СВЕРХБЫСТРАЯ ЭЛЕКТРОННАЯ КРИСТАЛЛОГРАФИЯ И НАНОКРИСТАЛЛОГРАФИЯ:
ДЛЯ ХИМИИ, БИОЛОГИИ И НАУК О МАТЕРИАЛАХ. ЧАСТЬ I. СВЕРХБЫСТРАЯ
ЭЛЕКТРОННАЯ КРИСТАЛЛОГРАФИЯ

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Прямое исследование и понимание динамики химических и биологических процессов, происходящих в конденсированных средах, в настоящее время находится на ранней стадии. Прогресс в этой области был вызван разработкой методов исследования структурной динамики матери в состоянии, далеком от равновесия, включая экстремальные состояния. Представленная информация служит основой для тестирования новых теоретических подходов к описанию вещества в причинно-связанной триаде «структура-динамика-свойство». Наблюдение динамического поведения материи в пространственно-временном континууме на сверхкоротких временных масштабах является необходимым первым шагом в объяснении и, в конечном счете, контроле неравновесных процессов и функциональности исследуемых систем. Метод сверхбыстрой электронной кристаллографии позволяет исследовать переходы неравновесные структуры, которые дают решающую информацию о структурной динамике фазовых переходов и когерентной динамики ядер в твердом состоянии, на поверхности и в макромолекулярных системах. В последние годы длительность электронного импульса в аппаратуре сверхбыстрой электронной дифракции значительно уменьшилась, а ускоряющее напряжение значительно увеличилось. В результате, были получены фемтосекундные импульсы электронов. Предложен метод радиочастотной групировки электронов для увеличения яркости электронного импульса. Для увеличения пространственной когерентности использовался метод электронной полевой эмиссии, а для уменьшения рассогласования скоростей светового и электронного импульсов и сжатия электронных пучков применялось пондеромоторное волновое фронтальное ускорение...
Эти достижения открыли новые возможности для изучения когерентной структурной динамики - атомно-молекулярного кино с фемтосекундным временным разрешением. В обзоре цитируются результаты нескольких всемирно известных исследовательских групп.

Ключевые слова: сверхбыстрая электронная кристаллография, промежуточные неравновесные структуры, поверхности и кристаллы, когерентная структурная динамика, динамика реакции с атомным разрешением, фемтосекундная спектроскопия

ULTRAFAST ELECTRON CRYSTALLOGRAPHY AND NANOCRYSTALLOGRAPHY: FOR CHEMISTRY, BIOLOGY AND MATERIALS SCIENCE. PART I. ULTRAFAST ELECTRON CRYSTALLOGRAPHY

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The direct probing and understanding of the dynamics of chemical and biological processes occurring in condensed matter, is currently in its early stages. Progress in this field has been pushed by the development of methods for the study of the structural dynamics of matter in a state far from equilibrium, including extreme states. The forthcoming information serves as the basis for testing new theoretical approaches to the description of the substance in casually connected triad "structure-dynamics-function". Observation of the dynamic behavior of matter in the space-time continuum on ultrashort time scales is a necessary first step in the explanation and, ultimately, control of far from equilibrium processes, and functionality of the systems studied. The method of ultrafast electron crystallography (UEC) makes it possible to investigate transient nonequilibrium structures, which yield decisive information about the structural dynamics of the phase transitions and coherent dynamics of the nuclei in the solid state, on the surface, and in macromolecular systems. In recent years, the electron bunch path length in the UEC apparatus diminished significantly, while the accelerating voltage increased considerably. Therefore, femtosecond electron pulses were obtained. A technique of radio frequency grouping of electrons was proposed to increase the electron pulse brightness. The method of electron field emission was used to increase the spatial coherence, and ponderomotive wave front acceleration was applied to reduce the mismatch between the velocities of the light and electron pulses and to contract the electron bunches. These achievements have opened up new possibilities for studying the coherent structural dynamics – atomic and molecular movie with femtosecond temporal resolution. The results of several internationally renowned research groups are included and cited.

Key words: ultrafast electron crystallography, transient nonequilibrium structures, surfaces and crystals, coherent structural dynamics, atomically resolved reaction dynamics, femtosecond spectroscopy
INTRODUCTION

The direct probing and understanding of the dynamics of biological, chemical processes, and processes occurring in solids (e.g., transport in metals, their alloys, rectification effects in semiconductors, magnetoresistance etc), which are amenable to study in real time, is currently in its early stages. Progress in this field has been pushed by the development of methods for the study of the structural dynamics of matter in a state far from equilibrium, including extreme states. The forthcoming information serves as the basis for testing new theoretical approaches to the description of the substance in casually connected triad "structure-dynamics-function". Observation of the dynamic behavior of matter in the space-time continuum on ultrashort time scales is a necessary first step in the explanation and, ultimately, control of far from equilibrium processes, and functionality of the systems studied.

The natural time scale of dynamic processes depends on the energy and length scales of the phenomenon of interest. For example, from attosecond to femtoseconds advancing time-resolved and control techniques from the molecular to the electronic time-scale; from femtoseconds to picoseconds for atomic, molecular dynamics relevant to chemistry; from picoseconds to nanoseconds for ferroelectric, magnetic diffusion, and polarization processes; from nanosecond and longer for nucleation, cellular functions, and growth of crystals. The corresponding natural length scale for these dynamic processes are in the range of atomic and molecular scale 0.1-10 nm for chemistry and biology, up to the micron scale of interactions between biological cells in grains and metal alloys.

A full description of the temporal sequence and interaction of dynamic characteristics requires the integration of information from different instrumental methods based on different physical principles, first and foremost, diffraction and spectroscopic methods with a time resolution and spatial resolution allowing the study to limit of the coherent dynamics of matter.

The quest to observe transient molecular structures and the chemical dynamics of molecules during reactions is greatly aided by employing two vantage points: the traditional spectroscopic view, which relies on the observation of energy levels and their populations, and a diffraction view, which directly images molecular structure (see more detailed discussion of this issue in the works [1-9]. Already in the 1980’ s, the diffraction paradigm was formulated: implementing electron diffraction with time resolution adds a temporal coordinate to the determination of molecular structures [10-13]. Time-resolved electron diffraction (TRED) rested on the concept of flash photolysis originally proposed by Norrish and Porter in 1949 [14]. Very recent advances in the generation of x-ray pulses have made possible the closely related time-resolved x-ray diffraction (TRXD). In both methods, short laser pulses create the transient structures and induce chemical dynamics that are subsequently imaged by diffraction at specific “points” in time.

The use of pico- or femtosecond bunches of electrons as probes, synchronized with the pulses of the exciting ultrashort laser radiation, led to the development of ultrafast electron crystallography and nanocrystallographic techniques [15], of dynamic transmission electron microscopy [1-3, 16, 17], to femtosecond electron diffraction of structural transitions [18] and of molecular quantum state tomography [19, 20]. One of the promising applications, developed by the electron diffraction methods, is their use for the characterization and the “visualization” of processes, occurring in the photo-excitation of free molecules and biological objects for the analysis of different surfaces, thin films, and nanostructures (see the recent review articles [6, 9, 21-37]). The combination of state-of-the-art optical techniques and diffraction methods, using different physical principles but complementing each other, opens up new possibilities for structural research at ultrashort time sequences. It provides the required integration of the triad “Structure-Dynamics-Function” in chemistry, biology, and materials science [1, 2, 5, 6, 8, 9, 38-40].

Since the 1980s, scientific laboratories around the world have begun to develop a new field of the research aimed at this goal. “Atom-resolved-molecular movies” will allow visualization of coherent dynamics of the nuclei in the molecules and fast processes in
chemical reactions in real time. Modern femtosecond laser sources have made it possible to significantly change the traditional approaches using continuous electron beams, to create ultrabright pulsed photodetector sources, to catch ultrafast processes in the matter initiated by ultrashort laser pulses and to achieve high spatio-temporal resolution to imaging molecular processes. Ultrafast laser spectral and electron diffraction methods complement each other and open up new possibilities in chemistry and physics to light up atomic and molecular motions involved in the primary processes governing structural transitions.

Ultrafast Electron Crystallography (UEC) is an extremely powerful tool for the investigation of the transient non-equilibrium structures. These studies provide rather important information on the phase transitions and coherent dynamics of the nuclei in the solid state. In the past few years UEC was widely used also for study of structural dynamics in different nanostructures. This variation of the technique is usually referred to as UEnC (Ultrafast Electron nano-Crystallography [5, 6, 15, 21, 22, 27].

The article discusses the application of time-resolved electron diffraction to investigate condensed matter and includes the discussion of the experimental techniques, the theory of both UEC (Part I) and UEnC (Part II), and some of the prominent results on the study of the structural dynamics of the surface, different phase transitions in the nanoparticles, and the dynamics of the molecular assemblies. By providing high spatial resolution (down to a few hundredths or thousandths of angstrom) and high temporal resolution (in pico-femtosecond range) it is possible to observe the structural dynamics and the energy flow in condensed matter in real time.

1. EXPERIMENTAL SETUPS

Schematic diagram of UEC experimental setup is shown in Fig. 1 [1, 41].

The experimental setup consists of three ultrahigh vacuum chambers with an electron source, LEED (Low-Energy Electron Diffraction) and Auger spectrometers and a batch box of gas that allows to control the preparation of a clean surface (see, e.g., [42]), its state during the adsorption of the atoms and the molecules and to determine the nature of its binding to the surface (chemisorption or adsorption). The device is equipped with a goniometer with an angular resolution of 0.005° and a cooling system capable of reaching temperatures of down to ~110 K. The accuracy of the inter-nuclear distances is ~ 0.01 Å, while the maximal resolved distance is about 50 Å. Here the laser pulse duration is ~120 fs with the penetration depth of the laser radiation being of 4 nm at 266 nm and 7 µm at 800 nm.

In the laboratory of prof. Ruan at the Michigan State University (USA) it has been used the latest achievements in the field of the generation, the control and the detection of ultrashort electron pulses for UEC-researches with sub-picosecond time resolution [15]. The apparatus is shown in Fig. 2. It consists of a femtosecond laser and electron gun with silver photocathode (thickness of silver layer is 40 nm). Here 800-nm laser radiation is utilized for the photo-excitation of the sample, while 266-nm-pulses are used to generate electron bunches from the cathode. Since the laser pulse (266 nm) that initiates the photoemission of the electrons is obtained by splitting of the radiation from the same laser setup, the temporal jitter between the exciting and the probe pulses is determined by the temporal interval of the photoemission process and, in principle, has a femtosecond level or below [43]. The arrival of the probing electron bunch with respect to the pump pulse is controlled by the optical delay line that allows observe the dynamics of the photo-induced reaction. The...
laser system (Ti: Sapphire laser with regenerative amplification from Spectra Physics) produced 45 fs laser pulses at 800 nm with an energies of up to 2.5 mJ and a repetition rate of 1 kHz. Using the frequency doubling or tripling, it is possible to get the 400-nm, or 266-nm pulses with the energies of up to 250 mJ and 40 mJ, respectively, which can also be utilized as a pump pulse.

The photocathode generates the electron pulses with kinetic energy of 30 keV (anode extraction field is about 6.5 kV/mm), which pass through the apertures and magnetic lenses and focused to the diameter of 5 microns (Fig. 3) on the sample located at a distance of 5 cm from the photoelectron source. The first short-focused femtosecond photoelectron gun was designed in Miller’s group [18]. It allowed to achieve high electron density (of up to $10^4$ electrons in the bunch) and to observe, in principle, the diffraction pattern in UEC using a single electron pulse. Note, that in UEC it is sufficient to have about $10^3$ e/pulse at the pulse repetition rate of 1 kHz.

The efforts of Ruan’s group [15] were trained on the focusing of the electron pulsed beam to a diameter of about 1 µm while maintaining femtosecond temporal resolution in order to reduce the mismatch of the velocities of the light and the electrons [44]. Such way ultimately allowed study individual (isolated) nanoparticles with high spatio-temporal resolution.
To demonstrate the possibilities of the existing apparatus, the photo-excitation of highly-ordered pyrolytic graphite (HOPG) has been studied. In HOPG, the phonon mode $E_{2g}$ corresponds to the inter-planar oscillations of the shear of graphite layers (Fig. 4). It is known that these oscillations interact strongly with the femtosecond laser field [45, 46].

After the photo-excitation, it is possible to observe the generation of surface charges a subpicosecond time scale (Fig. 5). The oscillations are associated with the charge accumulation on the outside surface of pyrographite and are caused by photo-induced electron tunneling through a potential barrier from the substrate surface which is followed by the formation of two-dimensional (2D) electron gas. In this case, the positively-charged ions remain behind the photo-excitation and is modulated by the surface phonons. This optically-induced charge redistribution and the corresponding atomic rearrangements are discussed in Sect. 3.

Fig. 5. Discovery of femtosecond jumps of the surface charges of Highly-Ordered Pyrolytic Graphite (HOPG) after the photo-excitation. Solid line is the result of a sinusoidal approximation of $E_{2g}$ phonon mode oscillation with a period of 0.78 ps, which demonstrates sub-picosecond resolution of UEC experimental setup [15]
(700-800 fs) is about of one order of the magnitude higher than that of the laser pulse (60 fs). The electron pulse formed in this way arrives into the resonator, where the electron velocities are modulated.

The amplitude of rf oscillations in the resonator was selected such that the electron pulse was grouped at the distance of 100 to 200 mm from the cavity. The working frequency of this rf cavity was selected as 6.2 GHz. In addition to the target and photocathode, a part of the laser beam is fed to the photo-detector in order to synchronize the microwave oscillations in the resonator with laser pulses. The oscillation phase stability should be about $4 \times 10^{-4}$ rad to achieve a 10 fs time resolution. According to the calculations, the maximal amplitude of the voltage across the resonator gap should be 3 kV. It is necessary to have a microwave signal source with a –4W to 6W output power to obtain this voltage. The stability of amplitude oscillations in the cavity should be approximately ±0.25%.

A low-noise microwave signal source with two output frequencies (6.2 GHz and 6.1 GHz) was designed at the Research Center of the Siberian Branch of the Russian Academy of Sciences (Krasnoyarsk) to ensure the above requirements [47]. The power of the microwave signal (6.2 GHz) can be regulated in the range of 0.1-6.0 W, while the power of a 6.1 GHz frequency signal is $> 5$ mW. To obtain a low phase noise level and simultaneously ensure electron frequency tuning, the output signal is formed by mixing two signals, i.e., a signal of the generator across the dielectric resonator (GCV) and a signal of the generator controlled by voltage (GCV). The frequency is tuned by the GCV, which also has a low phase noise level at its fundamental frequency. In the scheme the power is controlled by changing a gain of the preliminary stage of the amplifier.

For the convenience of its operation, the device has an output signal power-meter and the frequency-meter, which shows the frequency detuning of the output signal from the rated value. As is known, GDR features a low phase noise level of about –(95-100) dB/Hz, when the offset from the carrier is 10 kHz. In the designed scheme, in order to obtain a lower phase noise-level, GDR is synchronized by a 100 MHz reference signal using the phase-lock (PL). The synchronization circuit is based on sampling of the GDR signal by the short pulses formed from the reference signal with a lower phase noise level in the stroboscopic phase detector (SPD). The signal from SPD arrives through the low-pass filter (LPF) and an amplifier (PL loop filter) at the control input of the generator and finely adjusts its frequency and simultaneously compensates the phase noises of the generator. As a result, it was possible to decrease the noises down to –(115-120) dB/Hz, when the offset from the carrier was 10 kHz. This setup for the preparation of femtosecond electron bunches was supplied to the Max Planck Institute (Garching, Germany). Using such scheme the first experiments were performed and the first results were obtained [47].

2. STRUCTURAL DYNAMICS IN CONDENSED PHASE

The method of ultrafast electron crystallography makes it possible to obtain the information about the dynamics of coherent structures in the photo-induced phase transitions in nanoparticles and macromolecules, on the solid surfaces, in thin films and the interfacial areas. It allows explore the dynamic processes at the level of the constituent elements [1, 15, 41, 49-51].

30-keV electron beam with the corresponding wavelength of ~0.07 Å (the wave vector is labeled as $k_{i}$) is projected at the angle of $\theta < 5^\circ$ on the crystal surface with adsorbed atoms. The resulting diffraction pattern gives information about the structure of the surface defined by both the substrate and the adsorbed particles. Here Ruan with the co-workers [50] changed temperature of the substrate using 800-nm 120-fs laser pulses. The reference point when the pump and the probe pulses hit the surface of the crystal simultaneously was defined as time-zero $t_{0} = 0$ (see the inset in the upper left of Fig. 3.7). Note, that the apparatus (Fig. 3.1) was pumped down to about 10$^{-10}$ Torr. The recorded diffraction patterns, $\Delta I(\theta, s; t_{ref})$, are the difference curves related to the underlying structure of the transient state:

$$\Delta I(\theta, s; t_{ref}) = I(\theta, s; t_{ref}) - I(\theta, s; t),$$

where the reference time $t_{ref}$ can be selected either before or after the arrival of the exciting pulse. It is clear, that the diffraction pattern reveals the structure in the reciprocal space, after Fourier transformation.

The scattering intensity $I(\theta, s; t)$ in Eq. 1 in the kinematic approximation can be represented as follows [51]:

$$I(\theta, s; t) = \Phi_{H} \Phi_{V} |F_{2} - F_{1}|^2 + \Phi_{H} \Phi_{V} |F_{3} - F_{5}|^2 + \Phi_{H} \Phi_{V} |F_{3} - F_{5}|^2,$$

where there is an explicit dependence on the scattering angle and the spatio-temporal changes. The phase coherence is also evident. For example,

$$\Phi = \sum \exp(-i s r),$$

where, for $\Phi_{H} r = na + mb$, but for $\Phi_{V} r = le$.

The structural factors of the unit cell $F$ are defined as follows:

$$F = \sum f_i(s) \exp(-B s^2) \exp(-i s r),$$

where $B = $ Debye-Waller factors and $f_i(s)$ – atomic scattering factors.
Eq. 4 has been utilized for different values of $\theta$ and $t$ in order to reproduce correctly the diffraction data and the rocking curves. Here it was taken into account the surface potential and the finite length of the coherent scattering of the electrons (see Sect. 3). In the structural analysis, the difference method is utilized which eliminates the influence of the scattering intensity from the ground state as well as the incoherent nuclear scattering, the inelastic scattering and the volume resonance [52].

In the reciprocal space for 2D monolayer of atoms the diffraction "rods", separated by a distances of $a$ and $b$ (Fig. 7), are observed. These rods correspond to the constructive interference of coherent waves. As the monolayer is embedded in a crystalline substrate, the rods in the diffraction pattern are changing due to the modulation of the distance between the planes (Fig. 7). For the electrons, the Ewald spheres defined by the vector $k_i$, have rather large sizes, and the diffraction pattern, which is dependent on $\theta$, has the bands at small scattering angles and correspondingly the Bragg spots at large scattering angles in the Laue zones.

The reflections reveal the static structure of both the surface and the lattice. But with high temporal resolution it becomes possible to carry out the additional impressive measurement. First, one can diagnose the structural changes of the surface layer and its re-structuring in real time. Secondly, there is a significant difference of the characteristic temporal scales for the processes occurring in the surface layer and in the orthogonal direction. Thus, it is possible to isolate and diagnose the initial non-equilibrium structures (but not the structures that arise during the propagation of the exciting pulse). Third, if the surface is used as the matrix, one can investigate the strengthening of the mutual influence of the substrate and the lattice and represent the structural dynamics of the process. The detailed theory of UEC is presented in the papers [53-55].

![Fig. 7. Schematic diagram of ultrafast electron crystallography. Electron beam is directed to the surface of a crystal Si (111) at an angle of $\theta<5^\circ$. The Ewald spheres and Laue zones (L0, L1, ...) are depicted (see text). The lower part of the figure shows the structure and the distance between the bilayers (3.136 Å). On the surface layer, the adsorbed atoms or molecules are presented [1,41,50]](image-url)
2.1. SURFACES AND CRYSTALS

Vigliotti with the colleagues have performed an investigation of the structural dynamics of crystalline GaAs surface after the rapid increase of its temperature [56]. Using the changes in Bragg diffraction (a shift, a line width and an intensity) it has been shown, that "contraction" and "expansion" occur from -0.01 Å to +0.02 Å and that "transient temperature" (its precise definition is not given in the above-mentioned paper) reaches its maximal value of 1565 K after 7 ps (Fig. 8). The structural changes are delayed from the increase in temperature, which is demonstrated by the evolution of non-equilibrium structures.

These results [1] were compared with non-thermal femtosecond optical sensing [57]. Here GaAs surface was covered with a monolayer of chemically bound chlorine atoms. On ultra-short time scales, the contraction followed by the expansion was observed, which was caused by the increase in temperature of the phonons. On larger time scales there was observed the restructuring and the corresponding evolution towards the equilibrium structure. The registered structural dynamics can be divided into three modes: the changes, which include electron redistribution without the motion of the nuclei (from femtoseconds to a few picoseconds), coherent non-equilibrium lattice expansion (starting from 7 ps), the restructuring and the diffusion of heat (from 50 ps to several nanoseconds).

Similar studies were performed for silicon crystals in the presence and in the absence of adsorbates. The selection of the ground state as the reference point showed the changes in the structure caused by the initial femtosecond pulse, compared with the sample in the ground state at "negative" temporal delay (Fig. 8). The structural dynamics was manifested by the temporal shift on the rocking curve of the Bragg in-phase peak, while the increase in the amplitude of the corresponding oscillations was displayed in the broadening of the peaks.

2.2. CONVERSION OF GRAPHITE INTO DIAMOND

The conversion of graphite into diamond using different nano-carbon materials is of great particular interest. Such process is believed to be associated with an intermediate state of the rhombohedral phase of graphite [58, 59]. While these two forms of graphite are energetically almost degenerate, they are separated by a large energy barrier [58, 60]. Thermodynamically, these transitions can occur only under very high temperatures and pressures. Recently performed theoretical studies have shown the possibility of reducing the energy barrier by doping of graphite with charged particles [60], or by electronic excitation [46]. This was demonstrated experimentally: the formation of nanodiamonds was observed at ambient temperatures by projecting a beam of argon ions with high charge (Ar^{8+}) on graphite following the charge injection from the tip of a scanning tunneling microscope [61].

Raman et al. investigated the possible paths of photo-induced lattice motion in pyrographite using UEC [62]. Remember, that highly-oriented pyrographite consists of 2D graphite micro-domains with randomly-oriented basal planes. The excitation laser pulse had its maximum at 800 nm.
This excitation is associated with shear oscillation of the \( E_{2g} \) phonon mode \([45]\) and can cause a change in the symmetry of the hexagonal graphite layers (ABAB) into the rhombohedral layers (ABCABC), which is required for the formation of sp\(^3\) bonds between the layers \([63]\). The diffraction pattern of the pyrolytic graphite ground state consists of (006), (008), (0010) and (0012) Bragg peaks. The analysis of time-dependent scattering intensities showed that the thermalization time is \( \sim 8 \) ps which corresponds to the relaxation of the excess energy from the originally-populated mode \( E_{2g} \) into the thermal reservoir of the phonon vibrations \([64]\). The reduction of the intensities of Bragg peaks (006) and (0010) with respect to (008) and (0012) was also observed (Fig. 9), which corresponded to the decrease of the lattice constant by a factor of two. This was accompanied by the formation of the peak at \( r = 1.9 \) Å on LDF (the Layer Density Function) of graphite which was due to the new distance between the layers.

Fig. 9. The intensity of the electron scattering \( M(s,t) \) and the corresponding layer density functions (LDF) for the selected times after the rapid photo-excitation of pyrolytic graphite. The appearance of the peak at 1.90 Å indicates to the bond formation between the planes \([15]\).

The disappearance of the peak at long temporal delays indicates to the formation of a reversible sp\(^2\) – sp\(^3\) hybrid structure similar to that found in the work \([65]\), where the graphite-diamond transformations were studied at high pressures. The process of diamond formation is likely caused by Coulomb attraction generated by photo-induced charge separation at the surface via injection of hot electrons into the surface of the substrate.

2.3. FATTY ACID CRYSTAL BILAYERS: MOLECULAR ENSEMBLES

Studies of the bilayer of fatty acids deposited onto a hydrophobic surface using Langmuir-Blodgett technique were performed using UEC. Langmuir-Blodgett technique makes it possible to control the molecular deposition of ordered molecular film layer by layer, and is rather often used for different membranes building.

Arachidic acid was used as an example. The unit cell for arachidic acid is characterized by the following set \( a_0 = 4.7-4.9 \) Å, \( b_0 = 8.0-8.9 \) Å, \( c_0 = 2.54-2.59 \) Å, which depends on the pH and the deposition procedure. The increase in the length of \(-\text{CH}_2-\text{CH}_2-\text{CH}_2-\) fragment by \( \sim 0.01 \) Å was experimentally observed. Fig. 10 shows the time-dependent intensity of diffraction signal.

Chen with the co-workers \([66]\) investigated by UEC a single bilayer (two chains \( \text{C}_{19}\text{H}_{39}\text{COOH} \)) of arachidonic (eicosanoic) acid. They determined the structure and the orientation of aliphatic chains as well as the size of the molecular sub-cell (–\(\text{CH}_2–\text{CH}_2–\)). The
studies made it possible to observe the structural dynamics in the bilayer placed on the substrate, the motion of atoms and self-organization of layers [67]. Fig. 10 shows the structure of the investigated bilayer and the corresponding sub-cell in two directions.

All diffraction patterns consist of spots (and/or bands), that demonstrates a high quality of two-dimensional crystal structure of the arachidonic acid bilayer and surface of Si (111) substrate functionalized with hydrogen. The diffraction patterns at "negative time delays" and a small electron incidence angle (at parallel and perpendicular incidence directions) suggest that the tilting angle of chains is close to zero and the parameters of the bilayer sub-cell should be given by the set of \( a_0 = 4.7 \text{ Å}, b_0 = 8.0 \text{ Å}, c_0 = 2.54 \text{ Å} \).

The symmetry of the bilayer corresponds to rhombohedral packing R (001) with (001)-plane being parallel to the Si(111)-surface. These experimental values of the lattice parameters differ from theoretical values of \( a_0 = 4.96 \text{ Å}, b_0 = 7.4 \text{ Å} \) [68]. Such discrepancy can be explained by the fact that the theoretical values were calculated for infinitely-long aliphatic chains and the calculations did not take into account the terminal carboxyl group of fatty acids. Furthermore, the bilayer is consists of two monolayers, so that the substrate and the conditions of its deposition (e.g. its pressure and pH) definitely play an important role. Note, that the distance \( c_0 \) between (CH\(_2\))-planes is in agreement with the theoretical value of 2.54 Å. In the experiment the selective structural dynamics of the bilayer was investigated. As shown in Fig. 3.10, already at the delay of about 1 ps, after the heating pulse, the weakening of the Bragg diffraction peaks was observed. The changes in the Bragg diffraction peaks become more prominent for larger temporal delays (10-100 ps). Intriguingly, both electronic and thermal pulses did not destroy the bilayer, that has been demonstrated in multiple experiments.

The observed behavior of the diffraction pattern reflects the initial stretching (\( \Delta c_0 = 0.1 \text{ Å} \)) of the sub-cell in the bilayer induced by the heating pulse, followed by the restriction caused by heat dissipation. The stretching is taking place with the time constant of \( \sim 25 \text{ ps} \), while the subsequent contraction occurs with the time constant of \( \sim 55 \text{ ps} \). Note, that the possible compression on nanosecond temporal intervals is restructure. With 150-fs temporal resolution in the plane, perpendicular to the molecular chains, no significant changes were observed [67]. Here it should be noted that the morphology and the lattice parameters of the surface layer depend on of the matrix and the modification of the surface [69].

3. PHOTO-SWITCHED SYSTEM FROM INSULATING TO METALLIC PROPERTIES

The development of high-brightness electron sources was key to opening up this class of study. To gain some appreciation of the quality of the diffraction data that made this possible to see the dramatic improvement with source brightness [27].

Improvement in source brightness enabled a dynamic observation of the photo-induced structural changes in the interesting charge-ordered organic system comprising ethylenedioxytetrathiafulvalene (EDO-TTF) and PF\(_6^–\) counterions, (EDO-TTF)\(_2\)PF\(_6^–\), shown in Fig. 11.

This system can be photo-switched from insulating to metallic properties [70] by means of a charge-transfer process strongly coupled to nuclear modes, stabilizing the change in charge distribution, as shown schematically in Fig. 11. Inspection of the differences between the insulating and metallic structures shows that the formation of the metallic state involves the flattening of the EDO-TTF moieties. The displacement of a bending mode toward this planar configuration would lead to an increase in the wave function overlap between molecules and increased electronic delocalization as part of the onset to metallic properties. Within a conventional transition-state picture, one would naturally expect the bending coordinate to be the dominant mode in this process. However, this simplified line of thinking only works for few atom systems. Considering just the molecules within a single-unit cell, this problem involves over 280 different degrees of freedom or dimensions. However, it was found that all of the diffraction orders could be fit by the displacement of just three reduced modes (Fig. 11C) in which the motion of the heavy PF\(_6^–\) counter-ion appears to be the key mode. In hindsight, this observation is understandable because the photo-induced change in electron distribution will lead to a change in the local field that will exert a force on the counter-ion [9, 21]. The PF\(_6^–\) ion is rather large, and its motion, through steric effects, couples the other modes. The projections along the three reaction coordinates (Fig. 11C) look like shadow projections of one another; the modes are strongly correlated.

One typically uses an approximate frozen slice of a many-body potential to discuss reaction coordinates and get a feel for the forces and types of motion involved in directing the process. However, these results show that the modes are dynamically coupled and that one cannot intuitively guess which modes are involved or the relative degree of coupling. In principle, time-dependent ab initio theory can provide the information on the relative degree of coupling between the different possible motions [9, 21]. There is a limit. Even
the highest level of time-dependent ab initio theoretical methods have to use highly truncated model systems to approximate typical chemical reactions. In this respect, theoretical calculations of reaction coordinates are generally projected along the modes found to be most strongly coupled to the reaction coordinate. Given the level of approximations required in treating electron correlation energies and highly simplified model structures, the observed reduction in dimensionality even within full modal basis calculations might be considered to be a consequence of the truncated moiety used to model the reaction coordinate [9, 21].

We now see that this approach can be experimentally justified for even very complex systems. There is in fact an enormous reduction in dimensionality that again is the key to how chemistry reduces to transferrable concepts in the form of reaction mechanisms [27, 71].

CONCLUSION

Integration of electron optics and pulsed laser techniques into a single experimental layout provides an effective tool for research into the structural dynamics of matter with high temporal and spatial resolution. The new techniques give rise to a very promising branch of modern physics and chemistry, advancing investigations in the 4D space-time continuum for a better understanding of both the dynamic properties of molecular systems with an intricate landscape of the potential energy surface and the mechanisms holds promise to provide a common understanding of diverse phenomena from chemical/biological process to larger length scales associated with phase transitions in condensed media, including full description of nanovolumes of solid particles.

The “pump-probe” experiments using ultrashort laser pulses provide the important capabilities for the investigation of the fast processes on the surface. However, the existing experimental techniques which are based on the probe of the spectra of the adsorbed molecules provide rather limited information on the structural dynamics. Yet, the understanding of the dynamics of the surface absorbed molecules is required for a number of the applications such as heterogeneous catalysis.

Fig. 11. Reduction in dimensionality. Dynamic observation of the photo-induced structural changes in the g charge-ordered organic system comprising ethylenedioxytetrathiafulvalene (EDO-TTF) and PF$_6^-$ counter-ions, (EDO-TTF)$_2$PF$_6$. (A) Molecular structure of EDO-TTF. (B) Representative electron diffraction pattern to illustrate the high quality of diffraction. (C) The structural changes can be mapped onto three reduced-reaction coordinates ($\xi_F$, motion of the PF$_6^-$ counter-ion; $\xi_B$, bending coordinate; and $\xi_B$, sliding motion of the rings) that stabilize the change in charge distribution, leading to electron delocalization and metallic behavior. The projections along these three normalized coordinates are highly correlated, indicating strong coupling between these nominal reaction modes. (D) Schematic depiction of the motion along these modes is given to provide a sense of the motions involved, from the insulating structure (LT), to a transient intermediate structure (TIS), to the final metallic-like structure (HT), with direction of motion indicated by the arrows and superposed structures for some sense of animation [70].
The applicability of UEC and UEnC (Part II) for the study of the structural dynamics on the surfaces, the study of the influence of the adsorbed molecules, the observation of the transitions from the crystal to the liquid phases and the investigation of the surfaces with a variety of adsorbed molecules, molecular assemblies, different nanoparticles has been experimentally demonstrated. Here femtosecond laser pulses cause the coherent restructuring of the surface layers with sub-angstrom displacement of the atoms and non-equilibrium dynamics of the surface structure is determined from Bragg diffraction spots and rings.

The electron capture by the surface causes the change of the potential and subsequently affects the motion of the nuclei inside the adsorbed molecules. The adsorption of the molecules with a lower effective electronegativity leads to a decrease in the adsorption energy and reduces the characteristic time of the dynamic response of the system to the external excitation.

It was clearly demonstrated, that UEC and UEnC (Part II) can be successfully applied to study the redistribution of the charge and energy at the interfaces. For example, UEnC allows explore the samples with the surface densities as low as 6 particles per µm², that shows the potential possibility to observe the isolated nanoparticles. Upon reaching of the submicron transverse sizes of the probing electron bunches, it should be expected the fundamentally-new results in the field of 4D dynamics, studied with high spatio-temporal resolution.

In order to understand the underlying processes of the phase changes, it is required not only the structural imaging of the samples and their interfaces in steady conditions, but also the insight into their structural dynamics, the redistributions of their structure in real time. The development of ultrafast electron diffraction, ultrafast crystallography, ultrafast electron microscopy and the dynamic transmission electron microscopy (DTEM) allows combine atomic spatial resolution, achieved in the electron diffraction techniques, with high time resolution of the femtosecond laser experiments. In this context, the "optical pumping – electron diffraction probing" has led to an extremely powerful tool for the investigation of the structural dynamic processes in different materials.

The obtained experimental data demonstrate the capability for the direct observation of the structural dynamics of condensed phase and opens up new opportunities for the analysis of chemical substance on the surfaces in 4D space-time continuum with pico-femtosecond temporal resolution.

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Ищенко А.А., Гаграшвили В.Н., Голубков В.В., Згурыский А.В. Исследование короткоживущих интермейс- 
диатов и структурной кинетики фотоизлучения молекул с помощью стробоскопической электро 


42. Оура К., Лифшиц В.Г., Сарафин А.А., Зотов А.В., Катаима М. Введение в физику поверхности. М.: Наука. 2006. 494 с.


