Far-infrared picosecond time-resolved measurement of the free-induction decay in GaAs:Si

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By measuring changes in the photoconductivity induced by picosecond far-infrared pulse pairs from a free-electron laser, we have time resolved the free-induction decay of the 1s-210 and 1s-2p\textsuperscript{+} Si-shallow-donor transitions in bulk GaAs. The method frees us from the problem of measuring the optical emission of the transitions and allows us to obtain their dephasing times. We expect to be able to use the same method in the future to measure other coherent phenomena in these systems, such as photon echoes.

I. INTRODUCTION

In optical experiments in gases, liquids, and solids, ultrashort visible or near-IR laser pulses are commonly used to probe the dynamics of the system under study. Examples are free-induction decay measurements to obtain optical dephasing times of transitions, pump-probe techniques to measure excited-state lifetimes, and four-wave mixing to measure phonon echoes.\textsuperscript{1,2} When transitions have frequencies in the far-infrared region, it is sometimes possible to probe their dynamics with ultrashort visible pulses using nonlinear techniques such as Raman scattering.\textsuperscript{2} In many cases, however, the use of visible pulses also creates many undesirable simultaneous excitations such as the creation of electron-hole pairs by interband excitation. Worse, the high-intensity visible pulses usually needed for these nonlinear time-resolved experiments may cause irreversible damage to the material under study. As a result, direct excitation with far-infrared lasers is needed. However, time-resolved studies of the dynamics of these transitions with ultrashort pulses were hampered by the absence of suitable ultrashort-pulse far-infrared laser sources.\textsuperscript{3}

In this paper, we use picosecond, far-infrared pulses from the free-electron laser for infrared experiments (FELIX) at the FOM Institute Rijnhuizen, to time resolve the free-induction decay of the 1s-2p\textsuperscript{+} and the 1s-210 silicon (Si) shallow-donor transitions in GaAs. We avoid the problem of detecting the emitted far-infrared radiation by using a detection method based on the photoconductive response\textsuperscript{4-6} of the sample. Two pulses, separated in time, excite the system. We find that the second, delayed pulse induces changes in the photoconductivity of the sample that depend on the optical phase difference between the two pulses. We prove that these phase-difference-dependent changes reflect changes in the excited-state population of the Si donors, caused by the coherent manipulation of the second pulse of the optical polarization stored in the donor system by the first. By measuring how these coherent population changes decrease in magnitude for increasing pulse separations, we can obtain the picosecond free-induction decay time constant of the optical polarization of the Si donor. Thus, instead of a (difficult) time-resolved measurement of the far-infrared optical emission of the free-induction decay, we do a time-resolved measurement of the decay of the coherence that is the source of the emission. We expect that this method can also be used to measure various other coherent phenomena such as photon echoes.

II. EXPERIMENT

Our sample, obtained from Philips Research Laboratory, Redhill, U.K., is a 10-\mu m-thick Si-doped GaAs layer grown on a 400-\mu m-thick semi-insulating GaAs crystal. The doping density \(N_d\) is approximately \(5 \times 10^{14} \text{ cm}^{-3}\), and \(N_d \leq 1 \times 10^{14} \text{ cm}^{-3}\). Sn contacts have been diffused in for conductivity measurements. The experiments are done at 8 K in a magnet cryostat in magnetic fields up to 9 T. The magnetic field can be adjusted to tune the various donor transitions into resonance with a particular far-infrared frequency.

Tunable far-infrared pulses are generated by the FELIX. The light emerges from the FELIX in the form of pulse trains consisting of approximately 5000 micro-pulses. The pulse trains are generated at a 5-Hz repetition rate. The micropulses have an adjustable duration of a few picoseconds and are separated by a nanosecond. After passing through an attenuator, used to avoid saturation of the transitions, they are sent into a Michelson interferometer which contains a 6-\mu m-thick Mylar beamsplitter. One arm of the Michelson can be scanned with respect to the other so that the pulse pairs emerging from the Michelson have an adjustable time separation. The pulse pairs are focused onto the sample with a 50-cm focal-length polyethylene lens, and the radiation enters the cryostat through two polyethylene windows. Care is taken not to illuminate the contacts on the sample to avoid spurious signals. The magnetic field is perpendicular to the surface of the sample and perpendicular to the polarization of the laser pulses. The whole setup is enclosed in a box that is continuously flushed with dry ni-
trogen gas to reduce water vapor absorption. The high-
ohmic sample is voltage biased. The current response
of the sample is measured with a time resolution of 0.1 μs,
and follows the amplitude envelope of the FELIX pulse
train.

III. RESULTS

A. Si-donor optical spectrum

To identify the Si-donor transitions in the sample, we
measure the photoconductivity as a function of magnetic
field for an excitation frequency of 4.45 THz (Fig. 1). The
magnetic field shifts the transition frequencies, and for
fields larger than a few T the shift is linearly propor-
tional to the field. For this measurement, the pulse duration
was lengthened to > 10 ps to decrease the frequency
bandwidth of the pulse while one arm of the Michelson
was blocked. Figure 1 shows peaks in the photoconduc-
tivity whenever a Si-donor transition is resonant with the
excitation pulse. The transitions are visible in photocon-
ductivity measurements because most electrons in the ex-
cited states decay to the conduction band of the GaAs
crystal, where we can use conventional, relatively slow
electronics to detect them. The photoconductivity change
is linearly proportional to the excited-state popu-
lation. The electrons return to the 1s ground state of the
donor atom after roughly 100 ns. To avoid significant
depopulation of the ground state by the cumulative effect
of hundreds of micropulses, we keep the micropulse en-
gy in the following experiments lower than 10 nJ. In the
figure the 1s-2p⁺ and 1s-210 transitions are marked.
They are the two strongest transitions on which we will
focus our attention.

![Photoconductivity vs Magnetic Field](image)

FIG. 1. Measured photoconductivity vs magnetic field for an
excitation frequency of 4.45 THz. In the spectrum, the 1s-2p⁺ and
the 1s-210 transitions discussed in the text are marked.

B. Time-resolved measurements: resonant case

In Fig. 2(a) we plot the photoconductivity of the sample
as a function of the time separation between the two
pulses from the Michelson at an excitation frequency of
4.38 THz (corresponding to a wavelength of 68.8 μm) and
B = 0 T. The photoconductivity oscillates when the time
separation between the two pulses increases, and the os-
cillation period is given by the excitation wavelength.
At this magnetic-field strength, the far-infrared photons
directly excite electrons from the 1s ground state of the Si
donor to a broad continuum of conduction-band states.
As a result, the sample acts as broadband frequency-
dependent detector. The oscillations are explained by
the temporal overlap of the two pulses on the beam-
splitter of the Michelson, and Fig. 2(a) therefore
represents the field autocorrelation of the exciting pulses.
From the autocorrelation we deduce a pulse duration of
approximately 3 ps, assuming bandwidth-limited pulses.

When the laser is resonant with the 1s-2p⁺ transition,
the results are markedly different [Fig. 2(b)]. In contrast
to the results in Fig. 2(a), the oscillations in the photocon-
ductivity (seen more clearly in the inset) are still present
for large time separations when the two pulses have no
temporal overlap on the beamsplitter. The oscillations
eventually decrease in amplitude and, after 50 ps, they
can no longer reliably be measured. In Fig. 2(c) we plot
the photoconductivity of the sample versus time separa-
tion when the laser is resonant with the 1s-210 transition.
Here the measured oscillation amplitude decays much
more quickly for increasing time separation, although

![Oscillation Amplitude vs Time Separation](image)

FIG. 2. Measured photoconductivity as a function of pulse sep-
eparation for three different transitions: 1s continuum (a), 1s-
2p⁺ (b), and the 1s-210 (c). The excitation frequency was 4.38
THz. The inset in (b) shows a small section of the curve, magnified a few times.
significantly more slowly compared to the autocorrelation of the pulses in Fig. 2(a). Clearly, the above results provide evidence that the medium only has a memory for the phase of the exciting electric field when the laser is resonant with a (discrete) transition in the donor system. For pulse energies lower than 10 nJ, we observe no dependence of the decay times on the energy of the micropulses. Above 10 nJ possible energy-dependent effects are obscured by the cumulative depopulation of the ground state by hundreds of micropulses.

C. Time-resolved measurements: nonresonant case

If we detune the transitions with respect to the central wavelength of the pulses, by changing the magnetic field slightly, the results change dramatically. In Fig. 3 we show measurements of the photoconductivity for excitation of the 1s-2p\(^+\) transition for three different values of the detuning. When the laser is exactly resonant with the transition, we obtain the curve already shown in Fig. 2(b) and reproduced in Fig. 3(a). When we now detune the transition 0.08±0.01 THz away from the central wavelength of the laser, we obtain the result in Fig. 3(b). In addition to a decrease in the absolute value (not shown here) of the photoconductivity compared to Fig. 3(a), the signal also has a different shape. After an apparent fast decay of the oscillation amplitude for pulse separations up to 5 ps, a slower decay is seen for larger values of the pulse separation. When we increase the detuning to a value of 0.13±0.01 THz we measure the curve shown in Fig. 3(c). Again the photoconductivity is smaller than in Fig. 3(b) and the initial fast decay of the oscillation amplitude is followed by what seems to be a revival, with a node at \(t \approx 4\) ps. Both in Figs. 3(b) and 3(c), oscillations are still observed for pulse separations of several tens of picoseconds as in Fig. 3(a), with the difference that they are reduced in amplitude.

The oscillations in the photoconductivity shown in Fig. 3 have two contributions. This is clearly demonstrated in Fig. 4, where we plot the Fourier transforms of the signals. A broad peak represents the power spectrum of the laser pulses. Superimposed on this is a much narrower feature that shifts in frequency when the 1s-2p\(^+\) transition is detuned from the central wavelength of the pulses. It corresponds to the oscillations that are visible in Fig. 3 for time separations larger than 5 ps, when the pulses have no temporal overlap. This observation proves that the oscillations in the photoconductivity for pulse separations larger than the pulse duration are indeed caused by a memory effect in the Si-donor transitions.

IV. THEORY

To explain our results, we use a two-level system to model the Si-donor transitions. The oscillations in the photoconductivity as a function of the time separation between the two pulses represent oscillations in the population of the excited states of the Si donor\(^3\) due to the combined effect of the two pulses. When \(\tau\) is larger than the pulse duration, they are explained by the interference
of the electric field of the second pulse with the coherence or optical polarization, stored in the medium by the first pulse. For the complex electric-field amplitude of the exciting pulse pair, we have

$$E_0(t) = A_0(t) + A_0(t - \tau)e^{-i\omega_L \tau}.$$  \hspace{1cm} (1)

Using a density-matrix formulation we can write the following well-known coupled differential equations for the ground-state population $\rho_{11}$, the excited-state population $\rho_{22}$, and the slowly varying coherence amplitude $\rho_{12}$:

$$\frac{\partial \rho_{22}}{\partial t} = \frac{\rho_{22}}{T_{22}} - i \frac{\hbar}{\mu_{12}} (\rho_{21}E_0^* - \rho_{12}E_0) ,$$

$$\frac{\partial \rho_{11}}{\partial t} = -\frac{\partial \rho_{22}}{\partial t} ,$$

$$\frac{\partial \rho_{12}}{\partial t} = i \frac{\rho_{12}}{T_{12}} - \frac{\rho_{12}}{T_{12}} + i \frac{\hbar}{\mu_{12}} E_0^* (\rho_{22} - \rho_{11}) .$$  \hspace{1cm} (2)

Here $A_0(t)$ is the Gaussian envelope of the pulses, $\tau$ is their time separation, $\omega_L$ is the central frequency of the laser, $\mu_{12}$ is the transition-dipole moment, $\omega_{12}$ is the transition frequency, $T_{22}$ is the excited-state lifetime, and $T_{12}$ is the optical dephasing time. To obtain (2), the rotating-wave approximation was used.

We simulate the experiment by numerically integrating (2) for a large number of values of the optical pulse separation $\tau$. Figure 5 is a plot of the calculated final excited-state population $\rho_{22}$ as a function of $\tau$, for the three different values of the detuning shown in Fig. 4. The calculations are in good qualitative agreement with the experimental results for a value of $T_{12}$ of approximately 18 ps. The pulse duration was assumed to be 3.75 ps, and $T_{22} \gg T_{12}$. Similar calculations for the 1s-210 transition give us an approximate value of $T_{12}$ = 5 ps (not shown here).

V. DISCUSSION

The good agreement between the calculations and the experimental results, and a closer inspection of Eq. (2), lead to the following picture of the excitation and measurement processes. The first pulse induces a coherent time-dependent optical polarization (memory) $P(t) = 2N \text{Re}(\rho_{12}(t)\mu_{12})$, and transfers population into the excited state of the two-level system. After the first pulse, the optical polarization $P(t)$ decays with time constant $T_{12}$ due to dephasing processes. If the second pulse arrives before dephasing is complete, the electric field of the second pulse can coherently interfere with the optical polarization. This will lead to enhanced excitation to, or deexcitation of, the upper state depending on the phase difference between the electric field of the second pulse and the optical polarization. The final excited-state population therefore oscillates when the phase difference is varied, which is accomplished by varying the pulse separation.\textsuperscript{9,10} If the dephasing of the optical polarization is complete before the second pulse arrives, the excited-state population does not depend on the optical phase difference. Hence, by measuring the coherent population changes as a function of $\tau$, we obtain the decay time constant $T_{12}$ of the optical polarization. Note that in our experiment we can monitor the excited-state population by measuring the photoconductivity of the sample.

The apparent faster dephasing time of the 1s-210 that we find is consistent with the character of the 210 state as an autoionizing state with a lifetime much shorter than the $2p^+$ state.\textsuperscript{5,11} We would like to point out that the assumption $T_{12} \ll T_{22}$, used in the numerical integration of (2), is a realistic one for the 1s-2p$^+$ transition, justified by frequency-domain measurements on samples with a lower donor concentration. There it was found that the $2p^+$ excited-state population lifetime is at least several hundred ps.\textsuperscript{12} The lifetime $T_{22}$ of the 210 state, however, is probably short\textsuperscript{5,11} enough that we need its value in the future for an accurate determination of the dephasing time of the 1s-210 transition.

When the 1s-2p$^+$ transition is detuned with respect to the central frequency $\omega_L$ of the laser, the induced optical polarization oscillates at the transition frequency $\omega_{12}$, which is different from the central laser frequency. The second pulse interferes both with the electric field of the first pulse on the beamsplitter (for small $\tau$), and the induced optical polarization in the sample. The Fourier transforms of the photoconductivity measurements will therefore show two peaks (Fig. 4): a broad one that corresponds to $\omega_L$ and a narrow one that corresponds to $\omega_{12}$. We can also understand our time-domain measurements using time-domain arguments. For example, for a certain $\tau$ the electric field of the second pulse can be in phase with the electric field of the first pulse and simultaneously out of phase with the induced polarization. When we

\begin{center}
FIG. 5. Calculated excited-state population vs time separation for three different values of the detuning. We assume Gaussian pulses of 3.75-ps duration and a value of $T_{12}$ = 18 ps. The population lifetime is assumed to be infinite.
\end{center}
now change the pulse separation by approximately \( \pi/\omega_{12} \)
the situation is reversed: the electric field of the second
pulse is out of phase with the electric field of the first
pulse, but in phase with the induced polarization. Both
effects compensate each other, giving a zero net change of
the excited-state population. For pulse separations in be-
tween, both effects also cancel each other. Consequently,
a plot of the excited state population versus pulse separa-
tion will show a node as in Fig. 3(c).

Note that the parameters used in the calculations were
chosen for maximum qualitative agreement between the
calculations and the experimental results, and are slightly
different from those in the experiments. For instance, we
assume that the excitation pulses have a Gaussian tem-
poral envelope of 3.75-ps duration. In contrast, from the
autocorrelation measurement in Fig. 2(a), we can see that
the pulses used in the experiment clearly have a non-
Gaussian temporal envelope of roughly 3-ps duration. In
addition, the calculations assume that the transition is
homogeneously broadened, whereas strong experimental
evidence exists that the 1s-2p \(^+\) transition is inhomogene-
ously broadened.\(^{12}\) For these reasons, the value of the de-
phasing time \( T_{12} = 18 \) ps used in the two-level model
should be viewed as an estimate of the inhomogeneous
dephasing time only. The actual dephasing time can only be
determined by comparison of the data with a full-
model calculation that includes the exact pulse shape and
details of the (inhomogeneously broadened) line shape.
All essential features of the measurements, however, such
as the node in Fig. 3(c), are reproduced by the calcula-
tions since they do not strongly depend on either the line
shape or the exact pulse shape.

The time-dependent optical polarization radiates light
with the transition frequency and the efficiency of the
emission decreases when the polarization disappears.
This is the so-called free-induction decay from which the
optical-dephasing time can be obtained.\(^1\) Here, using the
induced photoconductivity, we have measured the decay of
the optical polarization which is the source of the
free-induction decay emission, not the emission itself, to
obtain the dephasing time. The method essentially frees
us from the difficult problem of doing a time-resolved
measurement of the (often) very weak far-infrared emis-
sion in the trailing wing of a pulse after it has propagated
through the sample. By implication, we may be able to
time resolve other optical coherent effects such as photon
echoes, by combining photoconductivity measurements
of the sample with the use of ultrashort pulses.

VI. CONCLUSIONS

We have used picosecond far-infrared pulse pairs from
the FELIX to time resolve the free-induction decay of
two Si-donor transitions in GaAs. Instead of detecting
the light emission of the sample to measure the decay, we
measure the decay of the optical polarization which is the
source of the emission. We have shown that we can mea-
sure the polarization decay by measuring the ability of a
second pulse to induce coherent changes in the popula-
tion excited by a first pulse. We detect these population
changes by measuring the photoconductivity of the sam-
ple.

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