Femtosecond time-resolved optical pump-probe spectroscopy at kilohertz-scan-rates over nanosecond-time-delays without mechanical delay line

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We demonstrate a technique for femtosecond time-resolved optical pump-probe spectroscopy that allows to scan over a nanosecond time delay at a kilohertz scan rate without mechanical delay line. Two mode-locked femtosecond lasers with approximately 1 GHz repetition rate are linked at a fixed difference frequency of $\Delta f_R = 11$ kHz. One laser delivers the pump pulses, the other provides the probe pulses. The relative time delay is linearly ramped between zero and the inverse laser repetition frequency at a rate $\Delta f_R$, enabling high-speed scanning over a 1 ns time delay. The advantages of this method for all-optical pump-probe experiments become evident in an observation of coherent acoustic phonons in a semiconductor superlattice via transient reflectivity changes. A detection shot-noise limited signal resolution of $7 \times 10^{-8}$ is obtained with a total measurement time of 250 s. The time resolution is 230 fs. © 2006 American Institute of Physics. [DOI: 10.1063/1.2167812]

Experiments in the field of ultrafast time-domain spectroscopy most commonly employ trains of pump and probe pulses derived from a single mode-locked laser. One pulse train travels a variable distance before arriving at the sample and data are recorded as a function of the time delay versus the other pulse train. In most cases the time delay is adjusted with mirrors mounted onto a mechanical translation stage or a vibrating membrane. Translation stages allow for several hundred picosecond delays but not for rapid scanning. Vibrating membranes allow for higher scan rates in the range of 100 Hz but are limited to approximately 10 ps time delay. A severe disadvantage of moving mirrors in general is the fact that it is impossible to eliminate residual variations of the beam pointing as time delays of several hundred picoseconds are realized by moving the stage by tens of centimeters. Also, spot size variations at the sample position are inherent due to the divergence of the laser beam. These effects are greatly unfavorable when phenomena with long decay and/or dephasing times are investigated over nanosecond time delays due to an unresolvable ambiguity between real sample dynamics and experimental artifacts. Examples of physical phenomena that require the detection of sample dynamics over nanosecond time delays are picosecond ultrasound pulses, transient spin-dynamics, long-living coherent lattice or charge carrier dynamics in solid-state high-resolution terahertz spectroscopy, terahertz ranging, or the characterization of high-speed optoelectronic devices. Furthermore, increased scan rates are desirable for imaging applications to reduce pixel dwell times or for experiments where data have to be acquired in a short period of time (e.g., pulsed magnetic fields).

Asynchronous optical sampling (ASOPS) is a technique demonstrated earlier with mode-locked picosecond lasers at low repetition rates (82 MHz) that enables ultrafast time-domain spectroscopy without a mechanical delay line. To this end, two mode-locked lasers are linked at a fixed repetition rate difference $\Delta f_R$, and serve as pump and probe lasers. Their relative time delay is linearly ramped between zero and the inverse repetition rate of the pump laser at a rate given by $\Delta f_R$. Our high-speed implementation of ASOPS operates with mode-locked Ti:sapphire femtosecond lasers at 1 GHz repetition rate and thus allows kilohertz scan rates over a temporal measurement window of 1 ns with femtosecond time resolution. High repetition rates in the range of 1 GHz are generally favorable for high-speed ASOPS because they reduce the required optical detection and data acquisition bandwidth $F$ to practically accessible values in the range of 100 MHz. The feasibility of high-speed ASOPS has recently been shown in the field of terahertz time-domain spectroscopy. However, the time resolution had been limited to 2.5 ps due to limitations of the laser stabilization and a restricted bandwidth of the detection system.

In this letter, we report the application of high-speed ASOPS to all-optical femtosecond time-resolved pump-probe spectroscopy. As an important prerequisite, we have significantly improved and simplified the electronics used to stabilize the laser repetition rate difference. To demonstrate the utility of the technique, we have chosen a GaAs/AlAs semiconductor superlattice sample as a challenging model system. The response of such structures to impulsive optical excitation is well known from earlier experiments performed conventionally with mechanical delay lines: Coherent longitudinal acoustic phonons with periods on the order of 1 ps and lifetimes exceeding 100 ps are observable through weak transient reflectivity changes.

We employ a dual-laser system (Gigajet TWIN, Gigaoptics GmbH, Germany) which comprises two Ti:sapphire femtosecond oscillators with repetition rates $f_{R,1}$ (laser 1) and $f_{R,2}$ (laser 2) of approximately 1 GHz. The oscillators are pumped by a diode-pumped solid-state laser at 532 nm wavelength whose 6.5 W output power is split into equal...
portions. The oscillators deliver approximately 500 mW of output power at a center wavelength of 820 nm with a pulse duration of approximately 30 fs. The repetition rate difference $\Delta f_{R} = f_{R,1} - f_{R,2}$ is fixed using active feedback. The time delay between the two pulse trains is then $\tau(t) = t \times \Delta f_{R}/f_{R,2}$ modulo $f_{R,2}^{-1}$ if laser 2 is used as the pump laser and $t$ is the real time. The quality of the feedback electronics determines the accuracy of the $\tau(t)$ relationship given above through a jitter of the time base which directly determines the experimental time resolution. Instead of employing two feedback loops to stabilize the lasers to their individual reference synthesizers with an offset of $\Delta f_{R}$, we use laser 1 as master laser and link laser 2 as slave laser to it at a difference of $\Delta f_{R} = 11$ kHz. This has the advantage that only the residual errors of one feedback loop enter the jitter of the time-base.

The experimental scheme is sketched in Fig. 1(a). We measure the $f_{R,1}$ by detecting laser 1 with a fast photodiode (PD1) and upshift the frequency of the electronic signal by 11 kHz using a frequency shifter. The frequency shifter consists of a single-sideband generator with a carrier suppression stage that allows a suppression of the initial carrier signal by better than 50 dB. Subsequently, $f_{R,2}$ is detected with a second photodiode (PD2) and phase locked to the frequency-shifted repetition rate signal of laser 1. To accomplish this, we use a double-balanced mixer (DBM) as phase detector and feed its output back to a piezoelectric transducer that supports a cavity mirror in laser 2 via an adjustable loop amplifier. The jitter of the time-base $\tau(t)$ is characterized by detecting a cross-correlation signal between the two lasers using noncollinear sum-frequency generation in a $\beta$-barium-borate crystal. The cross-correlation signal is detected with a 125-MHz bandwidth photoreceiver and recorded using a sampling oscilloscope (Tektronix TDS3032B). To generate a trigger signal for the oscilloscope at $\Delta f_{R}$, we use a second DBM with the signals from PD1 and PD2 as inputs. The real-time scale is converted to a time-delay scale by applying a factor $\Delta f_{R}/f_{R,2}$. Figure 1(b) shows a single scan measurement of the cross-correlation signal and an average over 512 measurements acquired over $\approx 1$ s, limited by the averaging function of the oscilloscope. The single scan trace with a rise time of 95 fs shows the limitation to the time-resolution $\Delta \tau$ given by the 100 MHz bandwidth of the sampling oscilloscope (determined by our specific chosen acquisition settings) according to $\Delta \tau = \Delta f_{R}/f_{R,2} F$, where $F$ is the effective bandwidth of the photodetection and data acquisition electronics. The detailed structure is attributed to the response of the photoreceiver. The averaged cross-correlation signal has a width of 230 fs, reflecting residual jitter of the time base. Averaging over longer times does not further broaden the signal, thus 230 fs represents the overall time resolution of our setup. This limit is mostly due to phase-noise intrinsic to the DBMs used for phase detection and trigger generation as well as a residual contribution at the carrier frequency at the output of the frequency shifter.

We envision that an improvement of the timing stability to the sub-100 fs regime is possible by performing the phase lock at a higher harmonic of the repetition rate or by using digital phase detectors.

All-optical pump-probe spectroscopy is performed on a GaAs/AlAs semiconductor superlattice sample with 40 periods of 18 atomic monolayers of GaAs and 18 monolayers of AlAs. The pump and probe beams are focused onto the same spot on the sample using a lens with a focal length of 35 mm. The spot size is approximately 30 $\mu$m. The pump and probe powers are 400 and 10 mW, respectively. The excitation wavelength is close to the interband transition between the first quantized electron and hole states. After reflection off the sample, the probe beam is focused onto the 125-MHz bandwidth photoreceiver; reference detection is not used. The signal is recorded with a 200 Ms/s 14 Bit analog-to-digital converter (ADC) on a PCI board (Compuscope 14200, GaGe Applied Technologies) that is triggered by the synchronization electronics. The data acquisition board has the capability of acquiring hardware-based averages over 1024 single scan traces. These averaged traces are displayed at a rate of 4 Hz on the computer. An arbitrary number of additional software based averages can be computed. The spatial overlap of pump and probe pulse can be optimized “on-line” by observing the signal transient on our data acquisition computer. An average over 1024x1000 single traces with a total acquisition time of 250 s is shown in Fig. 2(a). Data are calibrated to show relative reflectivity changes $\Delta R/R_0$, where $R_0$ is the unperturbed reflectivity.

The signal exhibits a rise in reflectivity of $1.5 \times 10^{-3}$ due to optically excited charge carriers in the superlattice followed by a decay due to relaxation, recombination, and diffusion. Sufficient signal-to-noise ratio to obtain the electronic dynamics can already be achieved without software averaging. Superimposed on the electronic signature is a modulation of the reflectivity with an amplitude of $\approx 2 \times 10^{-4}$ due to impulsively driven coherent acoustic phonons. Their oscillatory contribution has been extracted by subtracting a fit to the slow electronic dynamics from the original data and is shown in Fig. 2(b). A fast-Fourier-transformation (FFT) of the data is shown in Fig. 2(d) (left panel), revealing two dominating modes at 41.8 and 44.5 GHz. These frequencies agree well with those of propagating unfolded longitudinal acoustic phonons driven in the individual GaAs and AlAs portions of the superlattice.
phonons that cannot be graphically resolved on the 1 ns scale in Fig. 2(b). This contribution has therefore been extracted as above and is shown in Fig. 2(c). The FFT [Fig. 2(d), right panel] shows three modes that are attributed to a nonpropagating zone-folded acoustic phonon mode at 469 GHz and two propagating zone-folded acoustic phonon modes with 432 and 519 GHz. A detailed statistical analysis of data acquired at different numbers of software averages N shows that the signals are shot-noise limited with a noise dependence \(\propto \sqrt{N}\). At \(N = 1000\), the resolution for \(\Delta R/R_0\) is \(7 \times 10^{-8}\). In order to acquire equivalent data with an earlier demonstrated method based on a vibrating membrane, a total data acquisition time of \(\approx 10,000\) s would have been required. It is important to note, however, that this method would not have been capable to resolve the low frequency excitations around 45 GHz.

The presented data qualify high-speed ASOPS as a technique for femtosecond time-resolved optical pump-probe spectroscopy that is comparable to conventional methods in terms of time resolution and signal-to-noise ratio and is largely superior in a number of other respects: It allows for the elimination of mechanical delay lines and thus variations of beam pointing and spot size as well as acoustic noise due to mechanical movement. Furthermore, we have demonstrated a scan rate of 11 kHz over a measurement window of 1 ns with 230 fs time resolution and obtained a signal resolution of \(7 \times 10^{-8}\) within a total sampling window of 250 s. This is an improvement by more than one order of magnitude compared to the time required for mechanical delay scanning.

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