Nanobiotechnology and its role in the development of new analytical devices†

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Physical methods of molecule observation and manipulation will prove useful, not only as research tools for investigating biomolecular structure and behavior, but also for the creation of nanostructures. Supramolecular and self-assembling structures are able to generate nanostructures, with many such systems being of biological origin. They form the interface between nanotechnology and biotechnology. Whereas biotechnological processes usually involve populations of cells or molecules, nanotechnological methods operate at the level of individual molecule manipulation. This article considers what advances have been made through cross-fertilisation between nanotechnology and biotechnology to develop for the next millennium new analytical tools at the microscale, using nanostructures as the sensitive part and with the ability to detect individual molecules.

The convergence of technologies being developed for machining materials at the atomic scale with technologies achieved for observing nanostructures has promoted the emergence and rapid growth of nanotechnology. Supramolecular and self-assembling structures are able to generate nanostructures, many of such systems being of biological origin. They form the interface between nanotechnology and biotechnology.

Many biological systems—including viruses, membranes and protein complexes—are natural nanostructures. However, they may be analysed in the same way as non-biological structures, and their characteristics can be exploited in nanostructure design and development. It is becoming increasingly obvious that systems that integrate biological and chemical components with physical devices and electrodes have enormous potential for application, not only for developing analytical and monitoring devices, but also for molecular-scale bioreactors. Whereas biotechnological processes usually involve populations of cells or molecules, nanotechnological methods operate at the level of individual molecule manipulation.

This article considers what advances have been made through cross-fertilisation between nanotechnology and biotechnology to develop: studies of single biological macromolecules (i.e., proteins); biotransducers in new analytical devices; and micro-total analysis systems.

A. Studies of single protein molecule

Since many decades ago, characterization of the functional and structural properties of biological molecules has been made by conventional techniques that operate on a population of biomolecules. Catalytic parameters of an enzyme determined by spectrophotometric measurement or the 3D structure of the enzyme obtained from X-ray diffraction are always the average behavior of an enzyme population. The current hypothesis is to assume that a homogeneous population of molecules allows the extrapolation of the results to the single molecule. In the last years, techniques appear to characterize and manipulate individual biomolecules, such techniques are more precisely scanning probes and optical techniques (see the recent reviews†). With these techniques, heterogeneity of populations of molecules has been identified.

Scanning probe microscopy

Over these past years, scanning probe microscopy (SPM) and especially atomic force microscopy (AFM) has been intensively used to study biological samples at the nanometer scale. However, at the beginning, these techniques were not applied easily to biological specimens. Specific problems due to the technique were not known and some image interpretations were erroneous. Today the technique has matured and is reliable, the artifacts are well known and the image interpretation is generally not ambiguous. Concerning topographic imaging, some recent advances in sample preparations, operating conditions, probes and imaging mode have decreased the resolution to the nanometer level. For biological observations, the tip curvature radius (typically 10 nm to be comparable with the protein radius, i.e., albumin has a radius of 4 nm) and the protein compliance lead to great difficulties in observing the substructures of proteins adsorbed onto a solid surface. Nevertheless AFM has generated a growing interest due to the new capabilities it offers. It preserves the native state of molecules or cells because no sample preparations (fixation, dehydration, coatings, etc.) are needed and because proteins, e.g., bovine albumin, RNA polymerase or antibody can be explored in physiological media. Thus dynamic processes like antibody‐antigen complex formation, enzyme activity or the DNA transcription have been observed. Another important aspect of AFM is its capability to measure the force between a sharp tip and the sample surface. This offers the possibility of measuring a wide range of properties from mechanical properties (elasticity, viscosity, plasticity, friction, etc.) to forces (attraction, adhesion, interaction, etc.) using various imaging modes. Nevertheless it is very difficult to measure quantitatively these properties. Generally, the signal of the imaging mode is just used to elaborate an image with an arbitrary unit opening the door to over-interpretation of data.
For some imaging modes it is possible to realize a quantitative image of the tip/surface interaction (i.e., contact stiffness) but very difficult if not impossible to realize a quantitative image of the surface properties (i.e., elastic modulus). The atomic force microscope could be also used as a force-sensing instrument to measure the tip/surface force as a function of the distance and more interestingly the interaction energy (which is the integration of force with respect to distance). These two properties are important because they are responsible for dynamic behaviors like kinetic or affinity constants. Because the atomic force has been built for imaging it has several flaws concerning force-sensing measurement. The most notable of them is that the so-called force curve describes the tipsurface force as a function of the sample position instead of the tip/surface distance. One of the solutions to prevent this phenomena is to achieve an external force on the cantilever (via a magnet for example) to generate a true force tip/sample distance curve. But this has, to our knowledge, not been applied to biological experiment. Nevertheless this kind of experiment has been used to measure the avidin–biotin or antigen–antibody interactions or the elongation force of titin domains. Another interesting evolution of SPM is the development of the near field scanning optical microscope (SNOM). SNOM has been successfully used for biological investigations. Recent developments of probes and of shear force mode has decreased the resolution to a few nanometers. Future work could be the study of the luminescence activity of a single protein.

**Optical techniques**

Regrouping of Rhodamine molecules adsorbed on Si wafers was revealed as fluorescent spots with a measurable number of individual molecules. Fluorescence of the active site of an enzyme, i.e., FAD of cholesterol oxidase, allows one to follow the turnover of a single enzyme molecule. From the results of this kind of experiment, a new view of the population of enzyme molecules with a static and dynamic disorder of reaction rates appears. Individual enzyme molecules have different catalytic turnovers and a molecular memory phenomenon exists in the same enzyme molecule so that enzymatic turnovers are not independent of previous ones.

Recently, the rotation of F1 ATPase during ATP hydrolysis has been observed in an epifluorescence microscope. Visualization of the motion of the F1 subunit was effected by the attachment of a fluorescent actin to the F1 ATPase immobilized on a coverslip. Optical tweezers exploit the fact that light exerts attachment of a fluorescent actin to the F1 ATPase immobilized on a coverslip. Optical tweezers also allow the measurement of the binding force between two biological molecules or movements. Applications concerned the linear motor proteins that move along a polymer track such as kinesin, myosin or RNA polymerase. Understanding of the mechanisms of the linear motor proteins allows the possibility of using these proteins as nanomotors for motions at the nanometer scale.

**B. Biotransducers**

Many technology applications in analysis require the organization of atoms or molecules in a two-dimensional space. Two approaches may be used to create the initial framework. First, as demonstrated by the microelectronic industry, such frameworks can be generated by the manufacture of silicon-based materials, which may then be machined with the desired structure or pattern. This approach will find many applications in labs-on-a-chip fabrication. Second, it is possible to use the self-assembly abilities of synthetic and biological molecules. Self-assembled nano- and microstructures can be generated from natural amphiphilic molecules, and these molecular self-assemblies may be used directly as potential biomimetic systems (for more details see ref. 38). This approach is more related to the fabrication of biochips to combine the sensitivity of a physical detection and the biological specificity. Because of the importance of DNA-based assays for detection of diseases, genome sequencing and environmental control, a DNA chip is one of the major challenges.

**Supported layer**

Self-assembled monolayers (SAMs) may be generated by the spontaneous physisorption or chemisorption of molecules onto a surface. Fusion of phospholipid vesicles on hydrophobic SAM, i.e., octadecyl trichlorosilane (Fig. 1), results in the formation of a phospholipid monolayer on a SAM (Fig. 2A). Peripheral enzyme membranes, such as pyruvate oxidase, and membrane electron carriers, such as plastoquinone and ubiquinone, can be incorporated in the supported bilayer at the physiological level (Fig. 1). When this bilayer was laterally in contact with a built-in gold electrode, lateral diffusion of ubiquinone allows this electron carrier to shuttle between the electrode and pyruvate oxidase (Fig. 1). A catalytic current was measured both the electrochemical reaction of ubiquinone on the gold electrode, the lateral diffusion of ubiquinone along the alkane–lipid bilayer and the catalytic reaction of the ubiquinone with pyruvate oxidase.

![Fig. 1](image1.png) **Fig. 1** Reconstituted biological electron transfer. A lipid monolayer which incorporated a physiological amount of ubiquinone was added to the alkylated monolayer by fusion of lipid vesicles before the incorporation of pyruvate oxidase, a peripheral membrane enzyme. The electron carrier, ubiquinone, diffused along the biomimetic bilayer and made the shuttle between the gold electrode and pyruvate oxidase. The catalytic current measured both the electrochemical reaction of ubiquinone on the gold electrode, the lateral diffusion of ubiquinone along the alkane–lipid bilayer and the catalytic reaction of the ubiquinone with pyruvate oxidase.

![Fig. 2](image2.png) **Fig. 2** Schemes of supported lipid layer. A supported lipid monolayer that allows the incorporation of peripheral membrane proteins is obtained by fusion of lipid vesicles on the top of an alkylated layer (A). Several experimental methods were tested to obtain a lipid bilayer compatible with the incorporation of integral membrane proteins: lipid bilayer tethered by a hydrophilic spacer (B) or by an avidin–biotin complex (C); lipid bilayer onto a polymeric layer (D).
then measured which reflected the electrochemical reaction of the quinone on the gold electrode, the lateral diffusion of the quinone along the bilayer and the catalytic reaction of the ubiquinone with the enzyme.

During the last few years, technological efforts have been made to increase the efficiency of supported layers with comparison to the biological membrane. Formation of a new kind of lipid bilayer spaced from the surface by a linker has been done (Fig. 2). The bilayer was supported on a thin polymer layer 41 (Fig. 2D) or tethered to the surface by a hydrophilic spacer 42 (Fig. 2A), peptide 43 or avidin–biotin complex (Fig. 2C). In the spaced bilayer, an integral membrane protein such as ATPase 44 or rhodopsin 42 has been successfully incorporated. Moreover, the spaced bilayer creates an ionic reservoir between the bilayer and the support. 44 The ionic permeability of the bilayer can be modulated by the incorporation of peptides such as valinomycin 44 and gramicidin 45 that open the way to a new class of bicapture using ion-channel switches. 46 We can guess that the future biochip based on a lipid bilayer will integrate membrane receptors to perform a very specific and very sensible analysis.

2D and 3D imprinting

For some purposes, nanostructures cannot be created by self-assembly alone. In these cases, a technique called ‘templating’ can be used in two possible ways: a template can be used to create a more complex initial pattern for subsequent self-assembly; alternatively, the original structure can be used as a template that can be modified by chemical or physical means to stabilize, or tailor, the properties for a specific purpose (for more details see ref. 38). Molecular imprinting is one example of the templating approach (Fig. 3). A ‘print’ molecule interacts with a solution of molecules (Fig. 3A), which is subsequently ‘fixed’ (Fig. 3B) to form an impression of the print molecule 47 (Fig. 3C). Macroscopic polymer matrices are generally used to create the recognition sites (Fig. 3 upper part). The recognition sites of the polymer are related to a biomimetic approach to create synthetic receptors 48 or synthetic antibodies. 49, 50 In some cases, the imprint has catalytic properties, such as ATP hydrolysis. 51 Imprinting polymers appears then as a tool to create synthetic enzymes 57 in order to increase the stability of the catalyst. The imprinting polymer matrices are applied mainly for affinity chromatography, i.e., for nucleotides, 51, 52 steroids, 53 drugs 54 or herbicide analysis. 55 A thin film of the imprinting polymer can coat the surface of a physical transducer to create substrate-selective sensors. Examples of transducers used in imprinted polymer sensors are various: capacitance, amperometry, conductometry, potentiometry, pH, fluorescence, colorimetric, ellipsometry, surface plasmon resonance, surface acoustic wave and quartz microbalance. 56

Besides macromolecular polymers, 2D imprinting can be done using a SAM to create functionalized monolayers assembled on a surface, i.e., with size and shape specific molecular recognition sites (Fig. 3 lower part). Perforated monolayers that include molecular channels were then tailored and molecular components, i.e., tetradecanethiol and ‘trans’ quinone for photochemical imprinting, 57 were co-assembled in two-dimensional monolayers.

DNA chips

DNA-based assays have an increasing importance for detection of diseases, genome sequencing, forensic and environmental control. They are challenged by the detection of trace amounts of DNA fragments, characterized by a specific sequence in a huge background of other molecules and unspecific DNA. The polymerase chain reaction (PCR) can be used in conjunction with labeled probes to detect very small amount of DNA. The following examples illustrate the purpose in a non-exhaustive way considering the huge research in the field of DNA chips.

The diagnostic method is typically based on hybridization of oligonucleotides probes. Single DNA chips can consist of a matrix of dots coated with various oligonucleotides that simultaneously detect a range of DNA sequences. Deposition of DNA or oligonucleotides on the dot surface can be done by covalent linkage of activated oligonucleotide on amine functionalized surfaces, 56 attachment of thiol functionalized DNA directly on a gold surface, 57 electrochemical polymerization of oligonucleotide pyrrole, 58 intercalation of DNA with a self-assembled monolayer of acridin, 59 specific interaction of biotinylated oligonucleotide with streptavidin immobilized directly on the surface 60 or on a lipid supported layer 61 (Fig. 4).

Sensors are mostly based on radioactive labeled probes, fluorescent probes or electrochemically active probes. Electrochemical probes can be obtained by the linking of an electroactive unit such as ferrocene 62 with the oligonucleotide. The hybridization event can be also detected electrochemically by the intercalation of electroactive molecule such as Orange Acrinid or daunomycin. 63 A peak potential shift results from the hydrophobic interactions (intercalation) between DNA and intercalators. The properties of the intercalators to distinguish between double-stranded and single-stranded DNA are then very important to detect a gene specifically with high sensitivity. In the case of an oligonucleotide-functionalized polypropylene electrode, hybridization decreases the intensity of the current peak due to the oxidation (or doping) of the polypyrrole chain. 58

Fig. 3 Schemes of 2D and 3D molecular imprinting. The different steps for 2D molecular imprinting (lower part) and 3D molecular imprinting (upper part) are: specific interactions with the host molecule (A), polymerization or self-assembly of molecules around the host molecule (B) and removing of the host molecule from the molecular imprint (C). The molecular imprint has both specific interaction sites and a specific shape to recognize the host molecule.

Fig. 4 Schemes of oligonucleotide deposition. The deposition of oligonucleotide on the surface of a DNA chip can be performed in different ways: direct covalent linkage of the oligonucleotide on the surface (A), electrochemical polymerization of oligonucleotide pyrrole (C), immobilization with an avidin–biotin complex directly on the surface (B) or on a lipid supported bilayer (D).
Using fluorescent probes, such as fluorescein labeled oligonucleotides, a simultaneous detection of multiple DNA sequences is possible using a fiber-optic biosensor array system. In this system, several optical fibers were bundled together each with a different probe immobilized on its tip. With the possibility to detect single fluorophore molecules, ultra-sensitive fluorescence techniques allow the detection of individual pairing of oligonucleotides.

Another approach is based on the use of peptide nucleic acids (PNA), a recognition layer in DNA chips. PNA is a structural DNA analogue with a pseudopeptide backbone that mimics DNA by forming complementary duplexes with normal DNA. The PNA/DNA duplexes, that can be done using shorter probes, have higher thermal stability and can be formed at low ionic strength.

C. Micro-total analysis systems

Micro-total analysis systems (μTAS), also called ‘labs-on-a-chip’, must perform the functions of large analytical devices in small units. They must contain elements for the acquisition, pretreatment, separation, post-treatment and detection of the samples. Biochips could be included as a functional element in a μTAS device but a number of other processes are necessary to go from reactants to analysis. Several methods exist for carrying out chemistry confined to well-defined regions of a planar device. For silicon or glass devices, methods include photolithography based on beams of light, electrons or ions; these methods have been typically developed for microelectronics. A network of microfluidic channels can be generated using an elastomeric polymer, e.g., poly(dimethylsiloxane), in which analytes are transported, mixed and separated in μTAS. Patterning of molecules on a reactive surface can be done by microcontact printing of the active molecule, e.g., alkanethiol on a gold surface, with an elastomeric stamp. In this case, patterning of the surface with a hydrophobic and a hydrophilic SAM can also create microfluidic channels. Conventional syringe pumps or microfabricated pumps perform liquid flow in the microfluidic microchannel. Release mechanism of drugs can be for example based on electrochemical dissolution of a thin anode membrane covering microreservoirs filled with chemicals in solid, liquid or gel form.

Strong points of the use of μTAS include the reduction of the consumption of samples and reagents, reduction of the time of analysis, and the increase of sensitivity. These high performances of μTAS will be particularly valuable in the pharmaceutical industry for the screening of combinatorial libraries, in clinical analyses, in DNA-based diagnostics and genotyping. Ramsey’s recent work using microchip electrophoresis as a separation method illustrates the performance. Separations of 15 pM Rhodamine 6G and 30 pM Rhodamine B (200 pL of solution) were performed using microchip electrophoresis and were detected by counting fluorescence bursts from individual molecules. A binary mixture of Rhodamine and dichlorofluorescein were resolved in less than 1 ms using microchip electrophoresis. Polymerase chain reactions were carried out on as many as four DNA samples at a time on a microchip device and the PCR products analyzed on the same device by microchip electrophoresis.

D. Concluding remarks

The first major advances in nanotechnology were done by the microelectronics industry with the manufacture and machining of silicon materials. The second breakthrough was the emergence of the discipline of nanobiotechnology from attempts to understand friction, wear and lubrication at the atomic scale, and the atomic-force and friction-force microscopes were developed as investigative tools for this purpose. The third step in nanotechnology is the convergence between the field and the recent achievements in biotechnology.

There is not only a fertilisation of nanotechnologies through the use of biotechnologies but also a huge reciprocal impact of the nanococepts on biology.

The acquisition of expertise in these areas will soon become a prerequisite for those wishing to participate in the next phase of biotechnological evolution. Nanobiotechnology is important, not only for its research and development tools, but also for its potential commercial significance in the development of new analytical devices and preparative bioreactors.

References
