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Hydrated-Electron Relaxation Dynamics Studied with 5-fs Pulses

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10.30-13.00 **QFG - Condensed Matter Dynamics** Presider: K. Duppen, University of Groningen, THE NETHERLANDS

10.30 QFG1

Hydrated-Electron Relaxation Dynamics Studied with 5-fs Pulses Maxim S. Pshenichnikov, Andrius Baltuska, and Douwe A. Wiersma

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Excess electrons in condensed-phase media play a crucial role in the dynamics of important Access technical processes. Among those are solution photochemistry, non-radiative electronic transitions, and charge transfer reactions, Hydrated electrons, i.e. electrons solvated in water, are of special interest. They can be viewed as an exceptional instrument for extracting information about the solvation process in water that plays an outstanding role in nature. Another motivation for a detailed study of the hydrated electron stems from the unique possibility to confront the predictions of mixed classical-quantum mechanical molecular dynamics simulations. This presents a direct way to verify the basic a priori assumptions that radically influence the outcome of computer modeling.

In this Contribution we present an experimental study of the energy relaxation of the photo excited hydrated electron. The results of frequency-resolved pump-probe with 5-fs pulses provide sufficient evidence in support of an extremely fast initial energy relaxation. Our data show that this process is controlled by librations of the surrounding water molecules and has a decay time of ~50 is. We further demonstrate that the subsequent cooling of the hot-ground state proceeds on a

is time scale and exhibits no isotope effect. The overview of the fit results of the experimental transient spectra is shown in Fig.1. The contribution of maintent spectra is shown in Fig.1. The control of the stimulated emission becomes insignificant after first 100 is due to depletion of the excited state (open circles). The frequency of the excited-ground state crossing point is -9500 cm^3 which corresponds to -2/3 (the initial position for the excited state). of the initial excitation frequency. Therefore, a large on the animal extension respective, interclote a mage amount of energy deposited on the hydrated electron is rapidly absorbed by the water molecules with the characteristic transfer time of -50 fs. ha accordance with our previous findings [1,2], this decay time is dominated by the librations of water molecules as concluded from a substantial isotope effect. The following relaxation occurs in the h

the United with a characteristic time of 1 ps (Fig.1b). At this state with a characteristic time of 1 ps (Fig.1b). At this stage the water molecules surrounding the electron transfer the accumulated energy into a collective-type translational motion, most probably via the existing hydrogen-bond network. Finally, a nearly full equilibration of the hydrated electron is achieved by ~6 ps.

References 1. M.F. Emde *et al.*, Phys. Rev. Lett. **80**, 4645 (1998). 2. A. Baltuska *et al.*, *d. Phys. Chem*, **A103**, 10065 (1999).

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Fig.1. Summary of the experimental results: relative amplitudes of the excited (open circles) and hot ground (gold circles) states (a) and the spectral shift of the peak of the tol ground-state absorption with respect to the initial transition frequency (b). Solid curves in the inset show a mono-exponential decay and rise with a 55-fs time constant.

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10.45 QFG2

EUTERPE

Decay Times of Surface Plasmon Excitation in Metal Nanoparticles Determined by Laser-induced Persistent Spectral Hole Burning

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Surface plasmon excitation in small metal particles has found great interest in the Surface plasmon exclusion in small interar platicles has fould great interest in the past, in particular since this collective oscillation of the conduction electrons can be stimulated with light and since its resonance frequency can be tuned over a wide spectral range by varying the size and shape of the clusters, by changing the dielectric surrounding and choosing different netals. Even though numerous investigations on SP's have been reported, no systematic investigation of the decay time T_2 and the decay mechanisms of SP excitation are available. Main reason is that nanoparticles unsult have a branch citera discharge dis ticles usually have a broad size and shape distribution which introduces inhomogeneous line broadening and prevents the determination of T_2 by linear optical spectroscopy and time resolved experiments using femtosecond laser pulses [1,2].

Here, we describe a novel technique to determine the homogeneous linewidths of ice plasmon resonances of metal nanoparticles in the presence of inhomogenesurfa soluce plasmor resonances of mean nanopartices in the presence of the presence perimental results and a theoretical model of hole burning the linewidth of 260 meV perminant results and a reverticat mode of the bulling the memory of 200 me corresponding to a decay lime of 4.8 fs was extracted. This value is shorter than ex-pected for damping by bulk electron scattering. We contcude that additional damping mechanisms, in particular surface scattering, come into play if the electrons are confined in particles with sizes below 10 nm. Furthermore, an influence of the shape of the clusters on the decay time was observed.

The technique presented here holds great promise for investigating SP's as a function of particle radius, an essential step towards a detailed understanding of the importance of different decay mechanisms. This is also essential to optimize the local field enhancement at the particle surface in a systematic way by generation of aggregates with optimum axial ratio and by precise control of the chemical surroun-ding including choice of the substrate material.

T. Vartanyan, M. Simon, F. Träger, Appl. Phys. B 68, 425 (1999) B. Lamprecht, J.R. Krenn, A. Leitner, F.R. Aussenegg, Appl. Phys. B 69, 223 [2] (1999)