Ultrafast Electron Microscopy for Chemistry, Biology and Material Science

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ABSTRACT

For the past thirty years, intense efforts have been made to record atomic scale movies that reveal the movement of atoms in molecules, the fast dynamical processes in biological tissues and cells, and the changes in the structure of a solid confined to nano-scale volumes. A combination of sub-nanometer spatial resolution with picosecond or even femtosecond temporal resolution is required for such atomic movies. Additional important information can be obtained when the energy of the electron beam transmitted through the sample is measured. The four dimensional (4D) spatially and temporally resolved ultrafast electron microscopy method is made possible by the extremely high detection efficiency that is reached in 4D electron microscopy. Using ultra-short electron bunches for the visualization of biological tissue can also improve the spatial resolution compared to conventional electron microscopes, thereby enabling the study of complex biological samples of relevance to the life sciences. Of particular interest to a broad audience is the possibility to create a video, and in the future, a real atomic movie, using 4D electron tomography.

Keywords: Ultrafast Electron Microscopy; Dynamic Processes; Structural Dynamics; Atomic Movie; Femtosecond Temporal Resolution; Atomic Spatial Resolution; Electron Tomography; Spectral-Spatial-Temporal Resolution

1. Introduction

Since the 1980s, intense efforts have been made to record an atomic movie showing the movements of atoms within molecules, the fast dynamical processes in biological tissues and cells, and the changes in the structure of a solid in a nano-scale volume [1-3]. Such an atomic movie requires a combination of sub-nanometer spatial resolution with picosecond or even femtosecond temporal resolution.

The aim of any microscopy is to study the structure, composition, and a variety of physical and chemical characteristics of samples (usually solids) on a very small length scale, with linear dimensions of irregularities that are on the order of micrometers or nanometers. While in most microscopies, photon beams (in optical microscopy) or corpuscular beams (for example in electron microscopy) are used as the probes, tunneling microscopies use a very sharp tip that is scanned across surfaces. Electron microscopy and optical microscopy use light and a focused electron beam for imaging, respectively. Beyond those, other microscopic methods may use ions, protons, neutrons, or acoustic or microwave radiation in addition to other, less common, methods [3-6]. In each of them, the specificity of the interaction of a beam of the particles (or photons) and the molecules or atoms of a sample yield unique and rather useful information about the structure, composition and microscopic inhomogeneities of the sample, and the nature of their intermolecular interactions [3,4,7]. For example, the slow neutrons in neutron microscopy and spectroscopy barely interact with the electrons, but interact strongly with the nuclei of the atoms.

There are two main approaches to achieve imaging in classical microscopy: 1) In transmission mode, a large area of the sample is illuminated by a beam of light or particles. Specifically designed lenses project the transmitted beam onto a screen or detector to form an image of the structure of the sample. 2) In reflection mode, the beam reflected from the sample, or secondary particles or photons generated by the incident beam in the sample, form an image of the sample surface or a thin region underneath the surface [8].

Because of absorption of the optical radiation by matter, transmission optical microscopy is limited to thin
samples, typically less than 1 mm thick. In electron microscopy, the sample has to be much thinner, because the electrons interact with matter more strongly than light. In transmission electron microscopy it is therefore possible to get images only of very thin slices, with film thicknesses of much less than 1 micrometer. Such samples have to be prepared using special ultramicrotomes that can produce film thicknesses down to a few tens of nanometers.

There are two different ways to obtain an image [9,10]: 1) “all at once”, when the sample is irradiated entirely with light or electrons and the transmitted or the reflected beam is detected at once using, for example, a film, photographic plate, photosensitive recorder or digital camera; 2) in “scanning microscopy” mode, where the sample is scanned by a beam focused to a spot of rather small diameter, and the resulting image is assembled point by point.

2. Transmission and Scanning Electron Microscopy

In conventional scanning microscopy, images are formed by a beam of charged particles or photons that is focused to a very small diameter. This probe is moved across the surface of an object, and signal of any physical nature, but in particular reflected particles or photons, is detected and the image obtained “point by point”. The method is quite analogous to the generation of an image using a scanning electron beam in a traditional cathode ray tube (CRT), or in modulating the brightness of each luminous point in a LCD monitor [11]. Scanning microscopy is a rather convenient tool to study the morphology and topography of various geometric objects, to investigate elemental compositions, to measure the electric and magnetic fields in micro-volumes, (the so-called “micro-fields”, as introduced by Prof. GV Spivak), or to test electrical parameters of different semiconductor crystals (e.g., lifetime and diffusion length of charge carriers).

In scanning electron microscopy, the scan of the electron beam should be synchronous with a sweep of the probe on a sample. Like in a TV image, the raster image consists of individual small dots or pixels, the brightness and make-up of which, formed after signal amplification, define the image. Thus, the elements of the image appear one after the other, as if we move our fingers to spell a complicated book of life, but not all at once, as in the usual optical or transmission electron microscopes. This “point-wise” map of the object greatly facilitates the processing and interpretation of images [12].

In a transmission electron microscope, the electrons illuminate and pass through a very thin sample. An image of the sample is formed by means of electrostatic or magnetic lenses on the fluorescent screen or equivalent position sensitive detector [13]. In 1931, the first such device was demonstrated by M. Knoll and E. Ruska, who imaged a wire mesh using an incident electron beam and axially symmetric magnetic electron lenses with a narrow annular gap and a sharp maximum of the magnetic field on the axis. The first commercial transmission electron microscope was developed and released by “Siemens” in 1939 under the supervision of E. Ruska, who received the Nobel Prize for Physics in 1986 for his invention of the electron microscope.

The transmission electron microscope (TEM) is the now the most common tool, designed to study very thin samples with thicknesses of about 10 - 100 nm at electron accelerating voltages of up to 200 kV [14]. Its scheme is similar to the optical system of an optical light microscope in that it creates an image of the sample based on the transmitted radiation beam. Figure 1 compares the main units of the transmission electron microscope to those of the optical microscope and the scanning electron microscope.

The transmission electron microscope consists of an electron gun and electromagnetic lenses. Figure 1 shows the path of the rays in a microscope with three levels of zoom and one condenser lens, which is used to pre-focus of the electron beam. The two common modes of its operation are the imaging mode and the recording of micro-diffraction patterns. In imaging mode, the plane of a sample conjugates to the screen, where it is assumed that the sample is very thin compared to the focal length of the imaging objective lens. In the diffraction mode, the object of observation conjugates to the back focal plane of the objective lens, where the distribution of the amplitudes of the electron beam, diffracted off the sample, corresponds to the Fraunhofer’s diffraction of the electrons. The resulting picture is observed on a remote screen. It is possible to determine the composition of the sample by analyzing the diffraction pattern [16].

In the phrase “micro-diffraction”, the word “micro” implies that the diffraction signal is observed only from a rather small area of the sample. The area is chosen either by a special selector diaphragm, installed in the plane of the first intermediate image (see Figure 1), or by irradiation of the desired part of the sample with a sharply-focused electron beam with a diameter of about 1 micrometer or less.

In imaging mode, the so-called aperture diaphragm is installed in the focal plane of the object lens in order to limit the aperture of the beam, i.e. the opening angle of the cone of rays emerging from the sample. This helps to decrease the spherical aberration, a basic imaging error associated with the projection of a point in the sample to a point of the image.

Modern microscopes are usually equipped with a two-lens condenser, which allows for a rather high electron
flux to a small area of the object with characteristic size of 1 - 5 microns. This eliminates the growth of the films due to hydrocarbon polymerization of oil vapors as a result of electron bombardment of the unobserved sample areas in microscope where the column is pumped by diffusion pumps.

According to statistical data, tens of thousands of transmission microscopes with accelerating voltages of 80 - 100 kV have been produced. Hundreds of devices with an accelerating voltage of between 200 kV and 500 kV were created for the study of thicker objects, with a thickness of up to 10 microns. And there are only a few dozen microscopes intended for the voltage range from 1 MV to 3.5 MV.

For a typical transmission electron microscope with an accelerating voltage of 100 kV, the de Broglie wavelength of the electron is 3.7 pm. This is much smaller than the diameter of a hydrogen atom, which equals approximately 100 pm. This would seem to make it easy to observe any atomic particle, as long as the resolution of the electron microscope is on the order of the electron wavelength as is the case in optical microscopes. But in fact, it is extremely difficult to observe individual atoms because of the poor quality of the electron lenses compared to the light-optical lenses: at the same angular size of the light and electron beams, the electron lenses create almost of two orders of magnitude more aberrations and image distortion [13].

At a kinetic energy of 100 eV, the de Broglie wavelength of the electron is equal to 0.1 nm, i.e. on the order of atomic dimensions. Electrons with such energies are strongly diffracted by atomic lattices. This property is used in low-energy electron diffraction to investigate the structure and composition of surfaces and near-surface layers, because the penetration depth of the electrons is very small.

3. Stroboscopic Electron Microscopy

Electron microscopy takes advantage of two of the three main properties of the electron: the small de-Broglie wavelength of the electron, which enables very high spatial resolution; and the electric charge, which allows to control its movement by electric and magnetic fields. About 60 years ago, the third property of the electron, its small mass (i.e., low inertia) began to be used in so-called stroboscopic electron microscopy for the study of periodic processes [17]. The essence of the method is quite simple: a sample with periodically time-varying characteristics, with a fixed rate of change, should be irradiated by short bunches of electrons with the same frequency and fixed phase. The synchronicity freezes the image, which corresponds to the state of the sample at the time of arrival of the electron pulses.

In the early 60’s, in the Soviet Union as a result of the work of Prof. G.V. Spivak, and in other countries, stroboscopic techniques were developed for all major types of electron microscopes: translucent, raster, emission and mirror types [18,19]. These developments allowed the exploration of processes in rapidly varying thin films (transmission electron microscopy) and in the surface layers of solids (raster, emission and mirror microscope): re-polarization of various ferroelectrics; distinctions of the domain structure during re-magnetization of thin magnetic films; heterogeneity of alternating fields of...
magnetic heads; and local defects of p-n junctions during their fast switching between the locked and open states [20]. Additionally, from a theoretical point of view, it is possible to study the microstructure of solution-gels in periodic reactions such as the Belousov-Zhabotinsky reaction using electrons with low kinetic energies, less than 100 eV, and to investigate the periodic sorption-desorption processes of catalysis in a similar fashion [20].

The initial work on stroboscopic microscopy was performed with micro- and nanosecond time resolution. Later, several laboratories started to investigate periodic processes with picosecond temporal resolution, usually using stroboscopic scanning electron microscopy. The main difficulties arose from the gating, the interruption, or the modulation of the electron beam. Generally, one uses one of two methods: first, it is possible to change of the intensity of the beam by locking or unlocking it with appropriate voltage pulses applied across the cathode or across the modulator in the electron gun of the microscope; or secondly, one can deflect a continuous electron beam by an electric or magnetic field in the vicinity of a skipping aperture of small diameter. The blanked, locked or unlocked electron pulses should be properly frequency and phase synchronized with the signals on the sample in order to avoid a loss of resolution.

The larger the duty cycle of the blanking pulses, i.e., the higher the ratio of the repetition period to the duration of the front of the electron pulse, the better the temporal resolution of the stroboscopic microscope, but the lower the brightness of the image. Consequently, each application requires a careful assessment of the best trade-offs. Moreover, a very short front of the pulse causes the resolution of the microscope to deteriorate because of non-ideal edges of the pulse, the appearance of chromatic aberration in transmission electron microscopy and the manifestation of the Bursch effect, which describes the transition of the longitudinal component of the electron velocity to a transverse component due to Coulomb repulsion.

4. Dynamic Transmission Electron Microscopy

Transmission electron microscopy, with its rather wide range of its tools, has long been a powerful technique in many fields of science and technology. While allowing to reach a spatial resolution on the sub-nanometer scale, conventional transmission electron microscopy does not permit an ultrashort temporal resolution, i.e. a time resolution in the picosecond or femtosecond regime. In contrast, optical microscopy using fluorescent probes, such as green fluorescent proteins, made it possible to visualize the processes occurring in vitro [2]. However, despite the high temporal resolution of optical methods, their spatial resolution is usually limited by the wavelength to the range of 200 - 800 nm. A review article [21] and a recently published monograph [2] describe the development of the methods of 4D, dynamic transmission electron microscopy (DTEM) and ultrafast electron microscopy (UEM), respectively.

The images and diffraction patterns in the work [22] were obtained at an accelerating voltage of 120 keV for non-organic materials (single crystals of gold, amorphous carbon and polycrystalline aluminum) and for biological cells of the intestine of rats [23]. In those studies, the gated beam contains on the average only one electron per pulse, with the total dose being limited to a few electrons per Â². Note, that these electron pulses were completely controlled in space and in time. Conceptually, this works is based on the methods of time resolved electron diffraction (TRED) or ultrafast electron diffraction (UED) [1,24-26,78,112], ultrafast electron crystallography (UEC) [23] and the other DTEM studies [21], but with one important difference-namely, the implementation of synchronized pulses of single electrons to form an image. The sketch of this 4D DTEM is shown in Figure 2.

This single-electron approach differs quite remarkably from the one used in the work [27], which used single pulses with $10^8$ electrons and pulse durations of ~20 ns.

![Figure 2. The schematic of a dynamic transmission electron microscope (DTEM) [21].](image-url)
Those pulses, used to study laser-induced melting of different metals, contain so many electrons that high temporal resolution is not possible. Moreover, as noted in [28], the uncertainty in the spatial resolution in these experiments due to statistical interferences was of the order of micrometers, because the temporal window for imaging was on a nanosecond scale. It was found, that the use of amplified femtosecond light pulses, with much higher intensity of about $10^{12}$ W/cm², led, at first, to the formation of an electron beam that was insensitive to the focusing system of the microscope, and, secondly, to a significant lengthening of the pulse, which was due to electron-electron Coulomb repulsion.

The single-electron per pulse approach of 4D UEM [22,29] was used to obtain images from polycrystalline and single crystals [2]. To verify the authenticity of the experiment, some images were collected when the femtosecond pump laser pulses were blocked. In this case, the pattern was not observed, confirming that the time resolved signal generated in the electron microscope was indeed induced optically, and that the contribution of thermal electrons that are continuously generated by the cathode can be neglected [29]. It is useful to remember that electron microscopes can operate in UEM and DTEM mode. To reach atomic scale spatial resolution, the UEM diffraction regime should be used by introducing an intermediate lens to select the back focal plane of the lens as the object.

5. Controlling the Motions of Free Electrons by Femtosecond Light Pulses

Femtosecond lasers play a key role in 4D electron microscopy. Ultrashort pulses are necessary for the optical excitation of ultrafast processes in the sample and to form the ultrashort probe electron pulses. Additionally, pulsed laser radiation makes it possible to measure the temporal characteristics of the photoelectron bunches, to implement additional compression of electron pulses, and to generate electron bunches with durations that may potentially reach into the attosecond regime.

The last statements need to be amplified on, because they imply that is possible to control free electrons with a laser beam. At first glance, this statement conflicts with the well-known fact that free electrons cannot absorb the electromagnetic (EM) radiation because the laws of conservation of energy and momentum cannot simultaneously be satisfied. Nevertheless, free electrons can scatter EM radiation. From a quantum mechanical point of view, this is the result of stimulated Compton scattering in a strong laser field.

The possibility to control the translational degrees of freedom of charged particles, in particular of free electrons, by a spatially inhomogeneous electromagnetic field was demonstrated theoretically by Gaponov and Miller in the mid 1950’s [30]. This mechanism leads to the ejection of charged particles from the strong field. In the case of high-power laser radiation, the Gaponov-Miller, or ponderomotive (gradient) force (PF) [31], which is defined as the spatial gradient of the so-called ponderomotive potential $U_p$, can reach sufficiently high values to form a basis for the effective control of electron pulses in vacuum.

For non-relativistic electrons, the concept of the ponderomotive potential in a spatially inhomogeneous electromagnetic field $E = E_o (r) \sin(\omega t)$ can be introduced by averaging the Hamiltonian $H$ over the fast oscillations with frequencies $\omega$ and $2\omega$:

$$H = \left\{ (p + eE_o (r) \sin(\omega t)/\omega)^2 \right\}/2m_e$$

$$= p^2/2m_e + \left[ eE_o (r) \right]^2/4m_e \omega^2$$

Here, $m_e$ and $e$ are the mass and the charge of the electron, and $\langle \cdots \rangle$ denotes an averaging over optical cycles. As a result, the expression for the PF can be written as the following expression:

$$F^{(\text{pond})}(r,t) = -e^2 \lambda^2 / 64 \pi^2 m_e \epsilon_0 c^3 \nabla V(r,t)$$

Here, $\epsilon_0$ is the dielectric constant, $c$ the speed of light and $\lambda$ the wavelength of the laser radiation. It follows that for a tightly focused laser pulse with an intensity of $10^{12}$ W/cm² in the center of a focal spot of 2 µm diameter (at the level of 1/e) and $\lambda=800$nm, the force, $F^{(\text{pond})} \approx 10^{-11}$N, is approximately equal to the strength of the interaction between two electrons separated by a distance of 5 nm. In this example, the ponderomotive potential is $U_p \approx 60$eV.

It is evident that ultrashort laser radiation allows to control free electrons in vacuum on a femtosecond scale. This makes possible both the creation of ultrashort photoelectron bunches and the measurement of their duration [32,33]. The determination of the temporal characteristics of a pulsed electron beam is based on scanning the time delay between the laser radiation that forms the photoelectric bunch and the tightly focused laser beam that changes the velocity distribution of the photoelectrons as a result of the PF. Such a method compares favorably with a standard streak camera in its final characteristics and can be used for femtosecond electron beams [33].

In a streak camera, the electrons are deflected by a high-speed and high-voltage electric field with an amplitude of about 2 - 5 kV before reaching a position-sensitive detector. The rate of change of the high electric field determines the temporal resolution of the device. The currently best time resolution is in the subpicosecond range and reaches a value of $\sim 300$ fs. In addition
to this, it should be noted that the femtosecond temporal synchronization of the high-voltage electrical pulse in the streak camera and the femtosecond laser pulse that forms the ultrashort photoelectron bunch poses serious technical challenges.

The study of the behavior of the free electrons in spatially inhomogeneous electromagnetic fields began shortly after the appearance of the theory of Gaponov and Miller and the first experiments were devoted mainly to the possibility of creating traps using microwave technology [34]. Here, special attention was focused on the observation of the passage of the electrons with a certain kinetic energy through the ponderomotive potential, which allowed, for example, to determine the value of $U_p$ [34,35]. The first demonstration of scattering of the photoelectrons by the ponderomotive potential, created by an intense sub-nanosecond laser pulse, was done in the work [36]. Here the multi-photon ionization of Xe, especially bleeding in the vacuum system, was used to prepare the pulsed photoelectron beam with kinetic energies less than 5 eV. This experiment demonstrated the control of low-energy photoelectrons by an optical ponderomotive potential with the height of about 10 eV.

Special attention should be given to the measurement of the value of the $PF$ with which tightly-focused femtosecond laser radiation acts on the electrons propagating in vacuum. This may be useful for in situ space-time diagnostics of the laser fields of high intensity, and can be used to validate the measurement of ultrashort electron pulse durations based on the irradiation of the electron bunches with the laser pulses of high intensity. Here it should be noted that the use of laser radiation with intensities of more than $10^{14}$ W/cm$^2$ usually is accompanied by the photoionization of the residual gas in the vacuum system and thus by a possible deformation of the electron trajectories that pass through a cloud of charged particles.

As was noted above, electrons are ejected from their field-free paths by a strong electromagnetic field. This is illustrated in Figure 3, which shows two counter-propagating tightly-focused laser beams forming a standing wave from which the electron pulse is deflected. Such a scheme has a dual purpose. First, it allows to determine the duration of the initial electron bunch by scanning the timing of the laser pulses that create the standing wave, and using a position-sensitive scheme to detect the rejected, but in general, diffused, electron beam components. Secondly, the deflected electron pulses potentially have a shorter time duration compared to the incident electron pulses.

Two important remarks should be made here. The scheme in Figure 3 requires the spatial-temporal matching of two femtosecond laser pulses from two directions and a (sub)picosecond electron bunch from another direction. The use of a single laser beam for the electron scattering would, of course, be simpler. However, in the standing wave the spatial inhomogeneity of the electromagnetic field is about $\lambda/2$, a rather small value that is technically difficult to obtain with a single amplified femtosecond laser beam. For example, a parabolic mirror can usually focus 0.8 - micron amplified femtosecond laser radiation to a spot diameter of $d_p \approx 6$ microns. Therefore, for a given laser pulse energy, laser wavelength and focusing conditions, the scattering of electrons in a pulsed standing wave will be more pronounced.

Another important consideration relates to the ultimate possibilities of this approach. This is important because it is rather interesting to know the accuracy of the electron pulse duration measurement as well as the duration of the deflected electron bunch. These characteristics are directly related to the transit time of the fast electrons through the laser focal spot,

$$r_{fin} \approx r_p = \frac{d_{fin}}{v_e},$$

where $v_e$ is the speed of the electron beam. Using $d_{fin} = 6$ µm and $v_e \approx 10^7$ m/s for electrons with kinetic energy of 30 keV, we find $\tau_u \approx 60$ fs. Note that the use of a standing wave formed by two laser beams (Figure 3) will not lead to a decrease in $r_{fin}$ and hence the accuracy of measurement for this example will remain of the order of 100 fs. Therefore, to obtain a femtosecond electron bunch with $\tau \sim 10-50$ fs and its full diagnostics it is important to ensure the tight focusing of intense laser radiation, and/or to use electrons with higher kinetic energy. Of course, in order to deflect these ultrafast electrons it is necessary to implement intense laser fields.

An alternative way of creating ultrashort electron bunches also employs a standing light wave, but in a collinear geometry [2]. This approach is shown schematically in Figure 4.
Figure 4. Schematic diagram of the creation of attosecond electron pulses. (a) An electron bunch with sub-picosecond duration is irradiated by two counter-propagating laser beams (forward and after the bunch) with different wavelengths. In the reference frame of the electrons, the wavelengths are the same as a result of the Doppler effect, and a standing light wave is formed; (b) Ponderomotive forces push electrons away from the strong laser field; (c) As a result, the compression leads to the formation of electron bunches with attosecond duration. Strictly speaking, the envelope duration of the initial electron bunch remains constant, but importantly, the attosecond spikes appear inside the envelope [2].

In this context it should be recalled that on the basis of the uncertainty principle, $\Delta E \Delta \tau \geq \hbar$, the existence of ultrashort, in this case attosecond, bursts needs a wide spectrum of energy $\Delta E_e$. For example, a bunch with 100 as ($10^{-16}$ s) duration has at least $\Delta E_e \approx 6$ eV. The initial picosecond photoelectron pulse does not exhibit such spectral properties. As shown in Figure 4, the required broadening is due to the work of the ponderomotive forces [2]. Let us illustrate this with a simple estimate, by writing down the expression for the work in the traditional sense, where the energy gain of the electrons is equal to $\Delta E_e = F^{\text{(pond)}} \tau_p v_e$ ($\tau_p$ - the laser pulse duration), and the force is approximately defined as $F^{\text{(pond)}} \approx 2U_p / \lambda^* \ (\lambda^* - \text{wavelength of the laser radiation in the reference frame of the fast electron})$. In the conditions of the numerical experiment [2] $U_p \approx 0.2$ eV, $\lambda^* = 370$ nm, $v_e \approx 10^5$ m/s and $\tau_p = 300$ fs, one finds $\Delta E_e \approx 32$ eV. The calculated value $\Delta E_e$ exceeds by 2 orders of magnitudes the spectral width of “normal” sub-picosecond electron pulse, emitted from the solid photocathode under the action of femtosecond laser radiation.

As considered in the monograph [2], this scheme is rather elegant. Even so, it is important to emphasize that the measurement of the duration of the formed attosecond electron bursts is a quite a challenging experimental problem. It is likely that it belongs to the class of experiments in which it is easier to create an ultrashort bunch than to measure it.

6. Ultrafast Electron Microscopy

An early antecedent of modern methods to investigate fast processes is the English photographer Eadweard Muybridge, who used multiple cameras to temporally resolve fast motions. For example, in 1871 Muybridge captured the moment when a galloping horse separated all four hooves from the surface of the earth (Figure 5).

Of interest to the scientific world today would be movies showing the movement of atoms in molecules, the fast dynamical processes in biological systems, or the changes in the geometrical structures of nanoscale devices. To estimate the magnitudes of time scales involved, we can extrapolate from the observation of macroscopic objects by using as the maximum velocity the speed of sound. Thus, a time resolution of 1 μs suffices to freeze in time motions of objects with spatial dimensions of 1 mm. In a system with a characteristic length scales of nanometers, a picosecond time resolution is appropriate. And, for molecular systems, femtosecond time resolution is necessary to capture the motions of individual atoms and functional groups.
The ultrafast electron microscope built at the California Institute of Technology [2] is designed to explore a class of problems that occur on picosecond or sub-picosecond time scales. A schematic illustration of this instrument is shown in Figure 6.

It is convenient to conceptualize the setup as consisting of three main sections: 1) the formation and acceleration of the sub-picosecond pulsed electron beam; 2) the interaction unit, in which the sample is excited by femtosecond laser pulses and its spatial-temporal structure probed by ultrashort electron bunches; and 3) the position-sensitive detection of the electrons after their interaction with the sample. The use of electron beams of subpicosecond duration imposes strict requirements on the detector, because the probe pulse cannot contain a large number of electrons. As it will be illustrated below, the main cause of the electron pulse temporal spreading is the space-charge interaction between the electrons within each pulse on account of their Coulomb repulsion. Therefore, the detection efficiency for electrons in this microscope should be as high as possible. Much preparatory work was done to achieve this [2]. As a result, the detection efficiency for the time-resolved instrument is almost an order of magnitude higher than that of a standard electron microscope. This also provides an interesting opportunity to significantly reduce the exposure (the total number of electrons passing through the sample to form an image on the detector), which is particularly important for imaging biological samples that are very sensitive to radiation induced damage.
The ultrashort electron bunch is the essence of 4D electron microscopy. Therefore, the question of its duration should be given special attention. The duration of the photoemission of electrons from the surface of a solid cathode and therefore the initial duration of the photoelectron bunch are determined by the duration of the laser pulse. But during the propagation from the photocathode to the target, the electron beam is stretched in time. Here we consider three major factors that lead to a smearing of the electron bunch.

First, the acceleration of photoelectrons in a static electric field near the cathode causes so-called time-of-flight chromatic aberration for the electrons (TFCA). The temporal resolution limit is given by:

$$\Delta \tau_{EF} = \sqrt{2m_e \Delta E_e / eF},$$

where $\Delta E_e$ is the initial spread of the kinetic energy of the electrons as they emerge from the cathode and $F$ is the electric field in the acceleration gap.

The expression for TFCA can be obtained by solving a quadratic equation that describes the uniformly accelerated motion of the electrons in the acceleration gap of the length of $l$:

$$v_0 + (eF/m_e)t^2/2 = l$$

Differentiating the both sides of the equation with respect to time $t$, we get:

$$\Delta v_0 + (eF/m_e)\Delta t \approx 0$$

from which immediately follows the expression for TFCA. Let us consider a numerical example. At $\Delta E_e = 0.3$ eV and $F \approx 10^8$ V/m, which approximately corresponds to the maximum value of the field (higher values lead to vacuum breakdown even with special electrodes that underwent thorough mechanical, electronic and ionic treatment), this contribution is $\Delta \tau_{EF} \approx 60$ fs.

Secondly, the movement of electrons through the field free flight tube with slightly different kinetic energies is accompanied by an elongation of the time of flight, which is expressed as

$$\Delta \tau_{TOF} = t_{TOF}(\Delta E_e / 2K)$$

Here, $t_{TOF}$ is the time of flight of the electrons with kinetic energy $K$ from the photocathode to the sample. At $K = 30$ keV, $\Delta E_e = 0.3$ eV and $t_{TOF} \approx 3$ ns, corresponding to the length of 30 cm between the cathode and the target, we get $\Delta \tau_{TOF} \approx 15$ fs.

Thirdly, the Coulomb repulsion between the electrons within a pulse causes a swelling of the photoelectron bunch. To assess the effect of this mechanism on the temporal properties of the beam, we consider a simple model, in which the electron bunch is travelling through a field-free region. (Here the field-free region is taken for the simplicity.) The total energy of the bunch, consisting of $N$ electrons, is conserved:

$$V + W = 1/2 \left( \sum e^2 / 4\pi e_0 |r - r_j| \right) + 1/2 \sum m v_j^2$$

Therefore, the characteristic value of the distribution of the velocities, $\delta v$, resulting from the electrons pushing apart each other can be qualitatively estimated by using the expression

$$m_e \delta v^2 / 2 = e^2 N / (16\pi e_0 \delta r),$$

where $\delta r$ is the initial size of the photoelectron cloud. In this model, the velocity spread of the electrons causes the elongation of the order of $\Delta \tau_c \approx t_{TOF} \delta v / v_e$, which can be rewritten as:

$$\Delta \tau_c = (t_{TOF} / v_e) \sqrt{e^2 N / (8\pi e_0 \delta rm_e)}$$

At $v_e \approx 10^8$ m/s, $N \approx 10^8$, $t_{TOF} \approx 3$ ns and $\delta r \approx 10$ mm, corresponding to the size of the focal spot of the tightly focused laser beam, this contribution in the framework of the simple model is equal to $\Delta \tau_c \approx 5$ ps. This shows that the Coulomb repulsion of the electrons in the electron bunch is the main factor limiting the ability of ultrafast electron microscopy. Naturally, the use of ultra-high accelerating voltages, such as 500 kV instead of 30 kV, can reduce the swelling of the electron bunch due to Coulomb repulsion, but can also lead to destruction of the sample. Use of very fast electrons, in the MeV range, can further reduce the space-charge broadening because of the onset of relativistic effects [110,111]. It should also be noted that a system based on a TOF electro-static mirror, a so-called reflectron, can be used to re-compress the electron bunches after their original spreading [26,38]. At the present time, a radio frequency compression scheme is already in use for this purpose.

A serious challenges in the field of 4D electron microscopy remains the task of observing an image or a diffraction pattern produced by a single, ultrashort electron pulse. The solution to this problem is rather important, because it would enable the study of ultrafast irreversible processes. An early breakthrough has been achieved by a collaboration of scientists at Brown University and at SLAC, who used ultrashort electron pulses with 5.4 MeV of energy, generated using the Gun Test Facility at SLAC, to record single-shot diffraction patterns of 160 nm thick aluminum foils [110,111]. Their work showed that MeV electrons should make it possible to achieve sub-100 fs time resolution. Subsequently, physicists in Japan were able to collect diffraction patterns of thin gold films with single subpicosecond electron bunches [39]. They applied a magnetic sector in order to compress the electron beam in the experiment [39].
Non-Thermal vs Thermal Melting of a Solid Irradiated by a Femtosecond Laser

The study of ultrafast phase transitions in condensed matter is one of the most interesting fields of 4D electron microscopy, viz. ultrafast electron crystallography. To induce an ultrafast phase transition such as the melting of a solid, it is necessary to insert energy into the lattice within a small period of time and to reach the critical temperature as quickly as possible. The most suitable tool to accomplish this is a femtosecond laser pulse. In metals and semiconductors, the photons excite the electrons (and the holes) to high-lying electronic states, resulting in a change of the charge density. Initially, the absorbed energy is stored in electronic degrees of freedom. With the decrease of electron density between the atoms, the bonds in the system become weaker, and at some degree of excitation, the lattice is no longer fixed. Such processes may lead to non-thermal melting of a solid, provided that they are fast compared to the relaxation of the electron energy into the lattice vibrations (phonons). Preliminary studies of aluminum by a femtosecond laser pump-probe have shown that in this material, the transition from a solid to a liquid phase occurs within 500 fs, i.e. faster than the energy transfer from the excited electrons to the phonons [40].

The direct observation of melting of Al with subpicosecond temporal resolution has been achieved by ultrafast electron diffraction [41]. A 20 nm thick film of polycrystalline aluminum was used as the sample in this experiment. The results are shown in Figure 7.

Figure 7 clearly demonstrates, that the lattice is heated and remains in the solid phase with a distinct diffraction pattern on a time scale of up to 1.5 - 2.5 ps after the excitation by laser pulse, which is consistent with the thermal melting. Current research continues in this direction.

7. Phase Transitions in Nanoparticles

Vanadium dioxide, VO$_2$, undergoes a phase transition of the first type from a low-temperature monoclinic phase (M) into a high-temperature tetragonal phase of rutile (R) at $-67\degree$C. This phase transition has been the subject of intensive research since its discovery almost half a century ago. The work [29] describes the first study of this material using 4D UEM. Because single electron probe pulses were used, Coulomb (space-charge) repulsion was absent. The experiment demonstrated the possibility of obtaining a sequence of images (i.e. a movie) with atomic-scale spatial resolution combined with an ultrashort temporal resolution. In particular, it was shown that it was possible to investigate the ultrafast metal-insulator phase transitions in vanadium dioxide. A structural phase transition in VO$_2$ nanoparticles is exhibited in the diffraction patterns (at atomic scale) and UEM images (at nanometer scale) with a characteristic hysteresis with a temporal resolution on the order of 100 fs (Figure 8).

Figure 7. Electron diffraction patterns from a film of polycrystalline Al, recorded at different times after the excitation by a femtosecond laser pulse as labelled. The energy density of the laser pulse is about 70 mJ/cm$^2$ [41].
4D UEM with femtosecond laser irradiation of the sample in the near IR allowed to record the movement of atoms during the phase transition in all three dimensions and in time. This study showed that at first, the vanadium atoms were separated from each other for a short time period, and then began to move towards their final positions.

8. Visualization of Mechanical Displacements of a Cantilever

The development of ultrafast electron microscopy made possible the direct observation of vibrations of nanostructures in the ultrasound range, covering the range from kHz to GHz [42]. In this connection it should be recalled that the methods of optical interferometry cannot ensure complete three-dimensional spatial resolution, while the possibilities of standard electron microscopy are usually limited to a temporal resolution of about 30 ms.

The work [42] used a pseudo-one-dimensional molecular material, crystal Cu (TCNQ), which exhibits highly anisotropic electrical and optical properties, to prepare a nano- and microcantilever with a thickness of 300 nm and 2 mm, respectively. Opto-mechanical motions can be initiated in such a crystal by a charge transfer from the radical (TCNQ)− to copper Cu+.

In the experiment, the charge transfer occurred as a result of irradiation of the crystal by nanosecond laser pulses with a wavelength of 671 nm, a pulse intensity of about 160 mJ/cm² and a pulse repetition rate of 100 Hz. This exposure induced resonant cantilever vibrations in the megahertz frequency range. The tiny displacements of the tip were visualized using the 120 keV photoelectrons (Figure 9).

The results so obtained allowed to determine the Young’s modulus, measure the resonance frequency of the sample, and to calculate the energy stored due to the opto-mechanical expansion of the crystal Cu (TCNQ).

9. 4D Microscopy of Fatty Acids—Toward the Study of Cell Tissue Materials

The study of organic and biological structures is an extremely important field of electron microscopy. However, the investigation of such objects entails special
demands on the microscope. This is primarily due to the increased sensitivity of the bio-structures to possible irreversible deformation and damage that can be caused by a beam of fast electrons. Therefore, for successful experiments in this field it is necessary to ensure an extremely high detection efficiency of the electrons passing through the sample. Note that the efficiency of the detector is directly related to the time of exposure. In this connection it is helpful that the high detection efficiency in the 4D microscopy allows working with a smaller number of electrons in the bunch.

Amongst the most common biological structures are membrane-type layers of fatty acids. In the laboratory, two-dimensional crystals can be created from long fatty acid hydrocarbon chains. It is possible to use the Langmuir-Blodgett technique to prepare the monomolecular layers of organic material on a solid substrate. It is rather interesting to observe the transition from a two-dimensional to a three-dimensional crystal of the long molecular chains and to visualize the self-assembly processes.

C_{20}H_{42} chains with a length of 95Å on a silicon substrate were considered in computer simulations [2]. In the model, an ultrafast pulsed heating of the substrate leads to further changes in the inter-atomic distances of C_{20}H_{42}, reflecting the coherent motion of the atoms in the molecular chain. Each step of the calculation corresponds to a time interval of 0.5 fs, while the total number of steps reaches 200,000. Model calculations have shown an increase in the length of the fragment -CH_{2}-CH_{2}-CH_{2}- that is located in the immediate vicinity of the silicon surface, by about −0.08 Å during 5 ps.

The 4D UEM method should allow us to study more complex biological samples that are directly related to the life sciences. It should be recalled that the use of ultra-short electron bunches for the visualization of biological tissues can improve the spatial resolution of the device compared to conventional electron microscopy. In some ways the situation is similar to making a snapshot by a photographic camera with a long exposure time: the resulting image becomes quite fuzzy (Figure 10), but admittedly, can lead to a new artistic perception of reality.

The 4D electron microscopy succeeded in imaging stained cellular material in rat tissue (Figure 11), and of crystalline proteins, or cells in the vitreous bodies.

10. Musical Nanoscale Instruments: from the Drum to the Harp and the Piano

Ultrathin plates of graphite, gold, or other material, subjected to pulsed laser heating, can experience an ultrafast deformation (bending) as a result of anisotropic expansion of the atomic structure. As it turned out, the vibrations that accompany such deformations may be surprisingly interesting.

Electron bunches at 200 keV were used to visualize mechanical vibrations of graphite nano-sheets (the membranes) with thicknesses of 75 nm. Vibrations with submillisecond temporal intervals resulted from exposure to pulsed laser radiation with a wavelength of 532 nm and a repetition rate of 5 kHz [2]. The energy density of the focused laser pulses reached a value of about 7 mJ/cm².

Immediately following the pulsed laser heating of the
membrane, individual carbon atoms were found to vibrate in a random order. But after a few tens of microseconds, the corresponding modes of the membrane began to be synchronized (Figure 12), resembling a “drumbeat” with a frequency of around 1 MHz.

The processing of the images (Figure 12) used a cross-correlation function defined as

$$\gamma(t; t') = \left[ \sum C_{x,y}(t) C_{x,y}(t') \right] / \sqrt{\sum C_{x,y}(t)^2 S_{x,y}(t)^2}$$

where the contrast is

$$C_{x,y}(t) = \left[ N_{x,y}(t) - \langle N(t) \rangle \right] / \langle N(t) \rangle.$$

Here, $N_{x,y}(t)$ is the number of counts corresponding to the picture element (pixel) with the coordinates of $(x,y)$, and $\langle N(t) \rangle$ is the average number of the counts.

2000 images, collected with an interval of 50 ns, were used in Figure 12. The micrograph of the graphite membrane is represented in the lower left corner.

The success with the graphite nano-sheets inspired the search for other musical nano-instruments. A harp and a piano, constructed at the nanoscale, were made using arrays of cantilevers (Figure 13). They were prepared by micro-structuring of multilayer workpieces of Ni/Ti/Si₃N₄ using sharply-focused ion beams [44]. The workpieces consisted of 30 nm layers of nickel and titanium, consistently applied to 15 nm film of Si₃N₄. For the piano, the cantilevers had almost the same lengths, approximately 4.6 μm, but their width varied from about 400 nm to 2.3 μm. In the harp, the lengths and widths of individual elements varied in the range of 1.2 - 9.1 μm and −300 - 600 nm, respectively.

In the experiment, a sample (the harp or the piano) was heated by nanosecond 519 nm (or 532 nm) pulses from a Nd:YAG laser with a pulse repetition rate of 1 kHz. The energy density per pulse reached 2 mJ/cm². As a result of the heating of the sample, consisting of the layers with different coefficients of thermal expansion, the mechanical deformation of the cantilever triggered its vibrations. The mechanical movements were observed using the photoelectrons that were obtained by illuminating the photocathode with the fourth harmonic of a Nd:YAG laser. The results for individual cantilevers p1, p5, h1 and h5 (for convenience they are shown in Figure 13) are presented in Figure 14. The combined results of the observations are given in Figure 15.

In the next series of the experiments, the femtosecond laser with MHz pulse repetition rate was used “to play an instrument.” Here a fiber laser generator with pulse repetition frequency of about 25 MHz was used. The energy density per pulse was in the range of 20 - 30 mJ/cm².

Beyond their academic interest, the musical nano-systems can have very specific applications. For example, the authors [44] note that such layered nanostructures can be used to precisely measure the temperature of a device with a spatial resolution of 10 μm in combination with a microsecond temporal resolution.

11. 4D Electron Tomography

Electron tomography began to develop as early as the 1960-ies [45]. Progress in this field was mainly limited by the development of computers, because it is important to process a large number of 2-dimensional electron diffraction patterns and combine them in a single movie. A humorous illustration to these words is given in two images of (Figure 16), taken from the book [45].

The use of modern laser technology has allowed to add to electron tomography the fourth dimension, time, and thus to create a completely unique device (Figure 17).
Figure 12. Time-dependence of the cross-correlation function. The scale in the lower left corner corresponds to 5 microns. Five areas with different initial dynamics are marked in red, yellow, green, blue and purple. The transition from chaotic behavior to coherent oscillations is shown in the panels I-II-III. They correspond to different temporal intervals as indicated in the figure [44].

Figure 13. Images of musical nano-instruments. These frames were obtained with a scanning electron microscope. Note that two of the thinnest cantilevers, p7 and p8 on the left, and two pairs of cantilevers, h1/h2 and h3/h4 on the right, are partially melted in the central part as a result of micro-structuring by the focused ion beam [44].
Figure 14. The oscillations of the cantilevers were caused by the pulsed heating of the sample (top). The results of the fast Fourier transform are presented below. The experimental data obtained in the tilted position of the sample with the angle of 30° are given below (insert in the right). The tilted position made it possible to record the movements of the cantilevers, performed outside of the plane of the nano-harp [44].

Figure 15. The dependence of the oscillation frequency on the size of the respective cantilever. Only the vibrations that occur out of the plane of musical nano-instruments are considered in this case [44].
The temporal behavior of carbon nanotubes was studied by 4D tomography in the work [46]. The sample was twisted in the form of a bracelet. The measured length of the nanotube was \( L \approx 4.4 \) microns. The images in Figures 18 and 19 were obtained using the 200 keV electron pulses. Femtosecond laser excitation led to the heating of the nanotube and caused the subsequent structural changes that began on a picosecond temporal scale.

Of particular interest to a broad audience is that it is possible to create a video, and in the future, a real atomic movie, using the results of 4D electron tomography. It should be emphasized that the pioneering experiment [46] did not induce irreversible changes in the structure of the nanotube, because the total dose received by the carbon sample during the experiment was about 2 orders of magnitude less than the value at which the irreversible deformation occurs. This is a tribute to the extremely high detection efficiency reached in 4D electron microscopy.

12. Electron Microscopy with High Spectral, Spatial and Temporal Resolution

The combination of spatial nano-resolution and subpicosecond, eventually femtosecond temporal resolution, forms the basis of the atomic movie. Naturally, when the energy of the electron beam, transmitted through the sample, is measured in a similar geometry, additional and
Figure 18. (a) 3D images of multi-walled carbon nanotubes at different times. The lengths of the respective segments around a fixed area are denoted as L1 and L2; (b) Cross section of the 3D images. Two-dimensional slices in the plane xy are shown. Slice thickness is 4.6 nm. Dark areas of the arrows point to the nanometer-thick walls. According to these micrographs, the spatial resolution of the method is sufficient for the correct imaging of channels with a diameter of ~10 nm; (c) An image, taken with a transmission electron microscope [46].

Figure 19. Mechanical vibrations of a carbon bracelet. The arrows indicate the direction of the individual sections of a carbon nanotube [46].

very important information is obtained. Note that for most modern research laboratories, such a study would pose quite a serious experimental challenge. Therefore, the precedent in this field plays an important role.

This idea was realized in the work [47] using a single metal nanostructure. In this experiment, the silver particle of triangular shape with a characteristic length of 130 nm and thickness of 20 nm, placed on a substrate of graphene (Figure 20), was irradiated by femtosecond laser pulses with a photon energy of 2.4 eV. The plasma oscillations in silver, excited by the optical radiation, were probed by an ultrashort electron beam with a diameter of 10 nm. The electron beam could be moved over the surface of the sample. Here the energy gain of the electrons, passed through the sample, was measured in addition to the spatial and temporal characteristics.

It is important to emphasize that due to the field localized on the surface of a metal particle, the electrons can not only lose kinetic energy, as would be the case with a standard transmission electron microscope, but also to acquire energy (Figure 21). In principle, this process can be controlled by changing the wavelength of the laser radiation. The accuracy of this procedure is determined by the spectral width of the laser pulse and could reach values of about 1 meV.
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Figure 20. The process of ultrafast spectral imaging. A 10-nm electron beam was scanned over the silver particle that was previously excited by a femtosecond laser pulse. For each position of the probe, the change of the kinetic energy of the electrons (viz. electron spectrum) was measured as a function of the delay between the exciting, optical, and probing, electronic, pulses. The increase in the energy of the electrons is measured in units of the photon energy, being equal to $h\nu = 2.4$ eV [47].

Figure 21. Spectral and temporal characteristics of silver nanoparticles. The electron spectrum, obtained at a delay of 100 fs after the femtosecond optical pulse, is shown below. It is schematically indicated by the position of the electron nanobeam with respect to the triangular silver particle, which is depicted in gray [47].

The results of the measurement of the plasmic texture, generated on the surface of the silver particle, is shown in Figure 22.

It is clearly seen in Figure 22 that the field of plasma oscillations, excited by the laser radiation, is concentrated near the vertices (the cusps) of the triangular particle. There is a fairly good agreement with theory. Qualitatively similar results were obtained near the sharp edge of a copper surface irradiated by the laser [47].

13. Conclusions

Electron microscopy and diffraction with a high temporal resolution have opened the possibility of directly observing processes that occur in non-steady states of the studied substances. A temporal resolution on the order of 100 fs corresponds to the transition of a quantum system through an energy barrier of a potential surface, describing a chemical reaction in the processes of breaking and forming new bonds of the interacting agents. The advances thus opened the possibility of investigating the coherent nuclear dynamics of molecular systems and condensed matter [24,25,48-51].

In the past two decades it has been possible to observe the motions of nuclei in time intervals corresponding to the period of the oscillation of the nuclei. The observed coherent changes in the nuclear system at such time steps determine the fundamental transition from the standard kinetics to the dynamics of the phase trajectory of a single molecule, the molecular quantum state tomography [50-53].
Figure 22. Plasma oscillations, optically excited in the silver nanostructure, were visualized by a 10-nm electron beam. Here, the electron signal is proportional to the height of the corresponding cylinder. For each panel, the energy, acquired as a result of the interaction of the electrons with nanolocalized fields, is shown in the lower right corner [47].

Figure 23. Chronology of the basic works (the first steps) in Dynamic Electron Microscopy, done in the first two decades (1982-2004) [21]. The parallel development of UED, UEM, UEC and DTEM until 2010 is described in [2]. The references [58-109] demonstrate the worldwide experimental activity in this field.
At the present time, the method of ultrafast diffraction continues its rapid development. The methods of ultrafast electron crystallography and electron microscopy with temporal resolution from micro- to subpicoseconds provide great opportunities to the study of the 4D structural dynamics (see the review articles [21,54] and the monograph [2]). Very recent advances in the formation of ultrashort electron pulses allow us to reach an attosecond temporal resolution and observe the coherent dynamics of the electrons in molecules [55-57].

The development of dynamic electron microscopy and the first steps devoted to the progress in the study of structural dynamics of ultrafast processes are illustrated in Figure 23. It is all but certain that exciting developments in this field will continue into the future.

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REFERENCES

[25] S. A. Akhmanov, V. N. Bagratashvili, V. V. Golubkov, A.


[52] A. A. Ischenko, “The Study of Coherent Dynamics of the Nuclei by Electron Diffraction with a Time Resolution. II. Electron Scattering Coherently Excited Molecules,” Pro-


