP}}^\infty \nu A(\nu) A(\nu - \nu_{THz}) d\nu \quad (2.6) \]

The shot-noise amounts to:

\[ \Delta S \propto \sqrt{\int_{\nu_{HP}}^\infty \left( (1-\eta) |A(\nu)|^2 + \eta |A(\nu - \nu_{THz})|^2 \right) d\nu} \quad (2.7) \]

Equation (2.7) explicitly takes into account that for large quantum efficiencies \( \eta \) of the sum frequency process, a large number of incident gate photons may be converted from frequencies below to above \( \nu_{HP} \) and pass the filter. Figure 2.13(b) compares \( S \) (green curve) and \( \Delta S \) (orange curve) as a function of \( \nu_{HP} \) for \( \nu_{THz} = 45 \text{ THz} \), the gate spectrum of Figure 2.13(a) and assuming \( \eta = 1 \times 10^{-4} \). \( S \) remains unaffected for \( \nu_{HP} \ll \nu_g \), but starts to decrease when \( \nu_{HP} \) shifts through the overlap region between the fundamental and sum frequency spectra. In contrast, the noise drops faster with increasing \( \nu_{HP} \). A clear reduction already sets in when \( \nu_{HP} \) reaches the low-frequency wing of the incident spectrum, i.e. below the overlap region with the sum frequency spectrum. This leads to an improvement of the SNR (black curve), which exhibits a well-defined maximum before it drops towards zero. If no other noise source becomes dominant for a lower overall photon fluence, this peak identifies the cut-on frequency for maximum SNR.

To test this idea experimentally, spectrally filtered EOS is implemented with a series of commercial standard optical high pass filters. To this end, THz pulses are generated by OR of the 12-fs NIR laser pulses in a 200 \( \mu \text{m} \) thick AGS crystal. A 100 \( \mu \text{m} \) thick AGS crystal is used for EOS. With phase-matching angles of \( \phi = 45^\circ \) and \( \vartheta = 57^\circ \) THz transients centered at 45 THz are obtained. To implement the sum frequency generation process in EOS, the polarization of the incident THz and gate pulse is set to ordinary and extraordinary, respectively. The energy, bandwidth and center frequency of the gate pulses are \( E_g = 20 \text{nJ}, \delta_g = 42 \text{ THz} \) and \( \nu_g = 380 \text{ THz} \). The calculated shot-noise relative to the overall current in the photodiodes amounts to \( \Delta S/(I_a + I_b) = 9 \times 10^{-9} \text{ Hz}^{-1/2} \). In order to visualize the corresponding experimental noise floor in the electro-optic signal, the THz power is attenuated to an estimated number of \( \sim 10^6 \) THz photons per pulse and a short lock-in integration time of 100 \( \mu \text{s} \) for a given delay time \( t \) is employed. This way, the same multi-THz waveform with three different choices of \( \nu_{HP} \) is recorded (Figure 2.14(a)). As seen by comparison of the black waveform recorded with conventional EOS and the red waveform acquired with a HP filter in place (\( \nu_{HP} = 375 \text{ THz} \)), spectral filtering strongly improves the SNR. Increasing \( \nu_{HP} \) from 375 THz to 390 THz (blue waveform) yields further improvement. Quantitatively, the SNR increases by a factor of 2.9 for \( \nu_{HP} = 390 \text{ THz} \) as
Figure 2.14: (a) Multi-THz field transients recorded with phase-matched EOS in AGS without spectral filtering (black curve) or employing high-pass filtering with a cut-on frequency of 375 THz (red curve) and 390 THz (blue curve). For better visibility of the noise floor, the THz pulse is attenuated before EOS. The panel on the left magnifies the amplitude in the red shaded time window by a factor of 6. (b) Amplitude spectra of the transients shown in (a) with the same color coding on a logarithmic scale.

compared to conventional unfiltered EOS. Figure 2.14(b) depicts the amplitude spectra of the three waveforms. The strongly suppressed noise background due to spectral filtering is equally well visible in frequency domain.

For the experimental parameters (\( \nu_g = 380 \text{THz} \), \( \nu_{\text{THz}} = 45 \text{THz} \), \( \delta_g = 42 \text{THz} \), \( \eta = 1 \times 10^{-4} \), \( \nu_{\text{HP}} = 390 \text{THz} \)), the theory predicts an improvement of the SNR by a factor of 1.9. The experimentally observed gain in SNR thus even exceeds the theoretical value. This fact can be attributed to the suppression of a remaining fraction of technical noise induced by fluctuations of the gate light. Without the spectral filter in place, the actual experimentally observed noise on the recorded transient is above the calculated shot-noise level by a factor of \( \sim 1.5 \). Since the technical noise scales linearly with the power impinging on the photodiodes, reducing the overall photon fluence lowers the influence of technical noise as well. Accounting for both effects, the overall improvement by spectral filtering is perfectly explained.

Shot-noise reduction alone may even allow for larger improvement factors for other experimental parameters. For the assumptions made above, figure 2.15 summarizes the theoretical gain in the SNR as a function of the relative THz frequency \( \nu_{\text{THz}}/\delta_g \) and the relative difference \( (\nu_{\text{HP}} - \nu_g)/\delta_g \) and holds for \( \delta_g \ll \nu_g \). The black curve in Figure 2.13(b)
2.3. Shot-noise reduced electro-optic sampling

Figure 2.15: Calculated sensitivity $S/\Delta S$ of spectrally post-filtered EOS for the sum frequency generation process. The abscissa represents the THz frequency $\nu_{THz}$ normalized to $\delta_g$. The ordinate corresponds to the cut-on frequency of the HP filter $\nu_{HP}$ relative to $\nu_g$ in units of $\delta_g$. The region below the vertical scale break corresponds to the sensitivity of conventional EOS. Contour lines are spaced by a factor of 1.2. The Figure holds for $\delta_g \ll \nu_g$ and $\eta = 1 \times 10^{-4}$. An increase of $S/\Delta S$ by more than a factor of 5 can be obtained for configurations located in the purple shaded region. The coordinate corresponding to the experimental configuration with $\nu_{HP} = 390$ THz is marked with a black cross.

corresponds to a re-normalized vertical cut at $\nu_{THz}/\delta_g = 1$. For sampling of THz pulses centered at $\nu_{THz} = 1.3 \times \delta_g$, the SNR is expected to improve by a factor of 5.1 if the gate spectrum is high-pass filtered at $\nu_{HP} = \nu_g + \delta_g$.

In summary, this work introduces a universal method to enhance the sensitivity of EOS. It is shown, that this approach lowers the shot-noise significantly and is able to additionally suppress technical noise below a reduced shot-noise level. Existing EOS setups can be easily upgraded to enhance the SNR by up to 5 times saving as much as a factor 25 in measurement time, by adding a cost-effective spectral filter that can be selected with the presented analysis.

This novel concept has been employed consistently for the high-sensitivity studies of ultrafast photoinduced phase-transitions presented in the next chapter.
Chapter 3

Transient separation of excitonic and structural order of the charge density wave in 1T-TiSe₂

This chapter presents a femtosecond multi-THz study revealing the microscopic nature of the charge density wave (CDW) in 1T-TiSe₂. Ultrabroadband THz pulses are employed to simultaneously trace the ultrafast evolution of coexisting lattice and electronic orders. It is demonstrated that two components of the CDW order parameter – excitonic correlations and a periodic lattice distortion (PLD) – respond vastly differently to 12-fs NIR photoexcitation. Even when the excitonic order of the CDW is quenched, the PLD can persist in a coherently excited state. The underlying mechanism of the CDW formation is consistently identified as a cooperative coupling of excitonic correlations and a structural Jahn-Teller-like effect. A quantum-mechanical model [Wez10b] is employed to corroborate the interpretation of the experimental results. This part of the work has been published in Nature Materials [Por14a].

3.1. Transition-metal dichalcogenides and the role of 1T-TiSe₂

Transition-metal dichalcogenides (TMDs) are a family of layered compounds with chemical composition MX₂. M is a transition metal element (group 3 to 10 of the periodic table of the elements) and X is a chalcogen (group 16). Covalent bonds among the atoms result in layers of the form X–M–X with chalcogen atoms in two hexagonal planes separated by a plane of metal atoms. The layers themselves are weakly bound by Van der Waals interaction [Wil69]. The stacking of the layers is described by the polytype (e.g. 1T, 2H),
where the digit counts the number of layers per unit cell and the character abbreviates the symmetry of the unit cell (e.g. $T$ for trigonal, $H$ for hexagonal). Due to the weak interlayer coupling, TMDs can be considered as quasi two-dimensional structures.

The reduction of dimensionality has important consequences [Gru94]: Electronic correlations, random potentials and fluctuations have a more profound effect than in 3D systems. In TMDs, this leads to the formation of CDWs (e.g. in $1T$-TiSe$_2$, $1T$-TaS$_2$, $2H$-TaSe$_2$ or $2H$-NbSe$_2$) [Wil75], superconductivity (e.g. in $1T$-TiSe$_2$, $1T$-TaS$_2$ [Sip08] or $2H$-NbSe$_2$ [Yok01]), room temperature excitons (e.g. in WSe$_2$ or MoS$_2$) [Coe87] or a Mott insulating state (in $1T$-TaS$_2$) [Sip08]. A second asset of the weak out-of-plane interactions is the possibility to fabricate two-dimensional layers of single unit cell thickness via mechanical exfoliation techniques. Especially the rise of graphene [Gei07] attracted an increased interest in monolayered TMDs and their versatile electronic properties [Wan12]: One expects this class of two-dimensional systems to open up new possibilities in nano- and optoelectronics [Wan12] as well as in chemistry for batteries and supercapacitors [Chh13].

Among all TMDs, $1T$-TiSe$_2$ has attracted special attention. Its distinctive, semi-metallic band structure [Zun78, Wez10a] facilitates remarkably strong electron-hole correlations [Cer07]. In addition, strong electron-phonon interactions are present [Web11]. Although the material has been studied since the 1960s, the simultaneous existence of both correlation effects has so far hindered the identification of the microscopic mechanism behind the formation of a commensurate charge ordered phase below $T_c \approx 200$ K. Recently, $1T$-TiSe$_2$ regained interest due to the discovery of two superconducting phases and novel, chiral properties of the CDW [Ish10, Zen13]. A lattice incommensurate CDW phase constitutes the latest discovery in this material [Joe14]. Superconductivity in $1T$-TiSe$_2$ is accompanied by suppression of the CDW and can be induced by pressure [Kus09] and impurity intercalation with Cu [Mor06] or Pd atoms [Mor10]. It remains elusive, whether both superconducting phases are part of the same superconducting region in the 3D pressure/Cu-intercalation/temperature phase space or if they underly completely different mechanisms. Impurity intercalation between the layers may increase the interlayer coupling, thereby reduce the 2D confinement for electrons and tip the balance in favor of 3D BCS-like superconductivity [Mor06]. This mechanism cannot account for pressure induced superconductivity, which is suggested to be of a rather unconventional, excitonic nature [Kus09]. However, it is assumed that superconductivity in both cases is closely related to the CDW phase [Mor06, Kus09]. A recent study suggests that phase fluctuations in CDW ordered domains may be connected to pressure induced superconductivity [Joe14]. In any case, to ultimately unravel the nature of the possibly unconventional superconductivity in $1T$-TiSe$_2$, it is essential to understand the dominant correlation induced phenomenon first: The formation of a lattice commensurate CDW.
3.2. The CDW phase transition

3.2.1. Reconstruction of the crystal lattice and the electronic band structure

In its high-temperature phase, 1T-TiSe$_2$ belongs to the $P\bar{3}m1$ space group with threefold symmetry about the $c$-axis. Figure 3.1(a) shows the normal phase unit cell of 1T-TiSe$_2$. The unit cell parameters are $a = b = 3.54$ Å (in-plane distance of Ti atoms) and $c = 6.001$ Å (out-of-plane distance of Ti atoms) [Rie76]. Upon cooling below $T_c \approx 200$ K, 1T-TiSe$_2$ undergoes a second-order phase transition into a commensurate CDW. The space group changes to $P\bar{3}c1$. Charge ordering is accompanied by the formation of a periodic lattice distortion corresponding to a structural $(2 \times 2 \times 2)$ superlattice [DS76, Hol77]. The phase transition doubles the size of the unit cell in all three spatial dimensions and, equivalently, halves the Brillouin zone (Figure 3.2). Figure 3.1(b) shows the unit cell in the CDW phase. The PLD is visualized by blue arrows showing the direction of displace-
Chapter 3. The charge density wave in 1T-TiSe$_2$

Figure 3.2: Brillouin zone of 1T-TiSe$_2$ in the normal phase (red) and the CDW phase (light blue). The CDW wavevectors connecting the $\Gamma$ and $L$ points of the normal phase Brillouin zone are shown as blue arrows.

ment of the respective atoms (not true to scale). Figure 3.1(c) shows a top view of one Se-Ti-Se layer to highlight the displacement pattern. At a temperature of 77 K, the displacement amplitude of the Ti and Se atoms amounts to 0.04 Å and 0.014 Å, respectively [DS76]. The deformation pattern leads to a distortion of the Se octahedron (blue body in (b)) and implies three simultaneously present CDWs [Hol77]. In reciprocal space, the corresponding wavevectors $q_{\text{CDW}}$ (blue arrows in Figure 3.2) connect the $\Gamma$ and the three $L$ points of the normal phase Brillouin zone. In the CDW phase, these high symmetry points are mapped on top of each other via the CDW wavevectors and form equivalent zone centers $\Gamma'$ for the smaller CDW phase Brillouin zone (blue wire frame in Figure 3.2).

1T-TiSe$_2$ is a semimetal in both the normal and the CDW phase [DS76, Li07b]. Figure 3.3(a) illustrates the schematic band structure for $T > T_c$ in the vicinity of the high symmetry points related to the phase transition. At the $\Gamma$ point a valence band of Se4p character reaches slightly above $E_F$, leading to the formation of a hole pocket. Electron pockets at the three $L$ points originate from partially filled conduction bands of Ti3d character [Cer07, Roh11]. The periodic spatial reconstruction due to the CDW maps the bands at the $\Gamma$ and the three $L$ points on top of each other [Cer07, Roh11] and leads to the partial opening of an electronic energy gap, as illustrated in Figure 3.3(b) [Wez10b].

3.2.2. Possible driving mechanisms

On the search for the driving mechanism of the CDW phase transition various techniques have been employed. Superlattice peaks in neutron [DS76] or X-ray [Hol01, Web11, MV11] diffraction experiments allow one to identify the lattice distortion pattern and to study CDW induced changes in the phonon dispersion. Optical lattice resonances associated with the PLD can be observed with Raman scattering techniques [Sug80, Sno03] and, as explained in more detail below, via the THz optical response [Hol77]. The electronic degree of freedom was experimentally studied via DC transport measurements [Wil78b], the THz optical response [Wil78a, Lia79, Li07b] and time resolved [Roh11, Hel12] ARPES experiments [Pil00, Kid02, Ros02b, Cer07]. Evidence for both excitonic and phononic
3.2. The CDW phase transition

Figure 3.3: Schematic electronic band structure of 1T-TiSe$_2$ around the Fermi energy $E_F$ at a temperature above (a) and below (b) $T_c$. Back-folding of the three symmetry equivalent Ti$3d$-like conduction bands (represented by an orange curve in (a)) from the L points by the CDW wavevectors $q_{CDW}$ onto the zone centered Se$4p$-like valence bands (dark red curve in (a)) yields a gapped band structure (blue curves in (b)). $\Delta_E$ in (a) represents the small ‘negative’ band gap in the normal phase. Reconstruction of the band structure in the CDW phase as seen in (b) enables a direct transition from the valence to the conduction band across $\Delta_{CDW}$. Illustration adopted from [Mon10].

Contributions to the CDW phase transition was obtained in these experiments, leaving a controversial picture. Fermi surface nesting [Wha91, Joh08] was ruled out due to the lack of parallel Fermi surface contours between electron and hole pockets [Ros02b, Kid02]. In addition the latter mechanism usually leads to incommensurate CDWs. Ultimately, the following three scenarios remained under discussion:

(I) A band-type Jahn-Teller-like structural instability

The first model assumes electron-phonon coupling as the driving force and is based on a band-type Jahn-Teller effect [Hug77, Wha92]. This mechanism may be understood as the 3D analogue of the 1D Peierls instability: For an octahedral configuration of bound atoms, the Jahn-Teller effect describes the removal of degeneracy of electronic levels induced by a distortion of the octahedral structure. In 1T-TiSe$_2$, shortening of the Ti-Se bonds in the CDW phase [Wha92] is supposed to cause a Jahn-Teller-like energetic lowering of Se$4p$ hole-like bonding bands [Ros02b] that stabilizes the distorted octahedral configuration. Experimentally, the important role of electron-lattice interaction in the phase transition is evidenced by the existence of a Kohn anomaly, i.e. a CDW related soft phonon mode [Hol01, Web11]. One can consider the PLD as superposition of three statically excited (“frozen”) $L_T^{-}$ phonon modes with $k = q_{CDW}$ [Hol01]. The static amplitude of these modes and the electron-phonon coupling strength thus determine the amount of Jahn-Teller-like energetic lowering of the Se$4p$ bands. Above $T_c$, static deformation of the lattice costs more energy than is gained from the concomitant lowering of the Se$4p$ bands. Upon cooling
the system towards \( T_c \), reduced thermal fluctuations allow for a more efficient electronic energy gain from population of the PLD constituent phonons. As a consequence, the overall energy needed to induce the PLD is reduced. Equivalently, the eigenfrequency of coupled oscillations of the PLD constituent phonons and the Se4p band energy (i.e. the soft phonon mode) shifts red. This coupled mode ultimately softens to zero frequency at \( T = T_c \) [Hol01]. Below \( T_c \), a static deformation of the lattice is energetically more favorable. The former \( L_1^-- \) modes are now transformed into an \( A_{1g} \) CDW amplitude mode that describes the same deformation pattern but around the PLD equilibrium position. This mode stiffens when the temperature is further decreased [Bar08].

(II) The excitonic condensate scenario

This model assumes that the CDW transition is purely electronically driven [Pil00]. ARPES studies [Cer07, Mon10] revealed a remarkably strong back-folding of the electronic bands in the CDW phase. In addition, the opening of an electronic energy gap was observed. These findings could not be explained within the framework of a Jahn-Teller effect and were seen as evidence for spontaneous formation of bound electron-hole states (i.e. excitons) between the electron- and hole-like Fermi pockets. This excitonic instability is thought to drive a transition to a charge ordered state with a periodicity given by the spanning vectors connecting the valence band maximum at the \( \Gamma \) point with the conduction band minima at the three \( L \) points. The observations made in the ARPES experiments could be well reproduced by a theoretical model that accounts only for the presence of excitons [Mon09]. The PLD is considered as secondary effect induced via exciton-phonon coupling [Mon11]. However, the amplitude of the PLD and the soft phonon mode are not reproduced by this model. The term “condensate” does not refer to a macroscopic phase coherence of the excitons, but is used to account for the presence of free carriers [Mon10] that distinguishes the system from an excitonic insulator [Jér67].

(III) Combination of electron-phonon coupling and excitonic correlations

A third scenario proposes a combination of the above mechanisms. Van Wezel et al. [Wez10a, Wez10b] introduced a 1D model accounting for both a Jahn-Teller-like structural instability and exciton-like electron-hole correlations. This model directly captures the occurrence of the PLD and should in principle explain the observations made in the ARPES experiments [Wez10a]. In section 3.7, it is tested whether this model correctly reproduces the observations made in this work. A more detailed 3D model [Zen13] follows the same approach and additionally captures the chiral properties of the CDW. The latter is associated with different ordering amplitudes along the three CDW wavevectors.
3.2. The CDW phase transition

Figure 3.4: (a) Mid-infrared energy loss function \(-\text{Im}(1/\varepsilon)\) in the normal (\(T = 300\ \text{K}, \text{red curve}) and the CDW phase (\(T = 10\ \text{K}, \text{blue curve}) according to the model function by Li et al. [Li07b]. The curve for \(T = 10\ \text{K}\) results from a numerical adaption to the infrared reflectance of the sample under study (see Figure 3.7). (b) Corresponding curves of the real part of the optical conductivity \(\text{Re}(\sigma)\) in the same mid-infrared window.

3.2.3. The infrared fingerprints of charge and lattice ordering

The transition to the CDW phase modifies the THz optical response in three distinct ways, enabling one to separately study the electronic and lattice degrees of freedom:

(i) Electronic order manifests itself most accessibly as a dramatic modification of the free carrier plasma response. The physical eigenfrequency of collective plasma oscillations (i.e. the screened plasma frequency) is marked by a zero crossing of the real part of the dielectric function \(\varepsilon\). For a descriptive study of the plasma response, the energy loss function \(-\text{Im}(1/\varepsilon)\) is much better suited [Hub01, Li07b], since a zero crossing of \(\varepsilon\) emerges as a characteristic pole in \(-\text{Im}(1/\varepsilon)\). In the normal phase of 1T-TiSe\(_2\), collective plasma oscillations of free electrons and holes (compare Figure 3.3(a)) cause a broad plasmon pole in \(-\text{Im}(1/\varepsilon)\) at a central energy of \(\hbar \omega_{p,\text{norm}} = 145\ \text{meV}\) (Figure 3.4(a), red curve). The effective combined free carrier concentration \(n\) (see section 3.3.2) amounts to \(n_{\text{norm}} = 7.1 \times 10^{20}\ \text{cm}^{-3}\) [Li07b]. The spectral width of the plasmon pole is set by the free carrier scattering time \(\tau_{\text{norm}} = 40\ \text{fs}\) [Li07b]. Below \(T_c\), \(n\) strongly decreases due to renormalization of the electronic band structure (see Figure 3.3(b)), leaving behind a dilute electron gas with \(n_{\text{CDW}} = 9 \times 10^{19}\ \text{cm}^{-3}\). As a consequence, the screened plasmon pole shifts to an energy of \(\hbar \omega_{p,\text{CDW}} = 45\ \text{meV}\) (Figure 3.4(a), blue curve). The reduced phase space for scattering in the ordered state increases \(\tau\) to \(\tau_{\text{CDW}} = 0.9\ \text{ps}\), causing a dramatic narrowing of the plasmon pole to a linewidth of 7\ meV (FWHM).

(ii) The mapping of the \(L\) point onto the \(\Gamma\) point in the CDW phase enables a direct interband single-particle transition from filled Se4p to empty Ti3d derived bands across
Chapter 3. The charge density wave in 1T-TiSe₂

the CDW gap. While a collective plasmon mode is studied most accessibly via \(-\text{Im}(1/\varepsilon)\), a single-particle resonance appears most prominently as maximum in the absorption coefficient \(\alpha\) and in the real part of the optical conductivity \(\text{Re}(\sigma)\). As reported by Li et al. [Li07b] the transition across the CDW gap at \(T = 10\) K prominently manifests itself in \(\text{Re}(\sigma)\) as a broad maximum centered at an energy of 0.4 eV (Figure 3.4(b), blue curve). Importantly, the low-energy spectral wing extends to photon energies down to 0.1 eV. At \(T = 300\) K, this single-particle resonance is absent (Figure 3.4(b), red curve). Note here that in \(-\text{Im}(1/\varepsilon)\) the single-particle transition across the CDW gap emerges only as a relatively weak shoulder at an energy of 0.55 eV. Conversely, the sharp plasmon pole that dominates \(-\text{Im}(1/\varepsilon)\) at a \(T = 10\) K exhibits no tangible spectral feature in the corresponding spectrum of \(\text{Re}(\sigma)\). Only at higher temperatures, the conductivity of free carriers yields an increase of \(\text{Re}(\sigma)\) for photon energies up to \(\hbar\omega_{p,\text{norm}}\).

(iii) The PLD affects the far-infrared response by inducing modifications of the phonon spectrum, as seen best via the imaginary part of the dielectric response function \(\text{Im}(\varepsilon)\). Figure 3.5 depicts \(\text{Im}(\varepsilon)\) in a far-infrared spectral window measured at various temperatures. Above \(T_c\), a single transverse optical phonon resonance at 17 meV is observed (red curves). The mode of \(E_u\) symmetry involves an opposing in-plane motion of the Ti atom versus two Se atoms [Hol77]. Below \(T_c\), PLD induced back-folding of the uppermost acoustic branch of \(B_u\) symmetry from the Brillouin zone boundary to the \(\Gamma\) point [Hol77] yields the additional IR-active in-plane mode at 19 meV (indicated by black arrows). The weaker peak at 22 meV (gray arrows) most likely originates from a folded optical branch [Hol77]. The absence of both PLD induced modes in the normal phase is highlighted by red crosses. The far-infrared dielectric response of 1T-TiSe₂ in thermal equilibrium shown in Figure 3.5 was measured via the complex valued THz field transmission through an 80 nm thin film of stoichiometric 1T-TiSe₂ (see section 3.3.1).

Figure 3.5:
Imaginary part of the far-infrared in-plane dielectric function of 1T-TiSe₂ measured with THz time-domain spectroscopy. At all temperatures, an IR-active \(E_u\) transverse optical phonon mode at an energy of \(\approx 17\) meV is visible. In the CDW phase, at temperatures \(T < T_c\), (blue curves), additional resonances due to back folded phonon branches (indicated by black and gray arrows) attest to the presence of the PLD. For \(T \gtrsim T_c\) (red curves), the PLD induced peaks are absent (red crosses).
3.3. Paving the way for NIR-pump/THz-probe studies

3.3.1. Fabrication and properties of a thin film sample

The single-crystal of 1T-TiSe$_2$ under study is grown by iodine vapor transport$^1$. Excellent stoichiometry of the sample (i.e. the absence of significant Ti-excess) is confirmed by its temperature dependent DC in-plane resistivity [DS76] (see also section 3.4.3) and the narrow plasmon pole observed at $T = 10 \text{ K}$ (compare the corresponding curves in Figures 3.4(a) and 3.7 with the data presented in [Li07b] and [DS76]).

Spectroscopy of a non-transparent bulk crystal in thermal equilibrium is readily performed in reflection geometry. In contrast, THz probing of a photoexcited bulk sample of 1T-TiSe$_2$ in reflection geometry would imply several issues: Since the penetration depth of the pump photons is much smaller than that of the THz photons, extraction of the dielectric function of the photoexcited sample from the pump-induced changes in the THz polarization response would have to account for an inhomogeneous excitation profile. Furthermore, due to the semi-metallic character of 1T-TiSe$_2$, THz photons with energies above and below the plasma edge have dramatically different penetration depths and thus would effectively probe different excitation profiles. These complications can be avoided by probing a thin film in transmission geometry that is sufficiently transparent for all relevant photon energies.

$^1$Bulk samples have been provided by K. Rossnagel, Institute of Experimental and Applied Physics, University of Kiel, 24098 Kiel, Germany.
To this end, an 80 nm thin film of 1T-TiSe$_2$ with a lateral size of 0.2 mm × 0.2 mm is prepared. As a first step, a thin layer of the bulk sample is mechanically exfoliated onto commercially available adhesive tape. Repeated cleaving of the layer eventually results in thin, optically transparent films affixed to the tape. A selected film is then contacted to a CVD-grown diamond substrate by van der Waals bonding. Finally, the adhesive tape is dissolved with methanol. The single-crystal thin film sample of 1T-TiSe$_2$ on diamond substrate used for this work is depicted in Figure 3.6.

The film thickness of 80 nm is measured via optical transmission at a probe wavelength of 532 nm. The sample geometry and the refractive index at the probe wavelength [Bus93] are thereby taken into account via a transfer matrix formalism. Atomic force microscopy is employed to verify the optically measured thickness.

In the following, the NIR photoexcitation intensity is stated as absorbed pump fluence $\Phi$. For the thickness of the studied sample, 1 $\mu$J/cm$^2$ of incident pump fluence corresponds to an absorbed fluence of 0.44 $\mu$J/cm$^2$ (calculated via a transfer matrix formalism) and an average energy deposition per room temperature unit cell (UC) of $2.4 \times 10^{-5}$ eV/UC.

According to a finite difference frequency domain simulation (see chapter 5) that takes into account the refractive index at the pump wavelength [Bus93], the excitation density in the vicinity of the exit surface of the pump beam (i.e. the sample/diamond interface) is still 60% of the average excitation density throughout the sample. To ensure a homogeneous lateral excitation profile, the diameter of the pump focus is kept at twice the size of the THz spot.

### 3.3.2. Quantitative description of the plasma response

The screened plasma frequency $\omega_p$ and the free carrier scattering time $\tau$ provide a measure of electronic order, as discussed in section 3.2.3. To quantitatively link the above variables to the spectral shape of the plasma response, the following parameterization of the dielectric function introduced by Li et al. [Li07b] is employed:

$$\varepsilon(\omega) = \varepsilon_\infty - \frac{\omega_p'}{\omega^2 + \frac{i\omega}{\tau}} + \sum_{j=1}^{2} \frac{S_j^2}{\omega_j^2 - \omega^2 - \frac{i\omega}{\tau}} \tag{3.1}$$

The three terms on the right hand side describe the background dielectric constant $\varepsilon_\infty$, the Drude response of free carriers (second term, $\omega_p'$: unscreened plasma frequency) and the two energetically lowest interband resonances parametrized by a sum of two Lorentzian oscillators (third term). The interband resonance with the lower frequency accounts for the single-particle transition across the CDW gap centered at $\omega_{\text{gap}} = \omega_1$ with a spectral width

$^{2}$Best results with tesafilm®, ord. no. 57370-02, tesa SE.
3.3. Paving the way for NIR-pump/THz-probe studies

![Figure 3.7](image)

**Figure 3.7:** In-plane THz field reflection of $1T$-TiSe$_2$ at $T = 10$ K for s-polarization and an angle of incidence of 30°. Black line: Calculated spectrum based on the model function and parameters by Li *et al.* [Li07b]. Blue dots: Experimental data of the sample under study. Blue line: Model function after numerical adaption of the plasma frequency and the free carrier scattering time.

The free carrier density $n$ is inferred via

$$n = \frac{\omega_p^2 \varepsilon_0 \varepsilon_r m^*}{e^2}$$

employing the relation between screened and unscreened plasma frequency $\omega_p = \omega'_p / \sqrt{\varepsilon_r}$, using a relative permittivity of $\varepsilon_r = 60$ (extracted from [Li07b]) and assuming an effective free electron mass $m^* = m_e$ [Li07b].

For acquisition of the transient dielectric function $\varepsilon(\hbar \omega, t_D)$ as described in section 2.2.3, the model function is employed to describe the dielectric response of the unexcited sample for photon energies above the phonon reststrahlen band. Since most parameters of equation (3.1) are not explicitly stated by Li *et al.* in reference [Li07b], the model parameters are acquired by re-fitting the data presented in the reference. To account for the possibility of a slightly different equilibrium plasma response of the sample under study, its plasma edge in the reflectivity spectrum at a temperature of $T = 10$ K is measured (blue dots in Figure 3.7). To this end, the amplitude spectrum of a single-cycle transient reflected off a bulk sample is referenced against the amplitude spectrum of a transient reflected off a gold coated region of the sample surface. $\omega_p$ and $\tau$ are then numerically adapted so that the model function reproduces the measured reflectivity spectrum for photon energies from 40 meV to 85 meV (blue curve in Figure 3.7).
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Figure 3.8: 2D-maps of the transient energy loss function $-\text{Im}(1/\varepsilon)$ as a function of photon energy and pump-probe delay time $t_D$ following NIR photoexcitation at a temperature of $T = 10 \, \text{K}$. The figure depicts the situation for two representative excitation intensities: The dataset shown in (a) is acquired with an absorbed pump fluence of $\Phi = 20 \, \mu \text{J/cm}^2$ while (b) is measured with more intense photoexcitation with $\Phi = 0.33 \, \text{mJ/cm}^2$. White dashed lines are guides to the eye tracing the center of the plasmon pole.

3.4. The electronic degree of freedom upon photoexcitation

As elaborated in section 3.2.3, charge and lattice ordering of the CDW in $1T$-TiSe$_2$ manifest themselves as characteristic features in the THz dielectric response. This work harnesses these signatures for NIR-pump/THz-probe studies to separately monitor the femtosecond dynamics of electron and lattice order parameters after a controlled perturbation of the CDW ordered phase.

A first series of experiments targets the electronic degree of CDW ordering. It is analyzed on the femtosecond scale how the electronic order recovers from a perturbation induced by a NIR pump pulse and under which excitation conditions the electronic order is fully quenched. Ultrabroadband single-cycle transients (see Figure 2.8) are employed to probe the momentary dielectric response after photoexcitation. The temporal resolution of the experiments of 40 fs is set by the bandwidth of the electro-optic detector (see section 2.2.2). A reduced repetition rate of 400 kHz of the laser system ensures a complete thermalization of the sample with the diamond substrate between two laser shots.
3.4. The electronic degree of freedom upon photoexcitation

Figure 3.9: (a) Transient change in the free carrier density $\Delta n$ relative to $n_{CDW}$ and (b) transient free carrier scattering time $\tau$. The curves are extracted from the complex transient energy loss function acquired for a series of pump fluences and a temperature of the sample of $T = 10$ K. The curve in (a) showing the change in $n$ measured with $\Phi = 7 \mu J/cm^2$ is upscaled by a factor of 120 for quantitative comparison with the situation at $\Phi = 170 \mu J/cm^2$. The gray shaded time window in (a) highlights the rise time of $\Delta n$ of 100 fs (from 20% to 80% of the maximum change).

3.4.1. Non-thermal recovery dynamics of the free carrier response

For photoexcitation of the charge ordered phase, the sample is kept at a temperature of $T = 10$ K. Figure 3.8(a) shows the measured spectra of $-\text{Im}(1/\varepsilon)$ as a function of $t_D$ for an absorbed pump fluence of $\Phi = 20 \mu J/cm^2$ (the raw data of this experiment are shown in appendix A). At negative pump-probe delay times, the system is in thermal equilibrium characterized by the sharp plasmon pole at $\hbar \omega_{p,CDW} = 45$ meV. Upon photoexcitation, this peak rapidly blue-shifts and subsequently recovers within $1.5$ ps. The pole remains narrow during the entire dynamics, indicating that the electronic order remains largely intact while photogenerated free electron-hole pairs elevate the plasma frequency. An increased fluence of $\Phi = 0.33$ mJ/cm$^2$ induces a qualitatively different dynamics (Figure 3.8(b)). The plasmon pole shifts above the measurement window and stays strongly broadened during the subsequent relaxation. As discussed in more detail below, these signatures reflect the complete destruction of the electronic order. After a delay of $t_D \approx 2.5$ ps, a sharp plasmon pole re-emerges at its equilibrium position.

The model dielectric function denoted in equation (3.1) is also an excellent description of the non-equilibrium system. This enables a quantitative analysis of the photoinduced free carrier plasma response: For extraction of the transient plasma frequency $\omega_p(t_D)$ and the transient free carrier scattering time $\tau(t_D)$, the model function is adapted to the measured
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![Figure 3.10: Plasma frequency $\omega_p$ and scattering time $\tau$ plotted as phase space coordinates of the free carrier system. Black spheres in (a) and (b) show combinations of $\omega_p$ and $\tau$ occurring in thermal equilibrium at the indicated temperatures. Transient states during the relaxation of the electronic system after photoexcitation are color coded with the pump-probe delay time $t_D$. Excitation with (a) $\Phi = 20 \mu J/cm^2$ yields non-thermal states, while the free carrier response traverses through thermally accessible states after excitation with (b) $\Phi = 0.33 mJ/cm^2$.](image)

Transient dielectric response via both the real and imaginary parts of $1/\varepsilon(\hbar \omega, t_D)$ in an energy window from 40 meV to 100 meV (see appendix B for the quality of the fit curves). The transient free carrier density $n(t_D)$ is inferred from $\omega_p(t_D)$ via equation (3.2).

The transient change of the free carrier density $\Delta n(t_D) = n(t_D) - n_{norm}$ extracted from $1/\varepsilon(\hbar \omega, t_D)$ is shown in Figure 3.9(a) for a series of pump fluences measured at a sample temperature of $T = 10$ K. After photoexcitation, $\Delta n$ rises within 100 fs, as highlighted by the gray shaded time window. This delayed onset may be attributed to a cascaded relaxation of photoexcited charge carriers with high excess energy that leads to a measurably delayed generation of secondary free charge carriers. An exponential decay with a time constant of 0.5 ps follows. This dynamics of $\Delta n$ is virtually identical for all fluences tested. In contrast, as seen in Figure 3.9(b), the temporal evolution of $\tau$ critically depends on $\Phi$. Starting at $\tau_{CDW} = 0.9$ ps, $\tau$ drops rapidly after photoexcitation and recovers on
3.4. The electronic degree of freedom upon photoexcitation

a fluence-dependent ps timescale (a quantitative discussion follows in section 3.4.2). For $\Phi \leq 20 \mu J/cm^2$, $\tau$ remains distinctly above $\tau_{\text{norm}}$ for all delay times, indicating that the electronic order is not completely destroyed. In the case of $\Phi \geq 60 \mu J/cm^2$, the system temporarily reaches normal state conductivity with $\tau = 40$ fs and remains there for a duration of up to $\sim 2.5$ ps, depending on $\Phi$.

It is instructive to compare the photoinduced plasma response with the situation found in thermal equilibrium. In the latter, $\tau$ and the screened plasma frequency $\omega_p$ form a characteristic pair of values for a given temperature. Selected thermal combinations of $\omega_p$ and $\tau$ are shown as black spheres in the $\omega_p-\tau$ diagrams of Figure 3.10 (value pairs for $T \geq T_c$ are taken from [Li07b]). This way of visualizing the parametrization of the plasma response allows one to readily identify non-thermal combinations of $\omega_p$ and $\tau$ that may be observed after photoexcitation of the system. Figure 3.10(a) depicts the value pairs of $\omega_p$ and $\tau$ observed after photoexcitation of the CDW phase with $\Phi = 20 \mu J/cm^2$ color coded with the pump probe delay time in the range of $0.2 \text{ ps} < t_D < 4.9 \text{ ps}$. The characteristic trajectory obtained this way clearly reveals that non-thermal combinations of $\omega_p$ and $\tau$ occur during the entire relaxation process. $\tau$ remains elevated indicating a persisting lack of scattering channels due to the partial survival of the electronic order. At the same time, optically generated free electron-hole pairs elevate the plasma frequency. In contrast, as seen in Figure 3.10(b), strong excitation with $\Phi = 0.33 \text{ mJ/cm}^2$ fully quenches the electronic order as indicated by a normal phase plasma response. The relaxation of the free carrier system occurs only through thermally accessible states including $T > T_c$.

At the highest measured delay time of $t_D = 100$ ps, the plasma response has settled to a state found at an equilibrium temperature of $T = 108$ K (deduced by interpolation of $\tau$ as a function of temperature). At this stage, the electronic system and the crystal lattice have fully thermalized and the whole system remains heated due to the optical deposition of energy. The sample thermalizes with the diamond substrate to $T = 10$ K before the next pump pulse arrives.

3.4.2. Signatures of excitonic order

The analysis presented above mainly focuses on two regimes of photoexcitation of the CDW phase: A slight optical excitation that leaves the electronic order partly intact and a comparably strong excitation that fully quenches the electronic order for several picoseconds. To study the role of the excitation density on the suppression of the electronic order in greater detail, $n$ and $\tau$ are measured as a function of $\Phi$ (Figure 3.11). The delay

\[ \text{This is in excellent agreement with the temperature of 99 K expected as a result of optical energy deposition [Cra78] by the NIR pump pulse, proving the consistency of the analysis.} \]
Figure 3.11: (a) Free carrier density \( n \) and (b) free carrier scattering time \( \tau \) as a function of the pump fluence measured at a pump-probe delay time of \( t_D = 0.2 \) ps at a temperature of \( T = 10 \) K. Vertical shaded lines indicate the threshold fluence \( \Phi_{th} = 40 \mu J/cm^2 \) required for transient suppression of the excitonic correlations. Horizontal dotted line in (a): Critical free carrier density \( n_c = 4 \times 10^{20} \) cm\(^{-3}\). Gray dashed line: Linear regression of \( n(\Phi) \) in the high fluence regime (\( \Phi > 0.14 \) mJ/cm\(^2\)). The horizontal dotted line in (b) marks the normal phase free carrier scattering time.

time is fixed at \( t_D = 0.2 \) ps such that the measured pump-induced changes have reached their maximum levels (compare Figure 3.9).

When exciting the system in the CDW phase (\( T = 10 \) K), \( n \) first grows superlinearly with \( \Phi \), then changes its curvature at a threshold fluence \( \Phi_{th} \approx 40 \mu J/cm^2 \), and finally approaches a linear dependence for \( \Phi > 140 \mu J/cm^2 \) (Figure 3.11(a)). Simultaneously, \( \tau \) decreases with \( \Phi \), reaches the normal phase limit, and then levels off for the highest pump fluences (Figure 3.11(b)). The fluence required to switch to normal state conductivity coincides with the inflection point \( \Phi_{th} \) of the function \( n(\Phi) \). Excitation below \( \Phi_{th} \) leaves the electronic order partly intact, since \( \tau \) remains above its normal state value. As studied already for a representative pump fluence in section 3.4.1, the electronic system in this excitation regime cannot be described by a thermal states.

Figure 3.12 shows the results of an analogous study starting from the normal state (\( T = 205 \) K). The superlinear increase of \( n \) is not present in this case (purple dots) and \( \tau \) remains at its normal phase value for all fluences (cyan squares).

The observed curve of \( n(\Phi) \) for \( T = 10 \) K is consistent with the picture of charge-transfer excitonic correlations: The NIR pump generates primary electron-hole pairs with densities
3.4. The electronic degree of freedom upon photoexcitation

The electronic degree of freedom upon photoexcitation is crucial for understanding the behavior of charge carriers in materials. When a material is photoexcited, an excess number of charge carriers is created, leading to changes in the material’s electronic properties.

The dependence of the free carrier density \( n \) (purple dots) and scattering time \( \tau \) (cyan squares) on the pump fluence for photoexcitation of the normal phase \((T = 205 \text{ K})\) measured at a pump-probe delay time of \( t_D = 0.2 \text{ ps} \). The horizontal dotted line marks the normal phase free carrier scattering time.

Scaling linearly with \( \Phi \). Their excess energy is relaxed via cascaded electron-electron scattering, which multiplies the quasiparticle density. For low excitation density \((\Phi < \Phi_{th})\), the photogenerated carriers screen electron-hole Coulomb interactions and thus reduce the excitonic binding potential. As a consequence, part of the bound pairs break into free carriers leading to an additional fluence dependent increase of \( n \). This explains the superlinear rise of \( n \). As soon as the photoinduced carrier density exceeds a critical value \( n_c \) for \( \Phi > \Phi_{th} \), excitonically bound electron-hole pairs can no longer exist, \( \tau \) assumes its normal state value (Figure 3.11(b)) and the superlinear scaling of \( n \) stalls (Figure 3.11(a)).

Upon full suppression of all remaining excitonic Coulomb correlations (i.e. electron-hole fluctuations that are also present in the high-temperature phase [Cer07, Mon10]), only the linear dependence of \( n \) on \( \Phi \) remains (dashed curve in Figure 3.11(a)). Consequently, the superlinear increase of \( n \) is absent if the system is excited in the normal phase, since no additional carriers are released from excitonically bound electron-hole pairs.

The idea that Coulomb screening affects the strength of excitonic correlations is widely discussed in literature [Li07b, Mon10]. Here, the effects of photoinduced Coulomb screening are quantitatively analyzed via the Thomas-Fermi screening length \( r_{TF} \). This measure describes the effective range of a Coulomb potential in the presence of free carriers:

\[
r_{TF}(n) = \sqrt{\frac{\pi^2 \hbar^2 \varepsilon_0 \varepsilon_r}{m^* e^2} \left(3\pi^2 n\right)^{-\frac{1}{3}}}
\]

Using the same values for \( \varepsilon_r \) and \( m^* \) as in equation (3.2) and \( n = n_{CDW} \), a screening length of \( r_{TF} = 1.34 \text{ nm} \) is obtained for the system in the CDW phase. This value allows for efficient Coulomb attraction between electrons and holes on a length scale larger than the in-plane and out-of plane CDW wavelength of 1.23 nm and 1.2 nm, respectively. At the critical density of \( n_c \approx 4 \times 10^{20} \text{ cm}^{-3} \), the screening length is reduced to 1.05 nm. Thus, electron-hole attraction on a length scale related to the CDW period should be strongly

Figure 3.12:
Dependence of the free carrier density \( n \) (purple dots) and scattering time \( \tau \) (cyan squares) on the pump fluence for photoexcitation of the normal phase \((T = 205 \text{ K})\) measured at a pump-probe delay time of \( t_D = 0.2 \text{ ps} \). The horizontal dotted line marks the normal phase free carrier scattering time.
suppressed. This is in line with the experimental observation of quenched excitonic order for \( n > n_c \).

An analysis of the \( y \)-intercept of the linear fit of \( n(\Phi) \) for \( \Phi_{th} > 140 \mu J/cm^2 \) underpins the above picture. Extrapolation of the linear dependence to zero fluence yields \( n_{extrap} = 8.5 \times 10^{20} \text{ cm}^{-3} \). This value describes the hypothetical free carrier density after a quench of all electron-hole correlations without the presence of primary photogenerated carriers. If the above picture is valid, this state of the free carrier system should be comparable to a situation in thermal equilibrium when no electron-hole correlations are present, as it is the case for \( T \gtrsim 300 \text{ K} \). Indeed, the values of \( n_{norm} \) observed at \( T = 300 \text{ K} \) and \( n_{extrap} \) are in good agreement.

As mentioned in section 3.4.1, the recovery of \( \tau \) after photoexcitation of the CDW phase occurs on a fluence dependent timescale. Remarkably, the threshold behavior described above manifests itself also in the recovery dynamics of \( \tau \). Figure 3.13 shows the recovery time constant of \( \tau \) as a function of \( \Phi \) acquired by numerical adaption of an exponential decay to the curves shown in Figure 3.9(b). For each value of \( \Phi \), only those sections of \( \tau(t_D) \) are considered, in which \( \tau \) is rising towards its low-temperature value. The time constant \( \tau_r \) thus describes the recovery of the excitonic order once this process has set on. Importantly, \( \tau_r \) reflects the same threshold behavior as already discussed above. A fast recovery is found when the system is excited below \( \Phi_{th} \). In this case, part of the excitonic order remains present and long range order does not have to be newly established from spontaneous fluctuations. Above \( \Phi_{th} \), the recovery time features a plateau at \( \tau_r = 1.5 \text{ ps} \). This value reflects the recovery time constant of the excitonic order after a full quench.

In conclusion, the observed photoinduced modifications of the low-temperature plasma response discussed in this section can be consistently explained by the existence of excitonic charge ordering in 1T-TiSe\(_2\). The next section tests whether this interpretation is compatible with the well known DC resistivity anomaly of 1T-TiSe\(_2\).
3.4. The electronic degree of freedom upon photoexcitation

Figure 3.14: (a) Cyan curve: Equilibrium in-plane DC resistivity\(^4\) of the sample under study as a function of temperature normalized to the value at \(T = 300\) K. Red dots: DC resistivity calculated from the equilibrium values of \(\omega_p\) and \(\tau\) via equation (3.1). The dashed vertical line indicates the inflection point of the DC resistivity at \(T_c\). (b) Ultrafast evolution of the DC resistivity calculated from \(\omega_p(t_D)\) and \(\tau(t_D)\) measured with \(\Phi = 0.33\) mJ/cm\(^2\) at \(T = 10\) K.

3.4.3. The DC resistivity anomaly

The DC resistivity of 1\(T\)-TiSe\(_2\) exhibits an anomalous temperature dependence: Unlike other semimetals or metals, the resistivity does not monotonically decrease upon cooling from room temperature, but instead traverses a pronounced maximum below \(T_c\) [Wil78b, Lia79, Mor06, Kus09] (Figure 3.14(a)). The CDW phase transition occurs at the inflection point on the high temperature side [Wil78b]. Cooling from high temperature to \(T_c\) first increases the resistivity due to a reduction of the effective free carrier concentration. Upon further cooling, the loss of scattering channels [Li07b] due to stabilization of CDW long range order increases the free carrier scattering time and thereby reduces the resistivity. The anomaly vanishes together with the CDW order e.g. upon substitution of Se with S [Wil78b], Cu-intercalation [Mor06] or application of pressure [Kus09]. Here, the transient DC resistivity is extracted from the time resolved THz spectroscopic data by extrapolating the adapted free carrier plasma response to zero frequency [Lia79]. For the equilibrium spectra, this approach is able to qualitatively reproduce the DC resistivity anomaly (see Figure 3.14(a)). Since the optical conductivity after excitation with \(\Phi = 0.33\) mJ/cm\(^2\) mimics thermal states, including \(T > 300\) K, during its relaxation (see Figure 3.10), it immediately follows that the transient DC conductivity traverses the anomalous maximum, as indeed seen in Figure 3.14. At early delay times, the resistivity is reduced due to additional photogenerated electron-hole pairs. At \(t_D \approx 1.1\) ps, the transient resistivity corresponds to an equilibrium resistivity measured at a temperature

\(^4\)Data have been provided by C. Sohrt, Department of Physics, University of Kiel, 24118 Kiel, Germany.
Figure 3.15: Solid lines: Re(σ) in thermal equilibrium reproduced from Figure 3.4(b). Red dots: Re(σ) measured after photoexcitation with Φ = 0.33 mJ/cm² at a delay time of t_D = 1.05 ps and a temperature of T = 10 K. The two data points around a photon energy of 0.22 eV were acquired in a separate experiment using AGS as THz emitter and detector. Dashed line: Hypothetical conductivity of a crossover phase exhibiting a free carrier response as observed at T = 300 K in combination with a CDW gap transition as seen at T = 150 K.

of T = 300 K. The anomalous maximum is traversed at t_D = 2.5 ps. When one considers the DC resistivity anomaly as a fingerprint of excitonic charge order, this time trace can be understood in line with the interpretation of the recovery dynamics of the multi-THz plasma response (see section 3.4.1). Intense photoexcitation with Φ = 0.33 mJ/cm² quenches the excitonic order and thereby induces a DC resistivity characteristic for the normal phase. Recovery of the electronic order occurs through states that can be characterized by a quasi-temperature of the electronic system (compare Figure 3.10). The DC resistivity reflects this trend by traversing the anomalous maximum.

3.4.4. Stability of the CDW gap

As seen above, intense photoexcitation of the CDW phase at T = 10 K with a fluence of Φ = 0.33 mJ/cm² temporarily induces a free carrier response characteristic of the normal phase. To test whether this observation indicates a photoinduced transition to the normal phase of the system, it is analyzed if the single-particle transition across the CDW gap is suppressed under the same excitation conditions.

To this end, the effects of optical pumping and thermal heating on the single-particle CDW gap transition are compared via the real part of the optical conductivity Re(σ). Figure 3.15 reproduces the conductivity spectra for normal (red curve) and CDW phase (blue curve) from Figure 3.4(b). In addition, it shows the conductivity measured after
optical excitation of the CDW phase at $T = 10 \text{ K}$ with $\Phi = 0.33 \text{ mJ/cm}^2$ (dark red data points). The delay time of $t_D = 1.05 \text{ ps}$ is chosen such that the free carrier conductivity corresponds to a temperature of $T = 300 \text{ K}$ implying a full suppression of excitonic order. Intriguingly, the onset of the interband transition across the CDW gap remains clearly visible in this situation. The transient conductivity spectrum may be compared to a scenario, where optical pumping induces a hypothetical crossover phase (dashed line) that combines a free carrier response as expected at a temperature of $T = 300 \text{ K}$ with an interband transition across the CDW gap characteristic for $T = 150 \text{ K}$. The resulting conductivity spectrum closely resembles the measured data.

This observation gives a strong hint that part of the CDW order must remain present while the free carrier system exhibits an optical response characteristic for the normal phase. Thus, electronic correlations cannot be the sole cause for the band structure folding that enables the interband transition. To further corroborate this conclusion, it is analyzed how the lattice degree of freedom behaves under the same excitation conditions.

### 3.5. The periodic lattice deformation upon photoexcitation

While the plasmon pole witnesses the electronic order and the single-particle transition across the CDW gap is indicative for a folding of the band structure, the polarizability of back-folded phonon branches enables one to simultaneously follow the PLD [Hol77].

Figure 3.16 summarizes the phonon response of the photoexcited sample for $T = 10 \text{ K}$ and $\Phi = 0.33 \text{ mJ/cm}^2$, i.e. after complete quenching of excitonic correlations. Surprisingly, the PLD induced mode at $h\nu = 19 \text{ meV}$ (see also Figure 3.5) remains virtually unaffected at all delay times, the phonon resonance originating from the back-folded acoustic branch remains visible (black arrows), proving the persistence of the PLD. The elevated optical conductivity found at $t_D = 1.5 \text{ ps}$ (see Figure 3.8(b)) accounts for the increased background of the corresponding curve.
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times $t_D$. This unexpected observation demonstrates that the PLD remains stable, even though $\Phi$ is sufficiently high to keep the excitonic order molten for several ps (compare Figures 3.9 and 3.10). If excitonic correlations were the main driving force of the PLD, this would be enough time for the lattice to relax into its undistorted state. Yet this is not observed. In order to achieve a melt-down of the PLD with the maximal excitation density available, the sample has to be heated to $T = 150$ K (see Figure 3.17). In contrast, the electronic order is quenched with fluences lower by one order of magnitude, even at much lower temperature.

It is instructive to compare the energy per room-temperature unit cell (UC) required to optically and thermally melt the PLD. Optical excitation with $\Phi = 0.33 \text{ mJ/cm}^2$ corresponds to an energy deposition of $\sim 15 \text{ meV/UC}$. In contrast, it follows from the temperature dependent specific heat [Cra78], that an energy density of $30 \text{ meV/UC}$ is required to induce the phase transition thermally by heating the system from $T = 150$ K to $T = T_c$. This apparent discrepancy has been observed before in time resolved x-ray studies [MV11]. In light of the above results, such behavior may indeed be expected: Quenching of the excitonic order selectively excites the PLD related lattice modes to a temporarily increased quasi-temperature sufficient to melt the PLD. In addition, due to the suppressed excitonic contribution, the critical temperature of the remaining component of the CDW order may be temporarily lowered.

The far-infrared equilibrium dielectric response required for extraction of the transient phonon spectra is measured via the complex valued THz field transmission through the sample. The THz probe transients are generated and electro-optically detected using GaP as THz emitter and detector (see section 2.2.1). The time resolution in this configuration permits to trace photoinduced changes in the dielectric function occurring on a timescale of shorter than 1.5 ps (compare Figure 3.16).

![Image](image.png)

**Figure 3.17:**
Far-infrared spectra of $\text{Im}(\varepsilon)$ after NIR excitation with $\Phi = 0.33 \text{ mJ/cm}^2$ measured at $T = 150$ K. The suppression of the PLD-related phonon (red crosses) for $t_D = 0$ ps and 3 ps attests to a complete melting of the PLD within the corresponding time window.
3.6. Coupling of lattice and electronic orders

Both the remaining PLD and the CDW gap demonstrate that the CDW phase can exist without excitonic order. Nonetheless, lattice and electronic orders do couple, as seen next.

Optical excitation of the CDW phase at $T = 10$ K induces changes of the collective free carrier response as well as the single-particle excitations while the PLD appears unaffected. In the discussion so far, each degree of freedom is analyzed separately. In this section, the THz absorption coefficient $\alpha$ is studied, since it intuitively captures the absorption associated with both the free carriers and the transition across the CDW gap. Figure 3.18 shows 2D-maps of the absorption coefficient $\alpha$ as a function of the photon energy and the pump-probe delay time for three characteristic excitation densities. At low fluence with $\Phi = 3 \mu J/cm^2$ (Figure 3.18(a)), photoexcitation yields an increase of absorption both on the low and on the high energy side. These dynamics attest to modifications of the plasma response as discussed above and, additionally, to a red-shift of the onset of the interband transition across the CDW gap. The dynamics of $\alpha(h\omega, t_D)$ on the high-frequency side monitoring the single-particle excitation gap is even more interesting for larger $\Phi$ (Figure 3.18(b)): The spectral weight above a photon energy of 100 meV is increased further, indicative of a further reduction of the energy gap. In addition, the dynamics of the CDW gap is visibly superimposed with coherent oscillations at a frequency corresponding to the
A$_{1g}$ CDW amplitude mode (see below). A black contour line highlights this feature. Note that the low-frequency plasma response does not exhibit this coherent dynamics. Upon excitation with $\Phi > \Phi_{th}$ (Figure 3.18(c)) the free carrier absorption overlaps with the single-particle absorption of the CDW gap.

When one considers the CDW in 1T-TiSe$_2$ as the result of concerted action of excitonic correlations and Jahn-Teller-like effects, these observations can be well understood:

Optical excitation breaks the excitonically bound electron-hole pairs and thereby increases the free carrier density and decreases a jointly formed energy gap. Consequently, the plasmon pole shifts higher while the additional spectral weight from single-particle excitations extends towards lower frequencies. Upon approaching the critical excitation fluence, the collective plasma response should effectively screen low-energy transitions associated with the excitonic component of the CDW gap comparable to an excitonic Mott transition [Zim78]. The consequences are apparent in Figure 3.11(a): Excitonic correlations that contribute to the stabilization of the CDW gap are more effectively screened with an increasing density of free carriers $n$. Thus, charge carriers in this situation are more readily released from the bound states and add to the primary photogenerated free carriers, leading to the observed superlinear increase of $n$ with $\Phi$.

The coherent oscillations visible in Figure 3.18 complete this picture. Cuts through the data of Figure 3.18(b) along the time axis $t_D$ at two representative photon energies are shown in Figure 3.19. The coherent amplitude oscillations strongly modulate the mid-infrared absorption at a photon energy of 125 meV (blue curve), while the low-frequency response at 45 meV (red curve) is not affected. When optical excitation quenches part of the excitonic correlations, the equilibrium position of the remaining CDW is suddenly relaxed, such that a coherent oscillation of the A$_{1g}$ CDW amplitude mode around the new equilibrium position is triggered non-adiabatically. This coherent A$_{1g}$ phonon field forces the remaining CDW order periodically out of its momentary energy minimum and thereby modulates the Jahn-Teller component of the CDW gap.

Figure 3.19:
Cuts through the 2D-map $\alpha(h\omega, t_D)$ of Figure 3.18(b) recorded with an excitation fluence of $\Phi = 10 \mu J/cm^2$ at a temperature of $T = 10 K$ along the $t_D$ time axis at photon energies of 45 meV (red line) and 125 meV (blue line).
3.6. Coupling of lattice and electronic orders

Figure 3.20: Oscillatory component $\Delta A_{\text{osc}}(t_D)$ of the pump-induced THz transmission change $\Delta A(t_D)$ recorded at a temperature of $T = 10\,\text{K}$ and various pump fluences. $\Delta A$ results from a 1D pump-probe experiment and refers to an integrated spectral range covering photon energies from 40 meV to 140 meV. Vertical gray lines mark the maxima of the waveform recorded with $\Phi = 3\,\mu\text{J}/\text{cm}^2$. The black vertical line and the black arrow indicate the time difference required for three full oscillation cycles during persistent ($\Phi = 3\,\mu\text{J}/\text{cm}^2$, blue line) and suppressed excitonic order ($\Phi = 67\,\mu\text{J}/\text{cm}^2$, purple line).

With the physical origin of the periodic modulation for the THz response following NIR photoexcitation identified, a spectrally integrated time resolved THz transmission experiment [Kim12] is now performed to measure details of the transient oscillatory response as a function of the excitation intensity. For this study, the electro-optic sampling time $t$ is fixed to measure the maximum field transmission amplitude $A$ while the pump-probe delay time $t_D$ is varied. This procedure captures the spectrally integrated pump-induced transient transmission change through the sample $A(t_D)$. The oscillatory component $\Delta A_{\text{osc}}(t_D)$ superimposed in the dynamics of $\Delta A(t_D)$ shown in Figure 3.20 has been isolated by subtraction of exponentially decaying components from $\Delta A(t_D)$.

$\Delta A_{\text{osc}}(t_D)$ exhibits a frequency of 3.4 THz, for $\Phi = 7\,\mu\text{J}/\text{cm}^2$ (Figure 3.21(a)), which is characteristic of the well-known $A_{1g}$ CDW amplitude mode [Sno03, MV11]. While the oscillation is still observed for excitation densities above $\Phi_{\text{th}}$, its amplitude increases with $\Phi$ only up to $\Phi_{\text{th}}$ (Figure 3.21(b)). This behavior is indeed expected when the CDW gap is jointly formed by excitonic correlations and Jahn-Teller effects. Following impulsive weakening of the excitonic order, the remaining PLD oscillates around a slightly relaxed potential energy minimum. This coherent dynamics modulates the Jahn-Teller component of the CDW gap. Following a complete quench of excitonic correlations, the new equilibrium position of the PLD is stabilized by Jahn-Teller effects and the oscillation

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amplitude, given by the pump-induced shift of the equilibrium position, remains at its maximum level. This interpretation is supported by a theoretical model (see section 3.7) predicting a reduced, but finite PLD when the excitonic order is quenched. The precise traces of the coherent oscillations are recorded at different excitation densities (Figure 3.20). The initial phase of the coherent oscillation is constant for all fluences. Yet, its frequency slightly decreases when excited with $\Phi = 67 \, \mu J/cm^2$ as compared to $\Phi = 3 \, \mu J/cm^2$. The change in frequency eventually leads to a phase retardation of $\pi/2$ after three oscillation periods at $t_D = 1$ ps (black arrow in Figure 3.20). Since $\Phi = 67 \, \mu J/cm^2$ effectively quenches excitonic correlations within this initial time window (compare Figure 3.9(b)), the restoring force in this case is mainly given by a remaining Jahn-Teller mechanism. As discussed in the next section, the pump-induced softening of the $A_{1g}$ amplitude mode suggests a cooperative coupling between excitonic correlations and Jahn-Teller effects.

As a remark it is noticed here, that for low intensity photoexcitation, the center energy of the transition across the CDW gap may be extracted from the transient dielectric response (see appendix C). The observed behavior is in line with the above interpretation.

From the experimental point of view, all key observations can be consistently explained in the picture of combined action of excitonic correlations and a structural Jahn-Teller like effect in the formation of the CDW. In the next section it is tested whether this interpretation of the experimental results is compatible with a recent theoretical model that accounts for both contributions.

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**Figure 3.21:** (a) Frequency spectrum of the coherent oscillations observed with a pump fluence of $\Phi = 7 \, \mu J/cm^2$ (compare Figure 3.20). The position of the single peak at 13.8 meV identifies the well-known $A_{1g}$ CDW amplitude mode [MV11]. (b) Maximum amplitude of $\Delta A_{osc}$ as a function of $\Phi$. The curve flattens at a fluence coinciding with $\Phi_{th}$ (red shaded vertical line).
3.7. The results in the light of a 1D mean-field theory

Extensive theoretical work has aimed for a better understanding of the microscopic mechanisms that lead to the CDW phase transition in 1T-TiSe$_2$ [Mon09, Wez10b, Mon11, Cal11, Zhu12, Zen13]. This section compares the results obtained in this work to the predictions of a mean-field model introduced by van Wezel et al. [Wez10b]. Both Jahn-Teller and excitonic components of the CDW order parameter are considered in this model.

3.7.1. Description of the model

The equilibrium Hamiltonian

$$H = H_0 + H_X + H_{e-l}$$
(3.4)

models two coupled chains of atoms with one orbital per lattice site $i$ as schematically shown in Figure 3.22. The upper chain represents the interacting Se4p orbitals (green shapes), created by $c_i^\dagger$ (valence electrons) and the lower chain represents the Ti3d orbitals (gray shapes), created by $d_i^\dagger$ (conduction electrons). The energetic distance between the two atomic levels is denoted by $\Delta$. The noninteracting tight-binding Hamiltonian $H_0$ describes the band structure close to the Fermi energy, with nearest-neighbor intra-chain hopping of amplitude $t/2$ in both chains and inter-chain nearest-neighbor hopping of amplitude $t'$. $H_0$ also includes local phonon modes in the $c$- and $d$-chains (phonon energy $\hbar \omega_{ph}$), created by $b_i^\dagger$ and $a_i^\dagger$, respectively. The interaction terms $H_X$ and $H_{e-l}$ account for the nearest-neighbor couplings of the atoms of the two chains. Excitonic effects are induced by the screened Coulomb interaction potential $V$ between a $d$-conduction electron and a $c$-valence hole:

$$H_X = -V \sum_i d_i^\dagger d_i \left[ c_i^\dagger c_{i-1}^\dagger + c_{i-1} c_i^\dagger \right].$$
(3.5)

The Jahn-Teller electron-lattice local couplings are described by

$$H_{e-l} = \alpha \sum_i \left[ (X_i^a + X_i^b)(d_i^\dagger c_i + c_i^\dagger d_i) - (X_{i+1}^a + X_i^b)(d_{i+1}^\dagger c_i + c_{i+1}^\dagger d_{i+1}) \right]$$
(3.6)

where $\alpha$ is the electron-phonon coupling constant and

$$X_i^a = \sqrt{\frac{\hbar}{2ma_i^2 \omega_{ph}}} (a_i + a_i^\dagger)$$
(3.7)

is the displacement field operator of the atom $i$ with mass $m$. $a_i$ denotes the lattice constant.
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Figure 3.22: The one-dimensional double chain model introduced by van Wezel et al. [Wez10b] employed to analyze the influence of Jahn-Teller-like and excitonic effects on the CDW formation. Gray and green shapes correspond to Ti3d and selected Se4p orbitals, respectively. $i$ enumerates the lattice sites. $c^\dagger$ and $d^\dagger$ create electrons in the Se4p and Ti3d states, respectively. The $X$ operators describe the displacement amplitude of the corresponding atom.

Van Wezel et al. treat the many-body interactions $H_X$ and $H_{e-1}$ within the Hartree-Fock approximation and employ the Bloch theorem to derive a fully decoupled mean-field Hamiltonian that describes the charge density formation:

$$H_{MF} = \sum_k \left( \epsilon_k c_k^\dagger c_k + \eta_k d_k^\dagger d_k + \beta_k e^{i\phi_k} d_k^\dagger c_k + \beta_k e^{-i\phi_k} c_k^\dagger d_k \right)$$
$$+ \sum_k \hbar \omega_{ph} \left( a_k^\dagger a_k + b_k^\dagger b_k \right) + 2\alpha \tau_\delta \sqrt{N} \left( a_{k=0}^\dagger + a_{k=0} + b_{k=0}^\dagger + b_{k=0} \right)$$

The mean-fields are given by the average electron density per lattice site

$$\rho_c = \langle c_i^\dagger c_i \rangle \quad \rho_d = \langle d_i^\dagger d_i \rangle,$$

the charge transfer within ($\tau_{in}$) and between ($\tau_{out}$) the unit cells

$$\tau_{in} = \langle c_i^\dagger d_i \rangle \quad \tau_{out} = \langle c_i^\dagger d_{i+1} \rangle$$

and the mean-field lattice distortion

$$u = u_a + u_b = \langle X_i^a \rangle + \langle X_i^b \rangle.$$

The energies of the conduction and valence bands, including the excitonic energy contribution when inter-chain electron-hole pairs are present and $\rho_c \neq 1$, $\rho_d \neq 0$ are given by
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\[ \epsilon_k = t \cos(ka_L) - \frac{\Delta}{2} + 2V \rho_d \quad \text{and} \quad \eta_k = t \cos(ka_L) + \frac{\Delta}{2} + 2V(\rho_c - 1) \] (3.12)

The two chains are coupled via the tight-binding band structure described by \( t' \) and, most importantly, via the electron-lattice and excitonic interactions:

\[ \beta_k e^{i\phi_k} = t'(1 + e^{ika_L}) + \alpha u (1 - e^{ika_L}) - V(\tau_{in} + e^{ika_L} \tau_{out}) \] (3.13)

While the band structure contribution \( t' \) dominates at \( k = 0 \), close to the Brillouin zone boundary at \( k = \pi/a_L \), the coupling of the two chains is dominated by both excitonic and electron-lattice interactions characterized by \( \alpha u \) and \( V \tau_\delta \). Here, \( \tau_\delta = \text{Re}(\tau_{in} - \tau_{out}) \) describes the inter-chain electron-hole coherence.

The above mean-field Hamiltonian is diagonalized via a standard Bogoliubov transformation [Bog47] to obtain the quasiparticles (created by \( A_k^\dagger \) and \( B_k^\dagger \)) describing the hybridized excitations:

\[ c_k^\dagger = e^{i\phi_k/2} \left( \cos \theta_k A_k^\dagger - \sin \theta_k B_k^\dagger \right), \quad d_k^\dagger = e^{-i\phi_k/2} \left( \sin \theta_k A_k^\dagger + \cos \theta_k B_k^\dagger \right) \]

where

\[ \tan(2\theta_k) = \frac{2\beta_k}{\epsilon_k - \eta_k}. \] (3.14)

After introducing shifted phonon operators (\( \alpha_k^\dagger \) and \( \gamma_k^\dagger \)) the Hamiltonian can be expressed in diagonal form:

\[ H_{MF} = -\frac{8a_\Delta^2 \tau_\delta^2 N}{\hbar \omega_{ph}} + \sum_k \left( E_k^A A_k^\dagger A_k + E_k^B B_k^\dagger B_k \right) + \sum_k \hbar \omega_{ph} \left( \alpha_k^\dagger \alpha_k + \gamma_k^\dagger \gamma_k \right) \] (3.15)

The energy dispersion of the two quasiparticle branches is given by

\[ E_k^{A,B} = \frac{1}{2} \left[ (\epsilon_k + \eta_k) \pm \text{sgn}(\epsilon_k - \eta_k) \sqrt{(\epsilon_k - \eta_k)^2 + (2\beta_k)^2} \right]. \] (3.16)

In the zero temperature ground state, the lower branch is fully occupied and the higher branch is empty.

The values of the mean-fields are now determined by minimizing the total energy given by \( H_{MF} \) for each combination of the excitonic binding potential \( V \) and the Jahn-Teller coupling strength \( \alpha \) with the mean-fields as free parameters. Thereby, \( \rho_c, \rho_d, \tau_{in} \) and \( \tau_{out} \) describe two coupled qubits and are constrained by translational invariance along the chains and conservation of the particle number (see appendix D).
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3.7.2. The equilibrium phase diagram

The mean-fields $\rho = \rho_c - \rho_d$, $\tau_3$ and $u$ constitute the components of the order parameter of the model system and define the equilibrium phase diagram. Figure 3.23 and Figure 3.24 provide an overview of the different phases described by the model. As the first component of the composite CDW order parameter, the excitonic charge transfer $\rho$ from the valence to the conduction band as a function of $V$ and $\alpha$ is reproduced (Figure 3.23). The normal phase with electrons effectively only populating the valence band is characterized by $\rho \approx 1$ (red shaded region). For a sufficiently high value of $V$, an excitonic insulator is formed when $\alpha$ is sufficiently low. This phase is characterized by $\rho \approx -1$ (blue shaded region) and describes an almost complete charge transfer from the valence to the conduction band corresponding to the existence of excitonically bound electron-hole pairs.

To capture all components of the CDW order parameter, the analysis of van Wezel et al. [Wez10b] is extended by calculation of the PLD mean-field amplitude $u$ for each combination of $V$ and $\alpha$ (Figure 3.24(a)). Without excitonic effects ($V = 0$), a positive $u$ ($\alpha > 0.045 \text{ eV}$) describes a purely Jahn-Teller-like structural phase transition. For nonzero $V$, both cooperation and competition between excitonic and Jahn-Teller effects can occur: In the range $0 \text{ eV} < V < 0.75 \text{ eV}$, an increase of $V$ enhances the PLD and lowers the critical value of $\alpha$ for the structural CDW phase transition. Thus, in this region of $V$ with a nonzero PLD present, excitonic correlations and Jahn-Teller effects cooperate to form a combined Jahn-Teller excitonic CDW. In contrast, upon further increase of $V$,
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Figure 3.24: (a) 2D-map of the mean-field lattice distortion $u$ as a function of $V$ and $\alpha$. For $V > 0$, a finite value of $u$ characterizes a CDW resulting from both Jahn-Teller and excitonic contributions. (b) 2D-map of the CDW amplitude mode frequency $\omega_{am}$ normalized to the uncoupled oscillation frequency of the normal phase. The mode softens to zero frequency at the structural phase transition.

excitonic order and Jahn-Teller effects compete, as apparent from the formation of more excitons at the expense of a reduced PLD (compare Figures 3.23 and 3.24(a)). Ultimately, in the excitonic insulator phase, the PLD is zero since no direct exciton-phonon coupling is considered in the model.

To qualitatively compare the experimentally observed softening of the CDW amplitude mode, the frequency $\omega_{am}$ of the amplitude mode as a function of $V$ and $\alpha$ is calculated by variation of $u$ around its equilibrium value (Figure 3.24(b)). As one would expect for a mean-field transition, a softening to zero frequency occurs at the onset of the structural phase transition. When $u$ becomes nonzero, the mode changes its symmetry from a normal phase lattice mode to the CDW amplitude mode.

3.7.3. Modeling NIR photoexcitation

To compare the model predictions with the experimental data, the components of the order parameter are associated with experimentally accessible quantities. The free carrier density $n$ can be related to $\rho$ since both are directly connected to the exciton density (both $n$ and $\rho$ are reduced/increased by exciton formation/destruction). As discussed in section 3.6, the low-energy spectral wing of the single-particle transition across the CDW is observed in the experiment. In the model, this transition corresponds to the lowest energetic distance between the two quasiparticle branches, occurring at $k = \pi/a_L$. This
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Figure 3.25:
(a) $\rho$ as a function of $V$ for two selected values of $\alpha$. Reduction of $V$ by Coulomb screening due to optically generated free carriers leads to an increase of $\rho$. This holds for both the excitonic insulator ($\alpha = 0$, dashed line) and the coupled Jahn-Teller excitonic CDW ($\alpha = 0.06\,\text{eV}$, dark green line). The blue vertical line indicates a possible value of $V$ before a pump-induced quench to $V = 0$ (red vertical line). (b) Energy gap at $k = \pi/a_L$ as a function of $V$. For a cooperatively coupled CDW ($V \lesssim 0.8\,\text{eV}$), the quench of $V$ reduces the gap. (c) Frequency of the amplitude mode $w_{am}$ as a function of $V$. For an initial value of $V \lesssim 0.8\,\text{eV}$, the quench reduces $w_{am}$. At the structural phase transition from a combined CDW to an excitonic insulator ($V \approx 1.4\,\text{eV}$) the amplitude mode softens to zero frequency and changes its symmetry to a normal phase lattice mode.

energy gap, $\Delta_{k=\pi/a_L}$, also scales with both $V$ and $\alpha$. The frequency of the $A_{1g}$ amplitude mode is directly captured by the model via the PLD amplitude mode frequency $\omega_{am}$.

NIR photoexcitation creates high energy electron-hole pairs with an energy much larger than the CDW gap. These carriers relax by electron-electron scattering, exciting further low-energy quasiparticles via cascading effects. The following analysis assumes that photogenerated free carriers lead to a reduction of the excitonic electron-hole interaction potential by Coulomb screening. This scenario is modeled by a reduction of $V$ while keeping $\alpha$ fixed. The pump-induced change in the exciton energy $V$ may be thought of as an “interaction quench” with a finite duration. A direct change of the charge imbalance $\rho_c - \rho_d$ by excitation of charge carriers that do not directly participate in the formation of the CDW is not considered here.

Even with this simplest possible approach the model qualitatively accounts for all observations made in the experiment. A very general prediction is the release of free carriers induced by the destruction of excitons following a reduction of $V$, independent of the exact value of $\alpha$ (Figure 3.25(a)). In the case of $\alpha = 0$ (dashed line), a step-like increase of $n$ is expected upon optically induced suppression of $V$ below the critical value required to stabilize the excitonic insulator phase. For $\alpha = 0.06\,\text{eV}$ (green line), the increase of $n$ at a reduction of $V$ is more continuous. This aspect enables a qualitative understanding of the
3.7. The results in the light of a 1D mean-field theory

dependence of \( n \) on the excitation fluence \( \Phi \) (Figure 3.11(a)): A linear component in \( n(\Phi) \) accounts for the primary photogenerated free carriers, while the superlinear dependence seen at low fluences is a result of an increasing destruction of excitons. Upon approaching a full quench of \( V \) to zero, the number of additional free carriers released from excitons levels off until ultimately no more excitons can be broken and only the linear dependence remains. Furthermore, the approach correctly reproduces the behavior of the CDW gap. As shown in Figure 3.25(b), in the regime of a cooperative Jahn-Teller excitonic CDW \((V < 0.8\, \text{eV for } \alpha = 0.06\, \text{eV})\), a reduction of \( V \) continuously decreases \( \Delta_{k=\pi/a_L} \) (green line). Notably, even at a full quench of \( V \), indicated in Figure 3.25 by the transition from \( V = 0.8\, \text{eV} \) (blue vertical line) to \( V = 0 \) (red vertical line), the gap remains at an elevated value due to the remaining Jahn-Teller component of the CDW (compare Figure 3.15).

As shown next, with this remaining component of the CDW order, the model also explains the experimentally observed softening of the \( A_{1g} \) amplitude mode after photoexcitation. Figure 3.25(c) shows the frequency of the model CDW amplitude mode \( \omega_{am} \) as a function of \( V \) for \( \alpha = 0.06\, \text{eV} \). Excitonic correlations stiffen the amplitude mode in the regime of a cooperative CDW \((V < 0.8\, \text{eV})\). For higher values of \( V \), the mode softens due to the competition of Jahn-Teller effects and excitonic correlations. Thus, assuming photoinduced reduction of \( V \) in the cooperative CDW regime, the model correctly accounts for the observed softening of the amplitude mode after photoexcitation. Furthermore, the saturation of the oscillation amplitude at \( \Phi_{th} \) (see Figure 3.21(b)) can be identified as another consequence of a quench of \( V \). Since the maximum amplitude of the PLD oscillation is given by the photoinduced shift of the PLD equilibrium position, the increase of the oscillation amplitude stalls when the new equilibrium position, given by a purely Jahn-Teller stabilized CDW, becomes more robust against photoexcitation.

Finally, the coherent excitation of the \( A_{1g} \) amplitude mode can be modeled by forcing \( u \) out of its minimum energy value by \( \Delta u \) (with \( \Delta u \ll u \)). For minimized total energy with constrained \( u \), the relative change of \( \Delta_{k=\pi/a_L} \) is one order of magnitude larger than the relative change of \( \rho \). This is in line with the observation that the energy gap is visibly modulated by the coherent oscillations of the amplitude mode while the free carrier response remains unaffected (compare section 3.6). Thus, it is the Jahn-Teller component of the CDW gap that is modulated by the coherent oscillation of the amplitude mode.

In conclusion, a quench of the excitonic component of a cooperatively coupled excitonic Jahn-Teller CDW, modeled by a quench of the excitonic binding potential, qualitatively explains the key experimental observations. In perspective, the presented data may provide a foundation and benchmark for the development of advanced quantum-kinetic theories. If based on a full 3D model such as introduced recently by Zenker et al. [Zen13], even a quantitative agreement with the results presented in this work may be achieved.
3.8. The origin of the charge density wave in $1T$-TiSe$_2$

In conclusion, the microscopic origin of the lattice commensurate charge density wave in $1T$-TiSe$_2$ is consistently identified in this work by means of femtosecond NIR-pump/multi-THz-probe spectroscopy. Electronic ordering and a reconstruction of the crystal lattice, two coupled components of the CDW order parameter, are separately monitored via their individual THz fingerprints after controlled perturbation of the system with a 12-fs NIR pump pulse. The free carrier plasma response is employed as a measure of the electronic component of charge ordering. The observed photoinduced modifications of the free carrier system attest to an excitonic contribution to charge ordering. In addition, back folded phonon modes are harnessed as a measure of the structural component of the charge density order parameter. It is demonstrated, that photoexcitation can drive the system into a highly non-thermal phase in which the structural component of the charge ordered phase persists in a coherently excited state even when the excitonic order is quenched.

The experimental results strongly indicate, that the joint action of excitonic electron-hole correlations and a structural Jahn-Teller-like effect induces the charge density wave phase transition in $1T$-TiSe$_2$. A quantum-mechanical mean-field model [Wez10b], which is extended to describe the influence of photoexcitation on the excitonic and structural degrees of freedom, strongly corroborates this interpretation of the experimental data.
4.1. Unconventional superconductivity

4.1.1. A short overview

The strength of a purely phonon-based mechanism for the formation of superconducting Cooper-pairs is limited by the phonon frequencies in solids. In terms of the BCS theory, this insight fueled the widespread assumption that a transition temperature for superconductivity of $\gtrsim 30\,\text{K}$ should not be observable [Car03]. This conception was falsified in 1986, when Bednorz and Müller discovered high-temperature superconductivity in cuprates [Bed86]. The researchers expected superconductivity in these materials due to strong electron-phonon coupling indicated by a huge Jahn-Teller distortion. Initially, Bednorz and Müller found an inexplicably high critical temperature of $30\,\text{K}$ in the system
Ba-La-Cu-O. Shortly after, a transition temperature of 93 K was reported for Y-Ba-Cu-O [Wu87]. Currently, the highest transition temperature of 164 K is achieved in Hg-Ba-Ca-Cu-O under pressure [Gao94]. Bednorz and Müller were awarded with the Nobel prize already one year after their discovery. The latest experimental milestone was set in 2008, when Kamihara et al. [Kam08] reported a superconducting phase at 26 K in fluorine-doped La-Fe-As-O. This marked the beginning of worldwide efforts to investigate the new family of iron-based “pnictide” superconductors [Pag10, Dai12, Kim12]. Given the strong magnetism of iron and considering the usually antagonistic relationship between superconductivity and magnetism, the observation of high transition temperatures in these systems was completely unexpected. This again proved, that the fundamental understanding of the origins of unconventional superconductivity needs significant improvement [Pag10].

After almost thirty years of extensive research, the mechanism that leads to the formation of an unconventional superconducting condensate still remains elusive. Besides the relevance of a variety of many-body phenomena, such as spin-, charge- or stripe-ordered phases or the pseudo gap, also the role of electron-phonon coupling in the pairing mechanism remains debated. Some theories assume that high-temperature superconductivity may occur entirely without the influence of lattice vibrations. In these scenarios, the virtual exchange bosons that bind the electrons into Cooper-pairs are sought among purely electronic elementary excitations, such as fluctuating magnons [And07, Mon07] or interorbital pair hopping excitations [Ste11]. A certain role of phonons in the formation of a superconducting condensate in some cuprates is evidenced by various experiments, including ARPES [Lan01, Cuk04, Iwa08], inelastic neutron scattering [Rez06], tunneling [Lee06] and Raman [Ope99] spectroscopic techniques. The presence of an isotope effect on the transition temperature [Bat87a, Bat87b] and strong anomalies of certain phonon modes in cuprates across the phase transition [Ber00] indicate appreciable interaction between the condensate and the lattice vibrations. Time-resolved photoemission results have suggested that there is a subset of phonon modes (about 20% of total number) which exhibit stronger coupling to the electrons [Per07]. As this evidence was indirectly deduced from the evolution of a quasi-equilibrium electronic temperature, the character of the strongly and weakly coupled phonons could not be identified.

Here, femtosecond multi-THz spectroscopy is employed to study the ultrafast interplay of electrons and two specific phonon modes in optimally doped YBa$_2$Cu$_3$O$_{6.93}$. This experiment aims to answer the question of whether there exist separate timescales (i.e. a temporal hierarchy) of the electron-electron and electron-phonon scattering processes in the non-equilibrium system. It is shown, that both scattering processes occur at the same timescale of 150 fs, demonstrating extremely fast electron-phonon scattering. This stands in contrast to the assumptions of two- or three-temperature models [Kab08, Per07].
4.1. Unconventional superconductivity

\[ \text{BaCuO}_a^b\text{cY O}_1 \]

Figure 4.1: Crystal structure of orthorhombic YBa\(_2\)Cu\(_3\)O\(_7\). The black frame contains a single unit cell. The doping level \( \delta \) (here \( \delta = 0 \)) is controlled by the amount of oxygen ions occupying the chains along the \( b \) crystal axis (O1). Cu2 and O3 atoms form the two-dimensional superconducting planes (highlighted by blue surfaces).

4.1.2. Cuprate superconductors and YBa\(_2\)Cu\(_3\)O\(_{7-\delta}\)

Cuprates are the most thoroughly studied class of high temperature superconductors and currently host the compound with the highest observed transition temperature [Gao94]. Cuprates have one or more CuO\(_2\) planes in their structure which are separated by layers of other atoms (Ba, O, La, ...). In the planes, each copper ion is covalently bound to four oxygen ions. Unconventional superconductivity is most likely related to processes occurring in the CuO\(_2\) planes, with the other layers solely forming charge reservoirs. Their chemistry determines the type and the amount of charge carriers in the planes. The critical temperature scales with the number of CuO\(_2\) planes that are within a short distance of each other [Hot13].

Figure 4.1 depicts the unit cell of YBa\(_2\)Cu\(_3\)O\(_{7-\delta}\). The lattice constants are \( a = 3.82\, \text{Å} \), \( b = 3.89\, \text{Å} \) and \( c = 11.68\, \text{Å} \) [Sax09]. O1 – Cu1 chains along the crystal \( b \)-axis constitute the charge reservoir. The superconducting CuO\(_2\) planes (highlighted by blue surfaces in Figure 4.1) are formed by O3 and Cu2. Increasing of the amount of O1 leads to an increase in the effective valence of the plane coppers Cu2 due to a transfer of electrons from the planes to the chains. This is equivalent to a hole-doping of the superconducting planes (denoted by the hole concentration \( \rho \) per Cu2). When all O1 positions are filled (at \( \delta = 0 \), corresponding to \( \rho = 0.21 \)), the hole doping can be further increased by partial substitution of the trivalent Y\(^{3+}\) ions with bivalent ions, e.g. Ca\(^{2+}\).
Chapter 4. Phonon and quasiparticle dynamics in superconducting YBa$_2$Cu$_3$O$_{6.93}$

The value of $\rho$ decisively influences the electronic properties of the system. Figure 4.2 schematically depicts the various phases of YBa$_2$Cu$_3$O$_{7-\delta}$ that can be reached by tuning of $\rho$ and the temperature. Superconductivity occurs at doping levels $0.05 < \rho < 0.27$. The critical temperature strongly depends on $\rho$, forming the characteristic “superconducting dome” in the $\rho$–$T$ phase diagram [Cav90, Huf08]. An O1 oxygen deficiency of $\delta = 0.07$ is referred to as “optimal doping” since it yields the maximum transition temperature for superconductivity of $T_c = 93$ K. At doping levels that allow a superconducting phase, an incommensurate charge ordered phase exists in a temperature range above the critical temperature for superconductivity [Cha12]. At low hole-doping levels ($\rho < 0.05$), the system enters an antiferromagnetic insulating phase with the Néel temperature increasing with a further reduction of $\rho$. For doping levels $\sim 0.045 < \rho < 0.07$, an incommensurate spin ordered phase exists [Hau10]. This phase does not occur together with the antiferromagnetic phase but can be simultaneously present with superconductivity [Cha12]. The pseudogap “phase” is observed in a temperature region above the superconducting dome. For further details on the observed phases and quantitative phase diagrams of YBa$_2$Cu$_3$O$_{7-\delta}$ see e.g. [Huf08, Hau10, Cha12, Hot13].

4.2. The spectral hallmarks of superconductivity in the dielectric response of YBa$_2$Cu$_3$O$_{6.93}$

Due to the layered structure, the dielectric response of YBa$_2$Cu$_3$O$_{7-\delta}$ is highly anisotropic. The in-plane response for an electric probe field $E$ perpendicular to the crystal $c$-axis is dominated by the superconducting planes. Figure 4.3(a) shows the real part of the in-plane ($E \perp c$) optical conductivity $\sigma$ in the normal and superconducting phases as measured.
4.2. The spectral hallmarks of superconductivity in the dielectric response of YBa$_2$Cu$_3$O$_{6.93}$

Figure 4.3: Optical conductivity $\sigma_1(\omega)$ of YBa$_2$Cu$_3$O$_{6.93}$ in the superconducting ($T = 10$ K, blue curve) and normal ($T = 100$ K, red curve) states for (a) $E \perp c$ and (b) $E \parallel c$ as measured by ellipsometry$^1$[Ber04]. The modes centered at 39 meV and 71 meV in (b) correspond to the bond-bending and apex oxygen vibrations, respectively. The yellow shaded areas mark the loss of conductivity induced by the opening of a superconducting gap.

by ellipsometry [Ber04]. Above $T_c$, the metallic conductivity of the cuprate planes yields a Drude-like conductivity. When the temperature is decreased below $T_c$, $\sigma_1$ drops at energies $\hbar \omega < 110$ meV due to the opening of a superconducting gap [Bor04, Bas05]. A Drude-like contribution persists at the low-energy side of the response even at $T = 10$ K, as it is expected for a superconducting gap with $d$-wave symmetry [VH95, Tsu00], where nodal quasi-particles exist at finite temperatures. According to oscillator sum rules for the optical conductivity [Dre02], the spectral weight defined by the hatched area in Figure 4.3(a) is proportional to the density of condensed quasi-particles [Bas05].

While the in-plane response is dominated by electronic excitations, their contribution to the inter-plane optical conductivity ($E \parallel c$) is reduced by one order of magnitude, as seen in Figure 4.3(b). This is why two infrared-active phonon modes of $B_{1u}$ symmetry appear prominently on top of the electronic background: (i) The narrow resonance at an energy of 39 meV is caused by Cu-O bond bending involving an in-phase motion of all oxygen ions within the cuprate bilayers (O3 in Figure 4.1) against the central yttrium ion. (ii) Collective vibrations of the apical oxygen ions (O2 in Figure 4.1) located between the bilayers and the chains account for the conductivity maximum centered at an energy of 71 meV [Liu88, Bat89]. The bond-bending mode exhibits an anomalous blue shift and a slight broadening (see Figure 4.3(b)) as the system is heated towards $T_c$.

---

$^1$Data were provided by C. Bernhard, Dept. of Physics, Univ. of Fribourg, 1700 Fribourg, Switzerland.
[Lit92, Sch95, Ber00, Mun99]. At the same time the apex line changes from an asymmetric shape in the superconducting state to an almost symmetric one above $T_c$ [Sch95]. The physical origin of the latter phenomenon is explained by the coupling between optical phonons and the interlayer Josephson plasmon [Mun99]. This collective oscillation of the superconducting condensate along the crystal $c$-axis, renormalizes the phonon parameters by modifying effective local fields. In the case of the apex mode, the dominant effect is a strong increase of the phonon asymmetry which, thus, quantifies the coupling of this mode to the Josephson plasmon and scales with the density of the superconducting condensate. The eigenfrequency of the apex phonon $\omega_0$, on the other hand, shows a continuous decrease with increasing temperature [Sch95]. This red-shift is characteristic of an anharmonic lattice potential which softens for larger vibrational amplitudes. Consequently, the frequency shift of the apex phonon may be exploited as a sensitive probe of its transient vibrational occupation after photoexcitation.

4.3. Preparations for two-time THz studies

The single crystal under study of $\text{YBa}_2\text{Cu}_3\text{O}_{6.93}$ was grown by top-seeded solution growth [Yao97]. The sample was annealed for 72 hours in flowing oxygen at a temperature of 500°C and returned to room temperature by furnace cooling to obtain optimally doped crystals. NIR-pump/THz-probe experiments are performed in reflection geometry (see Figure 4.4) on as-grown surfaces of the crystal with an area of about $2 \times 2 \text{mm}^2$. One of these adjacent surfaces is oriented parallel, the other perpendicular, to the $c$-axis of the twinned crystal. The measurements are taken with $s$-polarized THz pulses and a $p$-polarized NIR pump, both incident at an angle of 30°. Thus, the pump is mostly polarized in the $ab$-plane. The polarization of the THz probe field can be chosen as $\mathbf{E} \perp c$ or $\mathbf{E} \parallel c$ by aligning the sample accordingly. A homogeneous lateral excitation profile is ensured.

![Figure 4.4:](image)

Experimental arrangement for reflection-type NIR-pump/THz-probe studies. The setup is integrated into the LHe flow cryostat (see Figure 2.5). The red curve indicates the model excitation profile of the sample.

\(^2\)Samples were provided by X. Yao, Dept. of Physics, Shanghai Jiao Tong Univ., Shanghai 200240, China.
4.3. Preparations for two-time THz studies

by setting the diameter of the NIR pump focus twice the size of the THz probe focus of \( \sim 75 \mu m \). In the following, the photoexcitation intensity is stated as incident fluence \( \Phi _i \).

A transfer-matrix formalism is employed to extract the optical conductivity \( \sigma (\hbar \omega , t_D) \) in the photoexcited state from the measured pump-induced changes in the complex THz reflectivity. An inhomogeneously excited surface layer is treated as a stack of much thinner layers with a homogeneous refractive index as shown by gray lines in Figure 4.4. The model excitation profile in \( z \)-direction is described by an exponential decay with a length scale of 0.1 \( \mu m \) (sketched by the red curve in Figure 4.4) to account for the absorption length of the pump light [Kab99]. The transient THz optical response is determined by numerical inversion of the expression for the complex THz reflectivity, as described in section 2.2.2. Thereby, the dielectric response measured by ellipsometry is employed to describe the reflectivity of the unexcited system.

In order to determine the pump fluence that ensures full suppression of superconductivity and simultaneously keeps thermal heating of the sample low, the dependence of the maximal pump-induced change of the reflected THz transients \( \Delta E_{\text{max}} \) on \( \Phi _i \) is studied prior to two-dimensional pump-probe experiments. As seen in Figure 4.5, the pump-induced change as a function of \( \Phi _i \) shows a saturation behavior combined with a linear increase. This dependence can be phenomenologically fitted by the following equation

\[
\Delta E_{\text{max}}(\Phi _i) = A \left( 1 - e^{-\Phi _i/\Phi _{\text{sat}}} \right) + B_{\text{lin}} \Phi _i.
\] (4.1)

The first term in equation (4.1) describes the depletion of the superconducting condensate characterized by the saturation fluence \( \Phi _{\text{sat}} \). The linear second term is related to a direct photoinduced change in the quasiparticle density and the suppression of a pseudogap (see section 4.4). Fitting the experimental data (red spheres in Figure 4.5) results in a saturation fluence of \( \Phi _{\text{sat}} = 0.1 \text{mJ/cm}^2 \). A value of \( \Phi _i = 0.3 \text{mJ/cm}^2 \) clearly exceeds the saturation threshold while it keeps thermal heating at a negligible level and thus is chosen for the detailed two-dimensional experiments presented in the next sections.
Figure 4.6: 2D NIR-pump/THz-probe data: (a) transient pump-induced changes of the real part of the optical conductivity $\Delta \sigma_1(\hbar \omega, t_D)$ as a function of the photon energy and the pump-probe delay time $t_D$ for $E \perp c$. (b) Corresponding changes of the absorption coefficient $\Delta \alpha(\hbar \omega, t_D)$, for $E \parallel c$. Both measurements are performed at $T = 20$ K with $\Phi_i = 0.3$ mJ/cm$^2$.

4.4. Phonons and quasi-particles after photoexcitation

Figures 4.6 (a) and (b) depict the ultrafast pump-induced change in the optical response induced by NIR photoexcitation with a fluence of $\Phi_i = 0.3$ mJ/cm$^2$ at a sample temperature of $T = 20$ K.

The transient change of the in-plane response shown in panel (a) is discussed first. The pump pulse barely affects the high energy side of $\sigma_1(\hbar \omega)$ whereas the conductivity is enhanced below an energy of 110 meV due to the contribution of photogenerated quasiparticles. It is instructive to compare the absolute size and the spectral shape of the pump-induced conductivity changes to the conductivity difference between the superconducting and normal states (figure 4.7(a)). A snapshot of $\Delta \sigma_1$ taken at a delay time of $t_D = 1.5$ ps (red curve) perfectly matches the conductivity difference between the superconducting and the normal state (black curve), indicating a pump-induced phase transition to the normal phase and a full suppression of the superconducting gap.
4.4. Phonons and quasi-particles after photoexcitation

![Graphs](image)

**Figure 4.7:** (a) Difference in the real part of the optical conductivity between normal and superconducting states (black curve) for $E \perp c$ and pump-induced conductivity change $\Delta \sigma_1(h\omega, t_D)$ measured at delay time $t_D = 1$ ps after photoexcitation (red curve). (b) Dynamics of the photoinduced quasi-particle spectral weight as a function the delay time $t_D$. Cyan curve: spectral weight for $E \perp c$ integrated from 40 meV to 130 meV. Purple curve: spectral weight (scaled by factor 170) for $E \parallel c$ between 45 meV and 60 meV. The experimental time resolution is indicated by the hatched area.

The temporal evolution of the change of the spectral weight, $S_{QP} = \int \Delta \sigma_1(h\omega, t_D)d\omega$, in an energy window from 40 meV to 130 meV is shown in Figure 4.7(b). Interestingly, $S_{QP}$ reaches its maximum with a delay of 150 fs, which is distinctly slower than the experimental time resolution of 40 fs. These data present a direct observation of the quasi-particle density generated during the pair-breaking cascade on the inherent time scale. The subsequent decay is described well by two exponential functions with time constants of 0.4 ps and 3.7 ps, respectively. The slow part corresponds to recondensation of quasi-particles into Cooper-pairs, whereas the fast component survives even above $T_c$ and was associated to the recovery of a pseudogap [Dem99, Kai00].

As seen in Figure 4.6(b), the situation for $E \parallel c$ is yet richer in detail. While all essential features discussed in the following are also evident in $\Delta \sigma_1(\omega, \tau)$, the 2D map of the transient absorption change $\Delta \alpha(h\omega, t_D)$ illustrates the photoinduced dynamics most obviously: Besides a broadband quasi-particle contribution featuring a similar spectral profile as in the case of $E \perp c$, two narrow minima are superimposed at energies of 38 meV and 72 meV. These are the fingerprints of pump-induced modifications of the two phonon resonances. As evident from the 2D data, the various contributions to the total response follow clearly different dynamics: The spectral weight of the quasi-particles integrated
in the energy range from 45 meV to 60 meV, i.e. between the phonon resonances, shows 
the same temporal trace as the in-plane conductivity (see Figure 4.7(b)). In contrast, 
the pump-induced changes of the phonons are delayed. The most striking difference is 
imprinted on the apex phonon which exhibits its maximum change as late as 1 ps after 
photoexcitation (figure 4.6(b)).

To go beyond this qualitative observation, the line shape of the phonons is analyzed to 
single out the microscopic mechanisms underlying the pump-induced phonon anomaly.

4.5. Analysis of the transient phonon lineshapes

4.5.1. Excitation dynamics of the apex mode

Figure 4.8 shows selected spectra of the real and imaginary part of the c-axis conductivity 
in the vicinity of the apex phonon resonance. Below $T_c$, $\sigma_1(\hbar \omega)$ features a characteristic 
asymmetric lineshape (Figure 4.8(a)).

For a quantitative description, the dielectric response is modeled as the sum of a 
broad-band background electronic conductivity $\sigma_{el}(\omega)$ and a phenomenological model [Hum00] 
of the phonon resonance:

$$\sigma(\omega) = \sigma_{el}(\omega) - \varepsilon_0 S \frac{(\omega A + i\omega_0^2)\omega}{\omega_0^2 - \omega^2 - i\omega \gamma}, \quad (4.2)$$

The phonon contribution (second term) is defined by the oscillator strength $S$, the eigen-
frequency $\omega_0$, and the damping $\gamma$. The parameter $A$ accounts for the peculiar asymmetry 
of the lineshape below $T_c$. The damping $\gamma = 2\pi \times 0.5$ THz is found to be almost tem-
perature independent and is not considered here further. In order to reduce the number 
of free parameters, the quasi-particle background $\sigma_{el}(\omega)$ is considered as a locally linear 
function of frequency (first term in equation (4.2)).

The requirement to reproduce both real and imaginary parts of the spectra simultaneously 
imposes strict boundaries on the three remaining fit parameters $S$, $\omega_0$, and $A$. As seen 
in figures 4.8(a) and (b), the equilibrium mid infrared response of YBa$_2$Cu$_3$O$_{6.93}$ can be 
described convincingly with equation 4.2.

Figures 4.9 (a) and (b) show the resulting fit parameters $A$ and $\omega_0$ as a function of 
temperature. The asymmetry sharply decreases upon heating the system towards $T_c$. 
Above $T_c$, no further change is observed, owing to the absence of the Josephson plasmon. 
The increasing thermal occupation of the phonon mode upon heating manifests itself as 
a continuous decrease of its eigenfrequency, apart from a small anomaly around $T_c$. 

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4.5. Analysis of the transient phonon lineshapes

Figure 4.8: Spectra of the optical conductivity $\sigma_1(\hbar \omega)$ and $\sigma_2(\hbar \omega)$ of YBa$_2$Cu$_3$O$_{6.93}$ for $E \parallel c$: (a) and (b) show the equilibrium spectra for selected temperatures. (c) and (d) show the transient spectra at selected pump-probe delay times $t_D$ measured at $T = 200$ K with $\Phi = 0.3$ mJ/cm$^2$. The solid lines show the numerically adapted curves according to equation (4.1).

The model function of equation 4.2 can also be numerically adapted to the transient complex conductivity spectra with high confidence, as shown in figures 4.8 (c) and (d) for representatively selected pump-probe delay times. This allows one to map out pump-induced modifications of the phonon parameters on a femtosecond scale.

Starting from the superconducting state (blue curve in Figure 4.9 (d)), the asymmetry of the apex mode decreases within $t_D \leq 150$ fs, following the depletion of the superconducting condensate. Analogously, the relaxation to the initial line shape asymmetry follows the recovery of the superconducting gap with a typical time constant of about 4 ps. This confirms a coupling to the Josephson plasmon as the microscopic origin of the asymmetry.

Note that unlike in a thermal transition to above $T_c$, ultrafast optical excitation suppresses the asymmetry entirely for $t_D < 1$ ps. This indicates an extremely non-thermal state of the system at early pump-probe delay times. The eigenfrequency of the apex mode (blue curve in Figure 4.9(c)) experiences an abrupt red-shift within 150 fs after photoexcitation, i.e. during the onset of the electronic response, and reaches its extremum within 300 fs. The
Chapter 4. Phonon and quasiparticle dynamics in superconducting YBa$_2$Cu$_3$O$_{6.93}$

Figure 4.9: (a) Eigenfrequency and (b) asymmetry parameter of the apex mode as functions of temperature. (c) and (d) show the corresponding quantities as functions of the pump-probe delay time $t_D$ after photoexcitation with $\Phi_i = 0.3$ mJ/cm$^2$. The blue and red curves are obtained by photoexcitation of the superconducting ($T = 20$ K) and the normal state ($T = 100$ K), respectively. The error bars indicate 95\% confidence intervals for the fitting parameters.

maximum photoinduced softening of the apex mode is comparable to the effect induced by a thermal phonon population at $T \approx 200$ K. Thus, it can be inferred that the optical pump causes a hot phonon population of the apex mode within 150 fs. Subsequently, $\omega_0$ relaxes to a value which is slightly reduced with respect to the equilibrium level at $T = 20$ K. The relaxation time of 1 ps (see Figure 4.9(c)) is significantly faster than the quasi-particle recondensation, which further confirms that the phonon softening is not governed by the coupling to the Josephson plasmon. It is suggested [Pas10], that the rapid recovery of $\omega_0$ is explained by a redistribution of the excess energy by phonon-phonon scattering into the large phase space of the entire Brillouin zone. The damping of the apex mode corresponds to a phonon lifetime of 2 ps which exceeds the observed relaxation time significantly. This finding shows that phonon-phonon scattering is faster during the hot non-equilibrium state after photoexcitation as compared to the situation in thermal equilibrium. Additional corroboration for a hot phonon population as the main cause of the red-shift of the apex mode is provided by the dynamics in the normal state (red curve in 4.9(c)) where $\omega_0$ follows a quantitatively similar trace. The asymmetry parameter in the normal state, in contrast, shows a substantially reduced change owing to the absence of a macroscopic condensate (red curve in Figure 4.9(d)). These observations prove, that in the non-equilibrium system scattering processes between the apex mode
4.5. Analysis of the transient phonon lineshapes

Figure 4.10: Temperature dependencies of (a) the eigenfrequency of the bond-bending mode and (b) the quasi-particle spectral weight for $E \parallel c$ integrated between 45 meV and 60 meV. (c) and (d) show the corresponding quantities as functions of the pump-probe delay time $t_D$ after photoexcitation of the superconducting state ($T = 20$ K) with $\Phi_i = 0.3 \text{ mJ/cm}^2$. The error bars indicate 95% confidence intervals for the bond-bending mode eigenfrequency.

and the electronic system occur on the same timescale as electron-electron scattering. In the next step, it is studied whether this also holds for the bond-bending mode.

4.5.2. Excitation dynamics of the bond-bending mode

Since there exists an anomalously strong influence of the Josephson plasmon coupling on the frequency evolution of the bond-bending mode [Ber00], it is not possible to disentangle it from the anharmonic population effect as clearly as in the case of the apex mode. Nevertheless, it is instructive to compare its equilibrium properties and pump induced dynamics to the behavior of the apex mode. To this end, a Lorentzian oscillator is locally fitted on the resonance to obtain its eigenfrequency as a function of temperature and pump-probe delay time. In contrast to the apex mode, the asymmetry of the bond-bending mode is small and only weakly affected by the Josephson plasmon.

The dominant effect of coupling between the Josephson plasmon and the bond-bending phonon is given by a change of the phonon eigenfrequency $\omega_0$. As temperature increases towards $T_c$, the density of the superconducting condensate and its coupling to the bond-bending mode vanish, leading to the blue-shift of $\omega_0$ as shown in Figure 4.10(a). Pump-induced suppression of the superconducting condensate also leads to a blue-shift of the
bond-bending mode (figure 4.10(c)). However, its dynamics does not reproduce the evolution of the quasi-particle response shown in Figure 4.10(d) although the temperature dependence of $\omega_0$ below $T_c$ clearly follows the quasi-particle spectral weight (compare figures 4.10(a) and (b)). The initial rapid increase of the eigenfrequency takes about 300 fs, which is two times slower than the depletion of the superconducting condensate. This stage is followed by a delayed growth of the eigenfrequency on the picosecond time scale. For $t_D > 2$ ps, the resonance frequency finally shifts back to its equilibrium value.

Above $T_c$, the bond-bending mode indicates a notable anharmonic red-shift (figure 4.10(a)) similar to the behavior of the apex phonon. Without Josephson coupling to the superconducting condensate, this phenomenon can be expected to persist also below $T_c$. The effective blue-shift of the bond-bending resonance below $T_c$, thus, results from the counteracting influence of Josephson coupling and lattice anharmonicities. The ultrafast dynamics depicted in Figure 4.10(c) may be understood as follows: Photoexcitation destroys the condensate and depletes phonon-plasmon coupling. The associated blue-shift of the phonon frequency is therefore expected to occur within 150 fs after the pump pulse. The delayed component of the onset dynamics of the eigenfrequency may be associated with the lattice anharmonicity induced by a hot phonon population. As the occupation number of the bond-bending mode decays, the red-shift relaxes and leads to a further increase of the phonon eigenfrequency on a picosecond time scale. Since the Josephson plasmon remains suppressed for a few picoseconds, the eigenfrequency may exceed the values observed in thermal equilibrium during this period, since the anomalous red-shift due to the Josephson coupling is suppressed while the phonon occupation number corresponds to a temperature below $T_c$. Together with the recovery of the Josephson resonance, the eigenfrequency ultimately starts to return to its equilibrium frequency.

Importantly, the relatively slow initial onset of the bond-bending mode anomaly within 300 fs indicates a different timescale of the electron-phonon interaction for this mode as compared to the apex mode.

4.6. Résumé: Electron-electron and electron-phonon scattering

The different timescales of the electron-phonon interaction observed for the two phonon modes under study permits one to exclude a scenario in which the phonon anomalies are caused by a photoinduced bond softening as observed, for example, in bismuth [Fri07]. In this case, the onset dynamics for all phonon modes would be the same since it is governed by the density of the photoexcited electrons. Such a behavior is in contrast to the experimental observation of the phonon dynamics in $\text{YBa}_2\text{Cu}_3\text{O}_{6.93}$. 

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Furthermore, the results clearly indicate that a major portion of the absorbed pump energy is transferred to the phonon subsystem within 150 fs after photoexcitation of the electronic system. Further evidence supporting this argumentation comes from an energy balance of the pump-induced transition into the normal state. The energy density of $\simeq 10 \text{J/cm}^3$ required to suppress the superconducting phase optically (see section 4.3) is much higher than the thermodynamically determined condensation energy of $1.2 \text{J/cm}^3$ [Lor93]. Following the lines of Kusar et al. [Kus08], only the phonon subsystem possesses sufficient heat capacity to dissipate the high excess energy of the pump pulse.

It is worth comparing this situation with the two- or three-temperature models [All87, Per07]. These theories have been established assuming the electron-phonon scattering rate to be negligible as compared to electron-electron interaction. In conclusion, this temporal hierarchy of scattering processes put forward in the context of normal metals [All87] or high-temperature superconductors such as Bi$_2$Sr$_2$CaCu$_2$O$_{8+\delta}$ [Per07] does not hold in case of YBa$_2$Cu$_3$O$_{7-\delta}$ where a hot phonon population and pair breaking occur on comparable time scales. In this work, it is found that for the case of YBa$_2$Cu$_3$O$_{7-\delta}$ electron-electron scattering and electron-phonon scattering processes occur on the same timescale.
Non-adiabatic switching of ultrastrong light-matter coupling

The two previous chapters reported new insights into the microscopic mechanisms behind the formation of macroscopic states in bulk solids resulting from strongly correlated low-energy excitations defined by nature. In this chapter, an artificial type of interaction between tailor-cut elementary excitations is studied: Ultra-strong coupling between a custom designed electronic resonance and a tailored optical eigenmode of a microcavity. The incentive to engineer the coupling strength between light and matter resonances is mainly based on the prospect to observe novel quantum-electrodynamical phenomena in these systems. Such effects are expected to occur when the light-matter interaction strength is modulated on a sub-cycle timescale. In this work, it is demonstrated that the matter part of an ultrastrongly coupled system can be activated non-adiabatically by a NIR control pulse while a femtosecond multi-THz pulse monitors signatures of the interaction strength. The development of this technology aims to ultimately provide a route towards experimental observation of a phenomenon reminiscent of the dynamical Casimir effect: The emission of photons out of the quantum vacuum [Ciu05, Lib07].

The first sections of this chapter introduce the physics of ultrastrong light-matter interaction and the technology of the optically switchable ultrastrong light-matter coupling device. Next, it is demonstrated via femtosecond multi-THz spectroscopy of the device’s photonic band structure that optical activation of an electronic resonance via a NIR pump pulse enables optical control of the light-matter interaction strength. Finally, it will be shown that optical excitation may be employed to switch ultrastrong light-matter interaction non-adiabatically. This part of the work has been published in Physical Review B: Rapid Commun. [Por12].
5.1. Regimes of light-matter coupling

One of the key challenges of modern quantum optics is to control the interaction of light with electronic excitations. A powerful technological approach to artificially enhance the strength of light-matter interaction is to couple an optical mode of a microcavity to a cavity-embedded matter resonance. Intuitively, the strength of light-matter interaction is quantified by the vacuum Rabi frequency $\Omega_R$, the rate at which a virtual cavity photon is absorbed and spontaneously reemitted by the quantized emitter.

Three regimes of light-matter coupling are distinguished (compare Figure 5.1): (i) If the loss rate of the cavity $\gamma$ and the dephasing rate of the radiative transition $\kappa$ (reflected by the linewidths of the cavity and the matter resonance, respectively) exceed $\Omega_R$, the system is in the weak coupling regime. In this case, the discrete density of photonic states modifies the lifetime of the matter excitation (Purcell effect [Pur46]). (ii) The system enters the regime of strong coupling, when $\Omega_R$ exceeds $\gamma$ and $\kappa$. The eigenstates of the coupled system are then given by hybrid light-matter mixed modes, called cavity polaritons. A first demonstration of these quasi-particle excitations was achieved using atoms in metallic cavities [Rem87]. A few years later, this regime was reached via coupling of
an excitonic matter transition to a semiconductor microcavity [Wei92]. The distinctive
dispersion of cavity polaritons [Kli07] described by two anti-crossing polariton-branches,
which are separated by the characteristic energy of $2\hbar \Omega_R$, has facilitated fascinating solid-
state quantum optics, including electrically driven inversionless lasing [Bha14] or Bose
condensation of exciton-polaritons [Den02, Kas06, Mén14]. (iii) One of the most in-
triguing aspects is the limit of ultrastrong coupling, where $\Omega_R$ becomes comparable with
the transition frequency $\omega_{12}$ of the matter excitation itself. This extreme case has been
reached by hybridizing discrete transitions between electronic subbands in semiconductor
quantum wells with the mid-infrared photonic mode of a planar waveguide structure
[Din03, Dup07, Ana09, Gün09]. By exploiting the strong field enhancement in plasmonic
metal structures, extremely high energy splitting of the two polariton branches by as
much as $2\Omega_R = 0.5 \times \omega_{12}$ has been achieved [Tod09, Tod10, Zan10, Gei10]. Alternative
approaches to reach the ultrastrong coupling regime involving superconducting circuits
[Nie10, FD10] or large molecules [Sch11] have followed.

An appropriate quantum-mechanical description of ultrastrong light-matter coupling has
to go beyond the rotating wave approximation. Such an approach includes the antireso-
nant terms of the light-matter coupling Hamiltonian and provides an adequate description
of the polariton vacuum ground state $|G\rangle$ of the ultrastrongly coupled system. With this,
it can be shown that $|G\rangle$, a two-mode squeezed vacuum state, is distinctly different from
the vacuum ground state $|0\rangle$ of the uncoupled system [Ciu05]. The defining property of
a vacuum ground state is that no eigenmode of the system is in an excited state. For
the uncoupled system, this means that neither a photon (created by $a^\dagger$) nor a matter
excitation (created by $b^\dagger$) is present, i.e.

$$
\langle 0\mid a^\dagger a\mid 0 \rangle = 0 \quad \text{and} \quad \langle 0\mid b^\dagger b\mid 0 \rangle = 0. \quad (5.1)
$$

In the polariton ground state, no polariton excitation (created by $p^\dagger$) exists:

$$
\langle G\mid p^\dagger p\mid G \rangle = 0. \quad (5.2)
$$

The expectation of non-classical effects mainly relies on the fact [Ciu05] that the polarit-
on ground state of the ultrastrongly coupled system $|G\rangle$ is occupied with virtual cavity
photons and that the ground state of the bare cavity mode $|0\rangle$ is populated with virtual
polaritons:

$$
\langle G\mid a^\dagger a\mid G \rangle \neq 0 \quad (5.3)
$$

$$
\langle 0\mid p^\dagger p\mid 0 \rangle \neq 0. \quad (5.4)
$$

These excitations are called virtual since they do not correspond to an excitation of
an eigenmode of the respective system and can only exist for a timespan allowed by the
energy-time uncertainty. They are unobservable since their detection would violate energy conservation. However, if the system is transformed non-adiabatically (i.e. on a timescale shorter than a single oscillation cycle of the involved modes) from one ground state to the other, the virtual excitations are predicted to suddenly become a real excitation of the system and should escape as entangled photons to the external space [Ciu05]. This process is comparable to the dynamical Casimir effect [Yab89, Ciu05, Lib07] and has been studied theoretically in detail for the case of a periodic modulation of $\Omega_R$ in an intersubband-polariton system [Ciu05, Lib07]. Recent publications report indications of quantum vacuum radiation emerging at an artificial event horizon [Bel10] and during rapid modulation of superconducting circuits [Wil11]. However, these remarkable findings remain controversially discussed and evidence of such non-classical radiation in the optical regime is still lacking.

Non-adiabatic dynamics of mid-infrared intersubband-polaritons entering the ultrastrong coupling regime has become accessible with field-resolved sub-cycle multi-THz spectroscopy in a seminal work by Günter et al. [Gün09]. This experiment demonstrated sub-cycle activation of ultrastrong light-matter coupling between a 2D waveguide mode and an embedded optically switchable intersubband [Yu10] resonance, for the first time. However, sub-cycle control of the light-matter interaction with NIR control pulses required elaborate tilting of the NIR pulse fronts. Furthermore, a complex prism-coupling geometry enabled only limited access on the relevant polariton modes. Since Casimir photons are expected to appear as excitations of the polariton modes, it is essential to obtain full femtosecond access to these modes in order to probe their population after the non-adiabatic switching event.

In the next section, a novel light-matter coupling device is introduced that enables to probe a coherent population of the relevant polaritonic bands by means of multi-THz electro-optic sampling in straightforward transmission geometry. This approach constitutes a further step towards the observation of the dynamical Casimir effect in intersubband-polariton systems.

### 5.2. A device for optical control of light-matter coupling

#### 5.2.1. Design of the switchable matter excitation and the photonic mode

Intersubband transitions in quantum wells [Ros02a, Yu10] are naturally suited for engineering the enhancement of light-matter interaction: Due to the spatial extent of subband wavefunctions of typically 10 nm given by the dimensions of the quantum wells, the intersubband dipole moment is much larger than the dipole moment of interband transitions,
5.2. A device for optical control of light-matter coupling

Figure 5.2: Design of the switchable ultrastrong light-matter coupling device. The refractive index contrast between vacuum, AlGaAs/GaAs heterostructure (quantum wells (QWs) and cladding layer) and GaAs substrate forms a planar refractive index waveguide. Together with a lateral gold grating, the structure constitutes a photonic crystal. Its optical eigenmodes with polarization components in the growth direction couple to an optically switchable intersubband transition located inside the quantum well region.

which is determined by the atomic dipoles of the underlying Bloch functions. To custom tailor also the light field for maximized light-matter interaction, photonic crystals ([Joa08]) provide an ideal solution. Furthermore, the dispersion of their optical eigenmodes can be precisely controlled to yield a design-cut photonic band structure. Zanotto et al. [Zan10] employed this concept to implement ultrastrong light-matter interaction between an intersubband transition and the custom tailored eigenmode of a one-dimensional photonic crystal. The photonic mode was engineered such that the polaritonic modes could be mapped out via equilibrium spectroscopy in simple reflection geometry.

This work combines the photonic crystal introduced by Zanotto et al. with a switchable intersubband transition [Gün09]. This way, full access to the polariton modes is obtained while light-matter interaction can be activated non-adiabatically with a NIR control pulse. The next paragraph summarizes the structure of the novel design. The specific choices of the structural and compositional parameters are explained in the subsequent paragraphs.

Figure 5.2 illustrates the switchable light-matter coupling device. The fabrication of the structure\(^1\) is separated into two steps: First, a 4 µm thick AlGaAs/GaAs heterostructure is grown by molecular beam epitaxy onto a 350 µm thick GaAs substrate. The heterostructure is comprised of a Al\(_{0.95}\)Ga\(_{0.05}\)As cladding layer with a thickness of 2 µm and

\(^1\)Samples were fabricated by S. Zanotto and G. Biasol (Laboratorio TASC, Trieste, Italy) as well as R. Degl’Innocenti, and L. Sorba (NEST, Instituto Nanoscience, Pisa, Italy).
a quantum well structure on top consisting of 50 repetitions of 8.3 nm thick GaAs wells separated by 30 nm thick Al\textsubscript{0.33}Ga\textsubscript{0.67}As barriers. In a second step, a one-dimensional gold grating (thickness: 40 nm, fill factor: 0.6) with period $a = 4.1 \mu$m is fabricated on top of the structure via thermal evaporation of the gold layer and electron-beam lithography. This architecture defines the intersubband resonance and the surface plasmon photonic mode while enabling ultrastrong coupling between both.

Figure 5.3 illustrates the design of the quantum well heterostructure. The width of 8.3 nm of each GaAs quantum well is chosen such that the energetic distance between the lowest electronic conduction subbands amounts to $\hbar \nu_{\text{ISB}} = 125 \text{ meV}$ corresponding to a photon frequency of $\nu_{\text{ISB}} = 30 \text{ THz}$. The width of 30 nm of the Al\textsubscript{0.33}Ga\textsubscript{0.67}As barriers confines

\[ \Delta E/h = 30 \text{ THz} \]

\[ \Delta E_{\text{min}} = 1.57 \text{ eV} \]

\[ |2\rangle_e \]
\[ |1\rangle_e \]
\[ |1\rangle_{hh} \]
\[ |2\rangle_{hh} \]

\[ k_{\parallel} \text{ (arb. u.)} \]

\[ z\text{-position (nm)} \]

\[ \text{Energy} \]

Figure 5.3: (a) Design of the multiple quantum well heterostructure. Upper and lower gray solid lines show the confinement potential given by the conduction and valence band edges as a function of the position $z$ along the growth direction. Subband energy levels for zero in-plane electronic wave vector $k_{\parallel}$ are indicated by horizontal lines. Dashed lines indicate the envelope wavefunctions of the electronic subband states\textsuperscript{2}. (b) Dispersion of the subbands. NIR optical pumping with a minimum photon energy of $\Delta E_{\text{min}}$ (red arrow) excites carriers from the uppermost heavy hole subband $|1\rangle_{hh}$ to the lowest electronic subband $|1\rangle_e$, activating the electronic intersubband transition $|1\rangle_e \rightarrow |2\rangle_e$ with a $k_{\parallel}$ independent transition energy of $\hbar \times 30 \text{ THz}$ (indicated by green arrows in (a) and (b)).

\[ |2\rangle_e \]
\[ |1\rangle_e \]
\[ |1\rangle_{hh} \]
\[ |2\rangle_{hh} \]

\[ k_{\parallel} \text{ (arb. u.)} \]

\[ z\text{-position (nm)} \]

\[ \text{Energy} \]

\[ \Delta E_{\text{min}} = 1.57 \text{ eV} \]

\[ \Delta E/h = 30 \text{ THz} \]

\[ a \text{ quantum well structure on top consisting of 50 repetitions of } 8.3 \text{ nm thick GaAs wells separated by } 30 \text{ nm thick } \text{Al}_{0.33}\text{Ga}_{0.67}\text{As barriers. In a second step, a one-dimensional gold grating (thickness: } 40 \text{ nm, fill factor: } 0.6 \text{) with period } a = 4.1 \mu\text{m is fabricated on top of the structure via thermal evaporation of the gold layer and electron-beam lithography. This architecture defines the intersubband resonance and the surface plasmon photonic mode while enabling ultrastrong coupling between both.} \]

\[ \text{Figure 5.3 illustrates the design of the quantum well heterostructure. The width of } 8.3 \text{ nm of each GaAs quantum well is chosen such that the energetic distance between the lowest electronic conduction subbands amounts to } \hbar \nu_{\text{ISB}} = 125 \text{ meV corresponding to a photon frequency of } \nu_{\text{ISB}} = 30 \text{ THz. The width of } 30 \text{ nm of the Al}_{0.33}\text{Ga}_{0.67}\text{As barriers confines} \]

\[ ^2\text{The spacing of the energy levels and the wavefunctions are calculated with the nextnano}\textsuperscript{3} \text{ tool.} \]
5.2. A device for optical control of light-matter coupling

Figure 5.4: Schematic photonic band structure of the device for inactive light-matter interaction. The eigenfrequency of the planar transverse magnetic TM0 refractive index guided mode as a function of the in-plane wavevector $k_{\parallel}$ is sketched as blue dashed curve. The Brillouin zone boundary at $k_{\parallel}a = \pi$ (solid vertical line) induced by the gold grating with period $a$ folds the guided mode into the first Brillouin zone, yielding the photonic band structure of the device (blue curves). The dispersion of the intersubband transition (not activated) is indicated by a dashed green line. The red dotted lines show the dispersion of external vacuum plane waves for two extreme values of the angle of incidence $\theta$. The orange shaded region can be probed experimentally.

the electronic envelope wavefunctions to a single quantum well, ensuring discrete subband states. Optical excitation of the undoped quantum well heterostructure with NIR pulses centered around a photon energy of 1.55 eV excites electrons from the uppermost heavy hole subband $|1\rangle_{hh}$ to the lowest electronic subband $|1\rangle_e$ (red arrow in Figure 5.3). Interband transitions are allowed only between hole and electron subbands with the same parity. Once the lowest energetic subband level is populated, the intersubband transition $|1\rangle_e \rightarrow |2\rangle_e$ (green arrows in Figure 5.3 (a) and (b)) is active. Its overall dipole moment is oriented in growth direction and scales with the density of photogenerated carriers $\rho_{1e}$ in the $|1\rangle_e$ subband. To efficiently populate the $|1\rangle_e$ subband states with nonzero in-plane wavevector $k_{\parallel}$ (compare Figure 5.3 (b)), the bandwidth of the NIR pump pulse needs to be sufficiently high.

For ultrastrong light-matter interaction, a confined optical mode with maximized coupling to the intersubband dipole moment is shaped inside the quantum well region. The first element for optical mode confinement is the planar refractive index waveguide formed by
Chapter 5. Non-adiabatic switching of ultrastrong light-matter coupling

the epitaxially grown layers and the substrate. The cladding layer serves to enhance the refractive index contrast. Transverse magnetic (TM) guided modes feature an electric field in growth direction and thus couple to the intersubband dipole moment. The second element for optical mode confinement is the one-dimensional gold grating on top of the planar waveguide. The purpose of this photonic crystal structure is two-fold. First, it enhances the field confinement itself (for details see section 5.2.2). Second, it tailors the dispersion of the guided modes in a way that they can be mapped out via straightforward transmission spectroscopy, as described in the following. Transmission or reflection spectroscopy can probe only photonic modes that share photon frequency and in-plane wavevector $k_{\parallel}$ with external plane waves. To illustrate the situation, Figure 5.4 compares the photon frequency dispersion of the lowest energy planar TM mode (blue dashed line) with the dispersion of external plane waves given by

$$k_{\parallel} = \frac{2\pi\nu \times \sin(\theta)}{c},$$

where $\theta$ is the angle of incidence. Red dotted lines show the plane wave dispersion for normal ($\theta = 0^\circ$) and streaking ($\theta = 90^\circ$) incidence. In the case of streaking incidence, the dispersion indicates the maximum values of $k_{\parallel}$ achievable with external plane waves. Thus, optical eigenmodes of the system with higher in-plane wavevector are totally confined within the structure and, in turn, cannot be probed via an external electromagnetic radiation (except via a disruption of the waveguide, e.g. by a prism-coupling geometry). In contrast, any eigenmode in the accessible “leaky” region of the photonic band structure (highlighted by an orange shading) can be fully accessed by appropriate tuning of the in-plane wavevector of the external plane waves by variation of $\theta$. The one dimensional photonic crystal on top of the structure induces a Brillouin zone boundary for the planar waveguide mode and folds its dispersion back into the first Brillouin zone. The period of the grating is chosen such that the polariton forming of the intersubband transition (green dashed line in Figure 5.4) and the back folded mode (blue line in Figure 5.4) lies inside the leaky region that couples to the external radiation field.

In the next sections it is shown that the polaritonic dispersion of the ultrastrongly coupled system can be mapped out with multi-THz time-domain spectroscopy on a sub-cycle timescale while light-matter interaction is activated non-adiabatically.

5.2.2. A switchable photonic bandstructure

Figure 5.5 shows typical multi-THz field transmission spectra of the device as a function of $k_{\parallel}$ captured by few-cycle THz-probe transients with a duration of 100 fs (generated and electro-optically sampled using 50 $\mu$m thick GaSe crystals). In the field transmission map of the unpumped device (Figure 5.5(a)), two photonic bands can be identified. Both modes are degenerate at $\nu = 26.6$ THz, in the center of the Brillouin zone, but split with
5.2. A device for optical control of light-matter coupling

![Figure 5.5](image)

**Figure 5.5:** Experimental field-transmission spectra of the photonic crystal structure \((a = 4.1 \mu m)\) as a function of \(k_{||}\) for \(0 \leq \theta \leq 40^\circ\). Results are measured (a) in equilibrium and (b) \(t_D = 1\) ns after photo-activation of the intersubband transition with \(\Phi = 1.6\) mJ/cm\(^2\). Black spheres: local transmission peaks indicating the eigenmodes. Solid black lines: numerically adapted linear dispersion in (a) and polariton dispersion in (b). Dotted black line in (a) and (b): dispersion of the lower photonic band. Dashed gray line in (a): Cross section indicating the spectrum measured at \(\theta = 20^\circ\) studied in Figure 5.6.

Increasing \(k_{||}\). Qualitatively these states arise from the planar waveguide mode which is Bragg scattered by the reciprocal lattice vectors \(\pi/a\) and \(-\pi/a\), respectively (compare Figure 5.4). Photoinjecting electrons into the lowest electronic subband \(|1\rangle_e\) of each quantum well with a NIR pump fluence of \(\Phi = 1.6\) mJ/cm\(^2\) induces a strong modification of the band dispersion, as directly mapped out in Figure 5.5(b) for \(t_D = 1\) ns. The interaction between the intersubband resonance and the upper photon band causes the formation of two polariton branches (black spheres and black solid lines in Figure 5.5(b)).

The minimum separation of the polariton branches occurring at the anticrossing point may be directly identified with the vacuum Rabi splitting \(2\Omega_R/2\pi\). Experimentally, the minimum energy splitting of the polariton modes is observed for \(\theta = 20^\circ\) and amounts to \(\hbar \times 3\) THz. Furthermore, both polariton bands are strongly flattened as compared to the unperturbed photon dispersion.

To gain further insight into the microscopic nature of the eigenmodes, the spectral shape of a typical transmission resonance is compared with the results of a FDFD calculation\(^3\)

\(^3\)The calculations are based on the total field/scattered field formalism [Rum06]. The simulations employ parts of a tutorial FDFD Matlab code provided by R. Rumpf, EM Lab, University of Texas.
Figure 5.6: (a) Field transmission of the plasmonic crystal structure ($a = 4.1 \mu m$) as a function of photon frequency measured (red dots) and calculated via a FDFD simulation (red line) for an angle of incidence of $\theta = 20^\circ$. The blue curve shows the corresponding calculated electrical field in growth direction integrated inside the quantum wells as a function of frequency. (b) Analysis of the electric field enhancement inside the structure. The color maps show the electrical field in growth direction that couples to the intersubband transition inside the quantum well (QW) region. The situation is shown for three characteristic photon frequencies (indicated by vertical dashed lines in (a)): (i) Off-resonance and at the upper photonic band at its transmission minimum (ii) and its maximum field enhancement (iii).

[Rum06]. Figure 5.6 (a) shows the experimental transmission curve of the unpumped structure measured at $\theta = 20^\circ$ (red dots) and the corresponding simulated spectrum (red solid curve). The simulation quantitatively recovers the Fano-like line shape [Zan12] and, additionally, reveals the electrical field enhancement inside the quantum well region. The blue solid curve shows the electric field in growth direction integrated inside the quantum well region. The peaks at 22.5 THz and 27.3 THz correspond to the field confinement caused by the lower and the upper photonic band, respectively. Furthermore, the FDFD simulation enables to link the transmission spectra with the underlying electromagnetic modes. Figure 5.6 (b) illustrates the spatial profile of the electric field $|E_z|$, oriented along the growth direction for three characteristic frequencies. Off resonance (i) at $\nu = 16$ THz, only a plasmonic field enhancement occurs at the edges of the gold stripes. The transmission minimum at $\nu = 27$ THz (ii) originates from the Fano-like resonance of the upper photon band. The field enhancement at this photon frequency is still mostly of plasmonic nature. Maximum field confinement occurs at $\nu = 27.3$ THz with the photonic mode strongly localized within the quantum well layers. Thus, the eigenmodes of the photonic
5.2. A device for optical control of light-matter coupling

Figure 5.7: Calculated field-transmission spectra of the (a) unpumped and (b) pumped sample for $0 \leq \theta \leq 89.9^\circ$. In (b), the intersubband transition is active with a center frequency of 29 THz and an assumed electron density in the lowest electronic subband of $\rho_{1e} = 2 \times 10^{11} \text{ cm}^{-2}$. Black dotted curves indicate the eigenmodes. Dashed gray lines in (a): cross section displayed in Figure 5.6(a). White tetragons: experimentally covered area shown in Figure 5.5.

Crystal are the result of the concerted action of surface plasmons and planar waveguiding by total internal reflection due to the refractive index contrast.

Repeating the simulation of the transmission spectrum for various angles of incidence, the full photonic band structure is recovered as seen in Figure 5.7 (a). In order to model the band structure of the strongly coupled system (Figure 5.7 (b)), the intersubband dipole moment is treated classically via an effective refractive index [Din03]. With a population of the lowest electronic subband of $\rho_{1e} = 2 \times 10^{11} \text{ cm}^{-2}$, good agreement with the experimental transmission map of Figure 5.5 (b) is achieved. The dispersion of all bands, the strength of light-matter coupling and the absolute value of field transmission are adequately reproduced. Furthermore, the FDFD analysis suggests that the surface plasmonic contribution increases $\Omega_R$ by 50% as compared to a structure supporting purely photonic propagation alone.

It may be noted, that under the action of extremely strong light-matter coupling, the slope of the polaritonic bands in the light matter coupling device is profoundly reduced as compared to the dispersion of the bare photonic mode (compare Figures 5.5 (a) and (b)). This effect gives rise to a slow-down of the group velocity $d\omega/dk_\parallel$ of the guided modes. For a quantitative discussion of this effect, see appendix E.


Chapter 5. Non-adiabatic switching of ultrastrong light-matter coupling

Figure 5.8:
Transmission spectra of the photonic crystal structure (substrate removed) showing the polariton formation at the anticrossing point ($\theta = 13^\circ$) measured at $t_D = 1\text{ ns}$ for various incident pump fluences $\Phi$. To ensure that additional membrane guided modes are far above the measurement window, the intersubband transition is now coupled to the lower photon band by setting $a = 3.6\mu m$. All curves are separated by a linearly increasing offset. Dashed vertical lines indicate the transmission maxima corresponding to the two polariton branches.

5.3. Optical control of the light-matter coupling strength

In preparation for time resolved experiments, the GaAs substrate of the heterostructure is locally removed by chemical etching with a citric acid/hydrogen peroxide mixture to obtain a free standing membrane with minimal dispersion for ultrafast laser pulses. Pump and probe beams are directed onto the rear side of the sample to inject a homogeneous distribution of carriers and prevent lateral diffusion effects from affecting the intrinsic dynamics. The increased refractive index contrast between the cladding layer and air gives rise to additional photonic bands of higher energy in the membrane. To ensure that the new resonances are far above the measurement window, the intersubband transition is now coupled to the original lower photonic band by setting the period of the gold grating to $a = 3.6\mu m$.

In a first step towards femtosecond control of ultrastrong coupling it is investigated how the strength of optically induced light-matter interaction scales with the intensity of NIR photoexcitation. To this end, transmission spectra through the sample are recorded for various values of $\Phi$ at a fixed delay time $t_D = 1\text{ ns}$ with $\theta = 13^\circ$ set to the anticrossing point (Figure 5.8). In the unpumped case, a single resonance with its field transmission maximum at 29 THz and a narrow width of 1.4 THz (FWHM) is observed. The width
5.4. Nonadiabatic switching of a photonic bandstructure

indicates a quality factor of $Q = 20$ of the uncoupled photonic cavity. With increasing pump fluence, the single peak splits into two polariton branches of comparable strength. The splitting of the polariton branches gradually increases with $\Phi$. Since the spectra are acquired at the anticrossing point, the splitting is a direct measure of $2\Omega_R/2\pi$. As quantitatively studied in Figure 5.9, the vacuum Rabi splitting (red dots) scales with the square root of the pump fluence for $\Phi < 120\mu J/cm^2$, analogously to the situation in planar waveguides [Ciu05, Gün09]. Yet $\Omega_R$ starts to saturate at $\Phi \approx 400\mu J/cm^2$.

A comparison with FDFD simulations suggests that a photogenerated carrier density of $\rho_{1e} = 3 \times 10^{11} cm^{-2}$ is present in this situation.

The results presented so far demonstrate full spatial control of the photonic eigenmodes of the light-matter coupled device. In addition, all optical control of the light-matter coupling strength between the custom tailored photonic mode and the intersubband excitation is realized. In the next section, it is shown that this control can be extended towards the temporal dimension.

5.4. Nonadiabatic switching of a photonic bandstructure

Nonadiabatic control of ultrastrong light-matter interaction requires the duration of the control pulse to be distinctly shorter than the oscillation period of the coupled resonances. The NIR pulses derived from the Ti:Sapphire amplifier system with a duration of 12 fs readily fulfill this condition. In order to explore how rapidly the hybridization of the photonic band structure with the intersubband transition proceeds, the pump-probe delay time $t_D$ is varied between $-120 fs$ and $1000 ps$ (see Figure 5.10). The pump fluence is kept constant at $\Phi = 1.6 mJ/cm^2$.

For negative delay times $t_D \leq -60 fs$, the transmission spectra are dominated by a single eigenmode at $\nu = 29 THz$, characteristic of the unpumped state. When the the pump
and probe pulses coincide at $t_D = 0$ fs, an abrupt qualitative change of the transmission spectrum manifests itself. Instead of the bare eigenmode, two resonances emerge: a strong peak centered at $\omega_L/2\pi = 28.2$ THz and a weaker one at $\omega_U/2\pi = 30.5$ THz, corresponding to the lower and upper polariton branch, respectively. Intriguingly, the upper branch exhibits a lower amplitude than the lower branch, it also remains spectrally much broader during the initial time window $0 \leq t_D \leq 340$ fs. The observed ultrafast build-up dynamics of the polariton bands stands in stark contrast to an adiabatic increase of light-matter interaction, as observed for a gradual increase of $\rho_{1e}$ (compare Figure 5.8). At $t_D = 340$ fs, a maximum splitting of $2\Omega_R/2\pi = 3.4$ THz is reached. This is as much as 12% of $\nu_{2SB}$ exceeding even values demonstrated with static doping [Zan10]. At this point, both polariton branches are comparable in intensity and width. Between $t_D = 5$ ps and 1 ns, the vacuum Rabi frequency reduces slightly by 10% due to interband carrier recombination.

The surprising and consistently reproducible asymmetric buildup of the polaritons stands in contrast to the dynamics seen in planar waveguides [Gün09]. This qualitative discrepancy clearly indicates that the peculiar band structure of the photonic crystal sets a characteristic switch-on time for the formation of ultrastrong light-matter coupling. Due to energy-time uncertainty, abrupt switching events have to involve a broad spectrum of
5.4. Nonadiabatic switching of a photonic bandstructure

eigenmodes. Contrary to planar waveguides, this spectrum is highly structured in the case of a photonic crystal. The simultaneous coherent interaction of intersubband transitions with higher photonic bands may lead to interference effects during the activation of ultrastrong coupling. It may be tentatively suggested [Por12] that nonadiabatic mixing of the upper polariton branch with the next higher photonic band plays a dominant role. A quantitative model of this fascinating new class of ultrafast quantum phenomena calls for a universal theory treating nonadiabatic quantum electrodynamics and quantum kinetic carrier dynamics, on equal footing.

In conclusion, non-adiabatic optical activation of ultrastrong light-matter interaction between a custom tailored plasmon assisted photonic mode and an intersubband transition of quantum wells is implemented. By means of sub-cycle multi-THz spectroscopy, the relevant photon and polariton modes of the novel device are fully mapped out in straightforward transmission geometry and their non-adiabatic switching dynamics is recorded on a femtosecond timescale. The novel light-matter coupling device presented in this work enhances the design introduced by Günter et al. [Gün09] by enabling full and simplified access to the polaritonic modes. Since Casimir photons are expected to emerge as excitations of the polaritonic modes upon non-adiabatic activation of ultrastrong light-matter interaction, the enhanced access to the polariton modes should help facilitate the detection of Casimir photons in future experiments based on shot-noise reduced ultrasensitive electro-optic sampling.
Summary and outlook

The objective of this thesis is to elucidate the ultrafast dynamics of coupled low-energy excitations in both strongly correlated materials and artificially engineered quantum structures. With the aid of NIR pump/multi-THz probe spectroscopy and a series of technological innovations such as a novel collinear four-pass Ti:Sapphire amplifier and shot-noise reduced electro-optic sampling [Por14b], fundamentally new insights into the many-body physics of two representative strongly correlated materials are obtained:

Two coupled order parameters associated with the charge density wave in the transition-metal dichalcogenide $1T$-TiSe$_2$ are transiently separated on a femtosecond scale, for the first time [Por14a]. Following intense photoexcitation with a 12-fs near-infrared laser pulse, a novel non-thermal phase is observed in which the structural component of the charge density wave persists without exciton-like electronic order. This proves, that excitonic order is not the sole cause for the formation of a charge density wave in this material, solving a long standing mystery. The comprehensive femtosecond study of the far- to mid-infrared spectral fingerprints of electronic and lattice order presented in this work allows for a consistent identification of the elementary microscopic mechanisms underlying the formation of a charge density wave in this material: The data provide strong evidence for the joint action of excitonic electron-hole correlations and a structural Jahn-Teller-like effect to cause the charge density wave transition. This interpretation is backed up by a quantum-mechanical mean-field model, which is extended to describe the influence of photoexcitation on the excitonic and structural degrees of freedom. The results presented in this work may also serve as a benchmark to quantitatively test and advance recent three dimensional models of the charge density wave phase transition in $1T$-TiSe$_2$ [Zen13]. Furthermore, the refined picture of the many-body effects in $1T$-TiSe$_2$ may ultimately assist the identification of the mechanism that leads to (unconventional) superconductivity in...
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1T-TiSe\textsubscript{2} [Mor06, Kus09] and possibly even other strongly correlated systems. From the experimental point of view, comparable studies in superconducting Cu\textsubscript{x}TiSe\textsubscript{2} could further enhance the fundamental understanding of the interplay between charge ordered and superconducting phases in strongly correlated materials [Mor06, Li07a, Kis07, Cha12]. The availability of high-quality monolayers of transition-metal dichalcogenides [Wan12] paves the way for NIR-pump/THz-probe experiments to target the many-body physics also in these systems. For example, room temperature excitons in monolayered WSe\textsubscript{2} [Coe87] could be investigated via the excitons’ internal degrees of freedom. With the introduction of silver thiogallate as THz emitter and detector reported in this work, the relevant spectral region that is expected to host internal transitions of the room temperature excitons (expected at ∼ 40 THz [He14a]) is now accessible with the experimental tools employed in this work. In general, the concept to disentangle the ultrafast dynamics of structural and electronic components of an order parameter by probing back-folded phonon branches together with a purely electronic response should be universally applicable to a broad variety of other strongly correlated materials.

Furthermore, this work presents novel insights into the microscopic interaction processes in one of the most intensely studied cuprate superconductors. This experiment represents the first resonant femtosecond observation of both electronic and phononic degrees in a high-temperature superconductor [Pas10]. By recording a photoinduced superconducting-to-normal phase transition in YBa\textsubscript{2}Cu\textsubscript{3}O\textsubscript{6.93} by means of femtosecond multi-THz spectroscopy, a qualitatively new insight into the electron-phonon interaction in this system is obtained. The lattice absorbs a large portion of the pump pulse energy while the photoexcited charge carriers thermalize and the condensate is depleted. This observation demonstrates, that there is no temporal hierarchy of electron-electron and electron-phonon scattering processes in this material. As a consequence, the applicability of two- or three temperature models [Kab08, Per07], which associate different timescales to the fundamental electron-phonon and electron-electron scattering mechanisms, can be ruled out for this case. Future experiments may target the novel class of iron-based pnictide superconductors in analogous studies. Such experiments could verify the current thinking that electron-phonon interaction does not play a primary role in the mechanism that yields superconductivity in these systems [Ste11]. Furthermore, some pnictide superconductors feature two simultaneously present superconducting gaps [Ste11]. Femtosecond THz spectroscopy appears to be ideally suited to disentangle these gaps via their intrinsic recovery dynamics and may provide valuable clues on their microscopic origins.

The last part of the work demonstrates full spatial and non-adiabatic temporal control of low-energy light-matter interaction in the limit of ultrastrong coupling [Por12]. The regime of ultrastrong light-matter coupling is of utmost interest for fundamental quan-
tum electrodynamics since it provides a unique test bed for intriguing quantum theoretical predictions. Thereof, the non-classical generation of photons out of the quantum-vacuum described by the dynamical Casimir effect [Ciu05, Lib07], which is related to Unruh-Hawking radiation of black holes [Yab89], is the most prominent example. This work takes a further step towards the observation of Casimir photons in intersubband cavity polariton systems [Gün09, Zan10]. It introduces a novel device that enables straightforward access to the photonic modes which are expected to be populated with Casimir photons upon non-adiabatic switching of ultrastrong light-matter interaction. The device features a refined control of ultrastrong light matter coupling in all four dimensions which is obtained by combining the spatial mode confinement and well-defined band structure of a photonic crystal with an optically switchable intersubband resonance. Femtosecond multi-THz spectroscopy is employed to demonstrate access to the relevant photonic modes of the device while ultrastrong coupling is activated non-adiabatically. The observed asymmetric formation dynamics of the light-matter mixed polariton modes calls for a novel theory accounting for both quantum kinetic carrier dynamics and non-adiabatic quantum-electrodynamics. Future experiments will primarily aim for the observation of the dynamical Casimir effect in this system. Thereby, the concept of shot-noise reduced electro-optic sampling developed in this work may prove helpful [Por14b].

In this work, NIR pump/multi-THz-probe spectroscopy has enabled the study of ultrafast low-energy dynamics of fundamentally different systems – strongly correlated materials and artificially engineered quantum structures. Recent advances in THz technology pave the way for entirely new types of experiments and can be anticipated to play a similarly outstanding role in the near future. High-field THz sources [Sel08a, Sch14] have been developed with the goal to not only study the low-energy degrees of freedom but to also coherently control them via both the electric and the magnetic fields of phase-stable THz radiation. This approach has already been successfully applied for the first observation of electric field induced dynamical Bloch-oscillations in a bulk semiconductor [Sch14] and may bring the observation of novel phenomena such as THz-magnetic-field-controlled spin switching into reach [Kam11]. Field resolved access to the oscillating THz near-field of nanostructures has become accessible recently with the novel technique of ultrafast near-field multi-THz nano-spectroscopy [Eis14]. One can envision to directly monitor how wavefunctions associated with eigenstates of nanostructures evolve on a femtosecond scale [Jac12] or to observe ultrafast formation processes of ordered domains in strongly correlated materials [Qaz07], just to name a few promising perspectives.
Appendix A

Raw data of a NIR-pump/THz-probe experiment

Figure A.1: Raw data of a NIR-pump/THz-probe experiment, corresponding to datasets shown in Figures 3.8, 3.9 and 3.10. (a) Electric field trace of single-cycle THz transients measured after transmission through an unexcited 80 nm thin film of 1T-TiSe2. (b) Photoinduced change of the transmitted THz electric field. The temperature of the sample is kept at $T = 10$ K and the absorbed pump fluence is $\phi = 40 \mu J/cm^2$. This 2D dataset was acquired in less than four hours. Note the excellent long term stability of the reference transients throughout the measurement.
Appendix B

Numerical analysis of the dielectric response of 1T-TiSe$_2$: Quality of the fits

As described in section 3.3.2, the transient change in the free carrier density $\Delta n(t_D)$ and the transient free carrier scattering time $\tau(t_D)$ of 1T-TiSe$_2$ after photoexcitation are extracted from the time dependent energy loss function by numerical adaption of the model function given by equation (3.1). Thereby, both the real and imaginary part of $1/\varepsilon(\hbar\omega,t_D)$ are fitted simultaneously, with high confidence. Figure B.1 compares the measured (left panels) and fitted spectra (right panels) of a 2D dataset acquired with a pump fluence of $\phi = 40 \mu J/cm^2$ at $T = 10 K$ to highlight the quality of the fit curves.

**Figure B.1:** Left panels: Imaginary and real part of $1/\varepsilon(\hbar\omega,t_D)$ of photoexcited 1T-TiSe$_2$ measured at $T = 10 K$ shown for a representatively selected pump fluence of $\phi = 40 \mu J/cm^2$. Right panels: Corresponding spectra of the numerically adapted model function (equation (3.1)) used to extract $n(t_D)$ and $\tau(t_D)$.
Extended analysis of the transient dielectric response of 1T-TiSe$_2$

As discussed in section 3.6, the multi-THz probe covers part of the interband transition across the CDW gap of 1T-TiSe$_2$. In this spectral region, coherent oscillations of the absorption coefficient are observed after NIR photoexcitation. To further corroborate the interpretation that the coherent dynamics reflects a modulation of the Jahn-Teller component of the CDW gap, the transient properties of the single-particle interband transition are studied quantitatively as far as possible. To this end, additional parameters of the model dielectric function (equation 3.1) are adapted to the measured dielectric response: The center energy $E_{\text{gap}}$ and the damping $\gamma_{\text{gap}}$ of the interband transition across the CDW gap. Even though only a small fraction of the resonance is captured, numerical adaption of four fit parameters proves possible for small pump induced changes at low pump fluence ($\phi \leq 20\,\mu J/cm^2$). Since numerical fitting of $1/\varepsilon$ strongly suppresses numerical weight of the single particle transition and allows for reliable extraction $n$ and $\tau$, the numerical fit shown here is performed via adaption of the dielectric function $\varepsilon$. To further extend the numerical weight of the interband transition, the energy window for the fit is extended to 150 meV. For convergence of the fitting algorithm, it is necessary to keep the spectral weight of the interband resonance fixed, which can be considered legitimate for low $\Phi$.

Figure C.1 shows the transient fit parameters extracted from $\varepsilon(\hbar\omega,t_D)$ measured with $\Phi = 20\,\mu J/cm^2$ at $T = 10\,\text{K}$. The transient free carrier density $n$ (a) and free carrier scattering time $\tau$ (b) reproduce the results shown in figure 3.9. Panels (c) and (d) additionally show the transient damping $\gamma_{\text{gap}}$ and center energy $E_{\text{gap}}$ of the interband transition across the CDW gap. The pump-induced change in $\gamma_{\text{gap}}$ mimics the change in $n$, suggesting that a broadening of the interband resonance is governed by the presence of photogenerated free carriers. The exponential decay dynamics of $E_{\text{gap}}$ is essentially the same as observed for $\tau$. Notably, the coherent oscillations of the absorption coefficient are entirely captured.
Figure C.1: Extended quantitative analysis of the transient dielectric response measured after photoexcitation with $\Phi = 20 \mu \text{J/cm}^2$ at $T = 10 \text{K}$. (a) and (b) show the Drude parameters $n$ and $\tau$ describing the plasma response. (c) and (d) show the damping $\gamma_{\text{gap}}$ and center energy $E_{\text{gap}}$ of the interband transition across the CDW gap. Dashed lines show an exponential decay numerically adapted to the measured data.

by $E_{\text{gap}}$. Taken together, these observations fully corroborate the picture of combined action of excitonic and Jahn-Teller like mechanisms in the CDW formation and clearly indicate that the CDW gap is jointly formed by both contributions: On the one hand, the exponential recovery dynamics observed in both $\tau$ and $E_{\text{gap}}$ can be associated with the recovery of the excitonic component of the CDW order. On the other hand, the oscillations superimposed to the exponential recovery of $E_{\text{gap}}$ reflect modulations of the Jahn-Teller like energetic lowering of the Se4p bands caused by coherent oscillations of the $A_{1g}$ amplitude mode. In fact, the modulation of $E_{\text{gap}}$ due to Jahn-Teller effects is reproduced by the theoretical model (section 3.7).
Appendix D

The model phase diagram of 1T-TiSe$_2$: Constraints for $\rho$, $\tau_{in}$ and $\tau_{out}$

As discussed in section 3.7, minimization of the total energy described by the mean-field model Hamiltonian (equation (3.8)) with the mean-fields $\rho$, $\tau_{in}$, $\tau_{out}$ and $u$ as free parameters yields the zero temperature model phase diagram. The parameters $\tau_{in}$ and $\tau_{out}$ are complex-valued, which, together with $\rho$, implies a total of five free parameters that describe the electronic degree of freedom.

To reproduce the results by van Wezel et al. [Wez10b], it turns out necessary to constrain $\rho$, $\tau_{in}$ and $\tau_{out}$ via conservation of the total particle number and translational invariance along the one-dimensional chain.

The following analysis is based on a general fermionic state describing the electronic degree of freedom of two neighboring unit cells with always two electrons present:

$$|\Psi_1\rangle = \alpha|1010\rangle + \beta|1001\rangle + \gamma|1100\rangle + \delta|0101\rangle + \epsilon|0110\rangle + \zeta|0011\rangle \quad(D.1)$$

The electron at the first position (Se orbital in unit cell $i$) is created with $c_1^\dagger$ the second (Ti orbital in unit cell $i$) with $d_1^\dagger$, the third (Se orbital in unit cell $i+1$) with $c_1^\dagger$ and the fourth (Ti orbital in unit cell $i+1$) with $d_1^\dagger$.

From translational invariance, the following constraints apply:

$$\langle\Psi|c^\dagger c|\Psi\rangle = \langle\Psi|c_1^\dagger c_1|\Psi\rangle \quad(D.2)$$
$$\langle\Psi|d^\dagger d|\Psi\rangle = \langle\Psi|d_1^\dagger d_1|\Psi\rangle \quad(D.3)$$

Following parametrization of $|\Psi\rangle$ fulfills these conditions:

$$|\Psi_2\rangle = \alpha|1010\rangle + \beta|1001\rangle + \gamma|1100\rangle + \delta|0101\rangle + \beta|0110\rangle + \gamma|0011\rangle \quad(D.4)$$
Again, from translational invariance, the phases of the coefficients can be either 0 or $\pi$. Taking into account that the state has to be normalized, three free parameters are left. The physically relevant choices of the phases are captured by the signs of the coefficients.

To obtain a parametrization that ensures normalization and is given by only three parameters, the state is described with real valued $x$, $y$ and $z$ as follows:

$$|\Psi\rangle = \cos\left(\frac{x}{2}\right) \cos\left(\frac{y}{2}\right) |1010\rangle + \frac{1}{\sqrt{2}} \left[ \sin\left(\frac{x}{2}\right) \left( \cos\left(\frac{y}{2}\right) \cos (z) + \sin\left(\frac{y}{2}\right) \sin (z) \right) \right] (|1001\rangle + |0110\rangle) + \frac{1}{\sqrt{2}} \left[ \sin\left(\frac{x}{2}\right) \left( \sin\left(\frac{y}{2}\right) \cos (z) - \cos\left(\frac{y}{2}\right) \sin (z) \right) \right] (|1100\rangle + |0011\rangle) + \cos\left(\frac{x}{2}\right) \sin\left(\frac{y}{2}\right) |0101\rangle$$

(D.5)

With this parametrization, $\rho$, $\tau_{\text{in}}$ and $\tau_{\text{out}}$ are:

$$\rho = \langle \Psi | c^\dagger c | \Psi \rangle - \langle \Psi | d^\dagger d | \Psi \rangle = \frac{1}{4} \cos (x - y) + \frac{1}{4} \cos (x + y) + \frac{1}{2} \cos (y)$$

(D.6)

$$\tau_{\text{in}} = \langle \Psi | c^\dagger d | \Psi \rangle = \frac{\sqrt{2}}{2} \cos\left(\frac{x}{2}\right) \sin\left(\frac{x}{2}\right) \cdot \left( \cos\left(\frac{y}{2}\right) + \sin\left(\frac{y}{2}\right) \right) \left( \cos\left(\frac{y}{2}\right) \cos (z) + \sin\left(\frac{y}{2}\right) \sin (z) \right)$$

(D.7)

$$\tau_{\text{out}} = \langle \Psi | c^\dagger d_1 | \Psi \rangle = -\frac{\sqrt{2}}{2} \cos\left(\frac{x}{2}\right) \sin\left(\frac{x}{2}\right) \cdot \left( \cos\left(\frac{y}{2}\right) + \sin\left(\frac{y}{2}\right) \right) \left( \sin\left(\frac{y}{2}\right) \cos (z) - \cos\left(\frac{y}{2}\right) \sin (z) \right)$$

(D.8)

Now, the parameters $x$, $y$ and $z$ substitute $\rho$, $\tau_{\text{in}}$ and $\tau_{\text{out}}$ upon minimization of the total energy. The values of the mean-field parameters $\rho$, $\tau_{\text{in}}$ and $\tau_{\text{out}}$ for minimum energy are then calculated with the above equations from the values of $x$, $y$ and $z$ for which the total energy is minimized.
Appendix E

Slow-down of light induced by nonadiabatic activation of ultrastrong light-matter coupling

The extremely strong light-matter coupling in the light-matter coupling device gives rise to a slow-down of the group velocity $d\omega/dk\parallel$ of its guided modes. As a quantitative measure, the effective group index of refraction $n_{\text{eff}}$ is extracted as a function of $k\parallel$ (Figure E.1) from a numerically adapted polariton dispersion curve [Hop58] on the experimental band structure (compare Figure 5.5), without resorting to a rotating wave approximation. While $n_{\text{eff}}$ reflects the theoretical value of 3.15 in the unpumped sample, the extracted $n_{\text{eff}}$ increases for the lower polariton branch to 41 at the highest $k\parallel$-values covered in the experiment (vertical dashed line in Figure E.1). A coherent photon population which is initially prepared in the bare grating waveguide by the multi-THz transients is, thus, abruptly transferred into a polaritonic band decreasing its group velocity by a factor of 13. Slow light reaching speeds down to only few tens of meters per second has been observed with a variety experimental methods [Big03, Ku04, Tur01]. Most of these techniques suffer from a limited bandwidth that hinders their practical applications. In this context, the present experimental scheme holds great potential because of its ability to slow down

Figure E.1:
Effective refractive index $n_{\text{eff}}$ at the upper photonic modes in the pumped and unpumped sample, as represented by the black curves in Figure 5.5. The solid curve is the theoretical $n_{\text{eff}} = 3.15$ of the light-mode propagating inside the unpumped quantum well region.
light over a bandwidth corresponding to a significant fraction of the carrier frequency. In this regards, the results presented here compare advantageously with state-of-the-art broadband generation of slow light in photonic crystal structures [Vla05, Set07]. The demonstrated control of the band structure may ultimately enable a broadband storage of light by transferring a coherent photon population from a leaky equilibrium photoplasmonic band to a non leaky polariton branch. Additionally, this scheme provides a novel possibility to investigate ultrafast switching of the slow light regime. One may even speculate that the presented concept may be employed to implement a spatially moving refractive index perturbation to generate an artificial event horizon that yields the emission of Hawking radiation, analogous to an experiment performed by Belgiorno et al. [Bel10].
Establishment of the femtosecond multi-THz laboratory

This work included the establishment of the femtosecond multi-THz laboratory at the Institute of Physics of the University of Regensburg. Parts of the NIR-pump/multi-THz-probe spectroscopy system (the cavity dumped Ti:sapphire oscillator and a series of mechanical actuators and optical components) were transferred from the University of Konstanz (Emmy Noether Group “Tera” of R. Huber at the chair of A. Leitenstorfer), where the experiments on YBa$_2$Cu$_3$O$_{6.93}$ were conducted. Building the experimental setup from scratch (compare Figure F.1 on page 112) provided the possibility to design the supporting infrastructure in the laboratory to facilitate long-term stability for high sensitivity experiments (compare Figure A.1 in appendix A). An air conditioning system stabilizes temperature and humidity in the laboratory (temperature stability: ±0.1 °C). Special care was taken to isolate the experimental setup from acoustic vibrations of the environment. To this end, the setup is placed on a vibration damped standard optical table. Acoustic noise induced by supporting hardware such as chillers and power supplies is minimized by placing these devices in a separate service room.

The optical table was installed in May 2011. First phase stable multi-THz pulses were generated and electro-optically detected in October 2011. Figure F.1(b) shows a current snapshot of the laboratory.
Figure F.1:

Impressions of the femtosecond multi-THz laboratory.  (a) Refurbished basement room in early 2011 designated to host the femtosecond multi-THz laboratory (picture by Martin Furthmeier).  (b) Laboratory in the present state.  (c) Close-up view of part of the experimental setup (Ti:sapphire four-pass amplifier).
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