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Femtosecond reactivity of electron in aqueous solutions

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Résumé. — Les techniques optiques femtosecondes qui permettent d'engendrer des impulsions intenses du proche UV au proche infrarouge sont utilisées pour étudier des réactions physicochimiques et photochimiques ultra-rapides dans des solutions aqueuses. Des résultats de l'étude dynamique de la solvatation d'un électron dans l'eau et une solution saline concentrée seront discutés.

Abstract. — Femtosecond optical techniques allowing the generation of intense optical pulses from the near ultraviolet to the near infrared are used in the monitoring of ultrafast photophysical and photochemical reactions in aqueous solutions. Selected examples for dynamics of electron solvation in water and aqueous solution of ferrocyanide are discussed.

1. Introduction.

Since its discovery more than twenty five years ago the hydrated electron continues to be of considerable interest and a matter of debate and controversy [1]. It is a species of prime importance in chemistry, biology and even technology because the energetics and the time dependence of the electron-medium interaction play an important role in the formation of this radical. It is also a theoretical challenge since the solvated electron presents the particular property that its structure is totally determined by the solvent. There has been continuing discussion over the relative role and chronological importance in the long-range and short-range electron-medium interactions in the sequence of electron trapping and solvation [2, 4].

Following a dielectric continuum picture of the solvent it has been suggested that the transition toward solvation in polar liquids is initiated by long-range dipolar interactions followed by self-trapping of the electron. This latter scheme has been favoured in liquid alcohols since a precursor of the solvated electron has been identified through an infrared absorption which extends to the visible and which decays within ten to a few hundred picoseconds, depending on the alcohol chain length. This has been interpreted as an electron localized in a shallow trap or a fluctuation of the surface potential, evolving from there toward the solvated state via a configurational relaxation of the medium in the electronic field of the charge. Until quite recently, only the fully relaxed solvated electron could be observed in liquid water, raising the assumption of direct localization of excess electron in pre-existing trapping sites [5].

Important theoretical breakthroughs have been realized recently by using computer simulations. Several groups have thus deduced some (conflicting) information on the structural aspect of the hydrated electron by employing path integral techniques while a molecular dynamic simulation of water has identified favorable sites for the initial trapping of electrons [6-9]. This fast evolution of computer simulations lacked however experimental grounds.

Employing femtosecond spectroscopy technique we have found unique information on the dynamics of both trapping and solvation of electron in aqueous solutions. While an unambiguous identification of the transient states cannot be reached directly by spectroscopy alone, these dynamical data demonstrate that electron solvation in water proceeds through at least one intermediate state whose lifetime is in the femtosecond regime.

In the following we will present the experimental scheme and results in the two cases of direct photoionization of a pure water molecule and photolysis of an aqueous solution of ferrocyanide ion. We will discuss our data in the framework exposed above.

The recent advances in ultrashort laser pulse generation permit the transfer to the picosecond and femtosecond time scale of most of the already developed optical methods making possible an accurate determination of the kinetics of electron solvation in polar medium. In such liquids, the electron solvation can intervene when an excess electron is ejected from a solvent or solute molecule. Spectral and kinetic investigations of the reactivity of an electron in polar media, have been carried out using a femtosecond pump-probe method [10, 11]. The experimental apparatus (Fig. 1) is based on a laser amplifier system producing high peak power tunable pulses of typically 100 fs duration. After passing through a variable delay line driven by a stepping motor, one beam is focused into a KDP crystal to produce up to 20 μJ, 100 fs at 310 nm (Fig. 1). The UV beam (pump or excitation beam) is then focused on a 300 μm diameter spot in the 2 mm long sample and initiates the ionization process. The other half of the 620 nm beam is focused into a 2 cm water cell and converted to a white light continuum. A part of the beam is selected with colored filters and split into two beams: one beam probes collinearly the excited part of the sample (probe or test beam) while the other (reference beam) misses the cuvette.

![Fig. 1.](image)

The two signals corresponding to the probe and reference pulse energy at a given wavelength are sent through an electronic chain (pulse shaping, sampling holder, digitizer) to a computer. This laser and amplifier set up coupled to this absorption spectroscopy detection scheme allowed us to measure changes in the optical density $A < 10^{-3}$ with femtosecond accuracy.

One key point in this experiment is the determination for each probe wavelength of both the instrumental instantaneous response and the zero time delay (coincidence) between the pump and the probe pulses. This has been done by observation of a very weak instantaneous induced absorption in a pure n-heptane solution excited at 310 nm in the same condition as water [11].

3. Results.

3.1 ELECTRON SOLVATION IN PURE WATER. — Up to now the great majority of electron solution studies in pure water has been realized by injecting electron using pulse radiolysis methods, providing at best a 10 ps accuracy. On the other hand, direct photoionization of water molecules by visible and near UV light does not seem possible since water is transparent in the UV spectral region up to 190 nm. However in our case of ultrashort pulse, light can be absorbed through a non-linear process, namely two photon absorption [12]. Indeed the two photon absorption coefficient of pure water at 310 nm has been estimated to be $4 \times 10^{-13}$ m/W yielding non negligible absorption when dealing with multi-gigawatt peak power pulses. In this case our 8 eV two-photon excitation is above the ionization threshold for liquid water, estimated to be around 6.5 eV (Fig. 2). So, each 100 femtosecond pulse produces a homogeneous solvated electron concentration of about 5 μM and furthermore permits the dynamics of electron solvation to be resolved with femtosecond accuracy.

![Fig. 2.](image)
accuracy, starting from initially quasi free state up to full solvation.

Induced absorption kinetics measured over the whole visible spectrum and in the near infrared up to 1250 nm have been used to reconstitute a transient spectrum. The high energy tail of an infrared band extending above 1250 nm appears within the excitation and is fully developed after a delay of 0.2 ps (Fig. 3). This IR band then decays concurrently with the rise of an absorption band which peaks around 720 nm and which coincides with the known solvated electron one. The kinetics at 1250 and 720 nm reflect the evolution of the infrared and solvated species respectively (Fig. 3). The infrared band appears with a time constant of 110 ± 20 fs and relaxes following a first order kinetics with a time constant of 240 ± 20 fs towards the solvated species.

Fig. 3. — Absorption spectra of the electron at 200 fs and 2 ps after femtosecond photoionization of liquid water at 294 K. Insert : time-resolved data obtained at 1250 nm and 720 nm following femtosecond photoionization of liquid water. The smooth lines represent the computed best fits assuming an appearance time of 110 fs for the infrared species (1250 nm) and its relaxation towards the solvated state in 240 fs.

These data are the first evidence of the existence of at least one transient intermediate of electron solvation in pure water. What is the nature of this state? If we follow the recent work of P. Rossky’s group [9] possible initial preexisting trapping sites are arising from statistical fluctuations and molecular clustering. They find a distribution of such sites monotonically in energy down to −1.4 eV, which is compatible with our IR data. We have proposed that this precursor is a state (presolvated state) where the electron is still spatially extended [13, 14]. Is it already a thermalized state? This would imply that the excess kinetics energy (1.5 eV at maximum) has been transferred to the water molecules in less than the risetime $T_1$ of the IR spectrum, i.e. 100 fs. Such fast thermalization is quite possible if we take into account the existing numerous modes of vibration of the solvent. These dynamics can in fact be compared to the dynamics of electron trapping in the shallow part of the exponential band as tail states in the amorphous materials process which has been demonstrated to occur in the femtosecond regime [15]. In this case the local temperature rise can be estimated to be negligible since the solvated electron spectrum taken only a fraction of picosecond after creation, and which is sensitive to temperature, is exactly as expected for $\theta = 294$ K.

Turning toward the transition to the fully solvated state, it has been suggested as mentioned earlier that once the electron gets localized, it evolves toward the solvated state via a configurational relaxation of the medium in the electronic field of the charge. Theories developed in this framework predict that this relaxation should occur through a continuous shift of the electron absorption with a characteristic time of the order of $T_L = 0.240$ ps where $T_L = \frac{T_D}{\varepsilon_0}$ with $T_L$ the longitudinal relaxation time, $T_D$ the Debye relaxation time, $\varepsilon_\infty (\varepsilon_0)$ being the high/(low) frequency limit of the dielectric constant [16]. Indeed we observe a decay of the IR species with a time constant $T_2$ close to $T_1$ and a concomitant rise of the known solvated electron spectrum. A picture of separate transient electronic states of electron would then be consistent with our data.

Fig. 4. — Schematic representation of the energy diagram and density states for presolvated and solvated electron in liquid water.
although we cannot completely rule out a dielectric relaxation of the medium. We may for instance assume a continuous shift of the energy of the electron associated with a decrease of Franck-Condon factors while the solvent has not reached its equilibrium. In this case the absorption of light by the electron would be negligible in the dynamic phase.

We may also notice that in our case the solvent is modified because we do not inject electrons but take them from a water molecule and create consequently a cation (H₂O⁺) which may self modify the medium properties. To study such a possible influence, we have extended our study to the case of electron solvation in ionic solutions.

### 3.2 Electron Solvation in Aqueous Ferrocyanure Solution.

The femtosecond ultraviolet photochemistry of aqueous solution of Fe(CN₆)⁴⁻ (0.45 M) permits the process of electron solvation in aqueous concentrated ionic solutions to be investigated. Femtosecond photolysis of an aqueous solution of ferrocyanide ion was performed using 310 nm pulses of 100 fs duration. At this wavelength, it is expected that the quantum yield of aqueous electron formation is near 0.1 [17]. Previous photochemical studies of the ferrocyanide ion have established that the excited state of Fe(CN₆)⁴⁻ in water may dissociate in a thermally activated process to yield a solvated electron in less than 0.3 ps [5] according to the following equation:

\[
\text{Fe}^{2+} + \text{hν} \rightarrow \text{[Fe(CN₆)³⁻]}^* \rightarrow \text{Fe}^{2+} + \text{CN}^{-} + \text{e}^- \text{aq}.
\]

This photochemical process competes with a reversible photoaquation reaction leading to the formation of a pentacyanoaquo complex:

\[
\text{[Fe(CN₆)³⁻]}^* + \text{H₂O} \rightleftharpoons [(\text{Fe(CN₆)H₂O})³⁻] + \text{CN}^{-}.
\]

The time-resolved spectroscopy data in the infrared and visible (Fig. 5) demonstrate that photodetachment of electron and subsequent solvation proceed through at least one intermediate state. This intermediate appears with a time constant of 120 ± 20 fs and relaxes towards a solvated state following a first order kinetics with a time constant of 240 ± 30 fs. At 720 nm, the signal follows the kinetics equation

\[
A'(0) = A_0^0(1 - (1/(T_2 - T_1))) \times (T_2 \exp(-(t/T_2)) - T_1 \exp(-(t/T_1)))
\]

as expected after a first-order kinetics of appearance of the precursor of the solvated species with $T_1 = 120 \pm 20$ fs as observed at 1 250 nm and a subsequent relaxation with $T_2 = 240 \pm 30$ fs.

This behaviour which is identical to what has been observed in pure liquid water demonstrates that electron solvation in concentrated solution of ferrocyanide ion does not proceed through a direct electron capture by pre-existing deep traps as suggested previously [5]. So, the presence of the negative ion Fe(CN₆)³⁻ at high concentration does not restructure the immediate water environment and does not provide more favorable spatial distribution of deep trapping sites than in pure liquid water. The reactivity of electron with concentrated ionic aqueous solutions proceeds through at least two transitions involving a transient localized state absorbing in the infrared (Fig. 4) and a subsequent fully solvated state. The time constant that we find for the appearance of the IR absorbing species and its decay toward solvated state are similar in pure water and aqueous ionic solution. This implies that the existence of the presolvated state of electron is neither influenced by the method of electron photoejection nor by the ionic strength of the polar medium in the range 0-0.45 M.

Assuming, from previous experiments [18], that thermalization and solvation distance of electron with exogenous energy around 1 eV is about 40 angström, it can be inferred that many of the quasi free electron with excess kinetics energy escape from the vicinity of the cation (H₂O⁺) or the anion Fe(CN₆)³⁻ and that long range Coulomb type interactions do not influence neither the initial presolvation nor the configurational relaxation of the medium in the electronic field of the charge. In particular we have no evidence of geminate recombination of presolvated or solvated electrons.

The very first risetime for the appearance of the presolvated state of electron in pure water and aqueous ionic solution implies that the efficient mechanism involved in the localization process does not require large structural reorganization of water molecules in the vicinity of the electron. We may
speculate that presolvation of excess electron in concentrated ionic solution is promoted, as expected for pure liquid water, by statistical fluctuations in the electronic charge density of the solvent molecules. However the complete understanding of the electron localization process implies knowledge of the distribution and the structure of initial shallow traps. The exact identification and understanding of bound surface states and microscopic trapping sites in concentrated ionic aqueous solutions need more theoretical investigation with tools like path integral techniques or molecular dynamics simulations by Monte-Carlo methods.

In conclusion, the femtosecond photoionization of water and inorganic anion by high power UV pulses induces an injection process of excess electrons in polar medium and furnishes useful information on the basic unresolved question of kinetics and mechanisms of electron localization and solvation in aqueous environment. This paper demonstrates that the electron solvation in pure liquid water and concentrated ionic solution at room temperature proceeds through at least two transient steps with, in particular, an infrared absorbing presolvated state. These data support neither the assumption of the predominant role of the high density of pre-existing deep traps into which the electron fall directly after photoejection nor the existence of a pure dielectric relaxation mechanism for the final energy rearrangement linked to the solvation step. Such data are fundamental for theoretical investigations exploring the obvious quantum mechanical character of the structural and dynamical properties of the electron in polar fluids with great ionic strength.

References