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COVER ARTICLE Bisquert Physical electrochemistry of nanostructured devices **PERSPECTIVE** Marangos *et al.* Dynamic imaging of molecules using high order harmonic generation



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Physical electrochemistry of nanostructured devices

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This Perspective reviews recent developments in experimental techniques and conceptual methods applied to the electrochemical properties of metal-oxide semiconductor nanostructures and organic conductors, such as those used in dye-sensitized solar cells, high-energy batteries, sensors, and electrochemical and optoelectrical measurements for semiconductor nanostructures (sintered colloidal particles, nanorods, arrays of quantum dots, *etc.*) deposited or grown on a conducting substrate. The Fermi level displacement by potentiostatic control causes a broad change of physical properties such as the hopping conductivity, that can be investigated over a very large variation of electron density. In contrast to traditional electrochemistry, we emphasize that in nanostructured devices we must deal with systems that depart heavily from the ideal, Maxwell–Boltzmann statistics, due to broad distributions of states (energy disorder) and interactions of charge carriers, therefore the electrochemical analysis must be aided by thermodynamics and statistical mechanics. We discuss in detail the most characteristic densities of states, the chemical capacitance, and the transport properties, specially the chemical diffusion coefficient, mobility, and generalized Einstein relation.



in mathematical physics. He joined Universitat Jaume I in 1991 and now is full professor of applied physics. He leads a research group of 8 physicists and chemists working on photovoltaic and electrooptic materials and devices based on nanostructured metal oxides and organic conductors. Current research activity is focused on dye-sensitized

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solarcells, organic LEDs, electrochromic displays, batteries and supercapacitors. He is specialist in theoretical modeling and interpretation of impedance spectroscopy of electroactive films, transport in disordered materials, interfacial charge-transfer, and the glass transition.

1. Introduction

The increasing need for the replacement of conventional energy sources in favor of carbon-neutral energies has boosted the research on new materials and devices for the production and storage of clean energies. Solar cells are able to directly convert sunlight into electricity. A great effort of research is currently devoted to develop solar cells that could lower considerably the production costs of current photovoltaic devices (mostly silicon-based) while maintaining a reasonable

Departament de Física, Universitat Jaume I, 12071 Castelló, Spain. E-mail: bisquert@fca.uji.es efficiency and stability. Also important would be devices able to produce hydrogen by splitting water with sunlight. But even if the production of huge amounts of clean energy, either in the form of electricity or hydrogen, would be reached, it is still necessary to store and transport the energy for solving the demands of energy use in society. Convenient hydrogenstorage systems currently constitute a bottleneck for the realization of the hydrogen economy. Cheaper fuel cells, and devices for energy storage with both a high energy and power density, like batteries and supercapacitors, are needed for electric cars, and also for leveling off the intermittent nature of several renewable energy sources. Other advances, such as large area light-emitting diodes (LEDs) for solid-state lighting, would also contribute significantly to the efficient use of energy.

The early 1990s witnessed a great development of a series of electrochemical devices.¹ The lithium ion battery entered the market, to soon become an essential component of portable electronics, and the group of M. Grätzel developed a successful dye-sensitized solar cell (DSC) based on nanocrystalline TiO₂.^{2,3} These breakthroughs showed the great potential of devices based on electroactive materials for energy production and storage applications. In addition, the demonstration of efficient LEDs with organic semiconductors⁴ (that have become a technological reality) showed that electroactive and optically active films for devices could be formed from either inorganic or organic materials, or with a combination of both. Research on nanostructured electrochemical devices has exploded in recent years, in relation with DSC, high-energy batteries, photocatalysis, chemical and biochemical sensors, and electrochromic devices.

This paper will focus on combined methods of physics and electrochemistry for analyzing the properties of electrochemical nanostructured devices in which a semiconductor nanostructure can be electronically addressed from a conducting substrate. Since organic, hole transport media are usually a key element of such devices, their properties will be addressed as well. The paper presents an introduction to the subject exposing an array of experimental methods and theoretical concepts that have emerged in recent years while studying such devices, particularly around the subjects of Fermi level displacement and electron transport. Recent reviews on physical properties of DSC, ^{5,6} electrochemical gating,⁷ and the electrochemical determination of the density of states (DOS).⁸ describe these related issues in more detail, and in another paper⁹ I discuss a broad range of applications of metal-oxide nanoparticles in electrochemical devices. Here, emphasis is made to relating the macroscopic quantities measured by physical-electrochemical methods to the electronic and ionic properties of nanostructured devices and in particular of DSC. First some introductory remarks will be made about nanoparticles, nanostructures and their application in electrochemical, photoelectrochemical and all-solid devices. Next, in section 2 the electrochemical measurement of the DOS and its shape in a variety of relevant materials is described. Section 3 considers the basic concepts of macroscopic transport in a semiconductor nanostructure, with several particular models that illustrate the general concepts. We finish with the conclusions and outlook in section 4.

1.1 Nanoparticles and nanostructures

There is a wide variety of nanostructures currently investigated for electronic and electrooptic devices with different morphology, size and composition. We will be mostly concerned with physical electrochemical properties of semiconductor nanoparticles and nanostructures that are prepared by wet chemical methods. These methods lead to colloidal nanocrystals present in dispersion, which are the basic building blocks for the preparation of larger architectures.¹⁰

Quantum dots are nanocrystals of size roughly between 1 and 10 nm. In these crystals, for example of CdSe, the electron wave functions are strongly confined, so that the electron energy levels are discrete and their separation is determined by the crystal size. Accordingly, electronic and electrooptic properties of quantum dots can be tailored by the dimensions.

Colloidal nanocrystals, in the range 10–50 nm, do not generally show the acute confinement effect, but are important for many applications, because they allow for electrochemical gating, *i.e.*, an increase in electronic density compensated by ionic density at the surface. Examples are metal oxides used in DSC such as TiO₂, ZnO and SnO₂.^{11–15}

Colloidal nanocrystals are usually deposited over a conducting substrate and thermally treated to form a connected array of nanoparticles that can be used as electroactive electrodes. Structures can be more or less ordered, and interparticle connection can also be controlled with molecular ligands, which can be used to promote the self-assembly of special architectures.¹⁶

Another approach towards electroactive nanostructured electrodes consists on growing nanofibers or nanowires with columnar, tubular, dendritic or other structures over a sub-



Fig. 1 TiO₂ nanotubes prepared by electrochemical anodization of Ti foil in dimethyl sulfoxide (top) and ethylene glycol (bottom) solution. This type of nanotube,^{17,18} when dye-sensitized, produces efficient DSC reaching 6.8%.¹⁹ Images courtesy of C. G. Grimes.

strate, see Fig. 1 and 2. These structures have the advantage of providing long and uninterrupted paths for electron transport while maintaining a high area density as that found in random nanoparticulate films.^{20–23} There has been recent progress in obtaining highly ordered transparent TiO₂ nanotube arrays of the type shown in Fig. 1 for DSC with high electron lifetimes and excellent pathways for electron percolation.^{17,18} Vertically aligned ZnO nanorods, formed by electrodeposition on a transparent conducting oxide,^{24,25} as those shown in Fig. 2, have also attracted much attention for applications in solar cells^{20,26–28} and optoelectronic devices such as light emitting and laser diodes.^{24,29,30}

An important step of nanostructuring is an additional treatment of colloidal nanoparticles or nanorods for governing the electronic properties at the surface. This can be realized, for example, with strategies of conformal coating of sintered nanoparticulate films. Insulating layers *ca.* 2 nm thick have been deposited over TiO_2 nanoparticles in DSC.^{31–37} It



Fig. 2 SEM microgaph of electrodeposited ZnO nanowire array. Image courtesy of R. Tena-Zaera.



Fig. 3 Conductivity plot of a bare, mesoporous TiO_2 film and molecular modified films with electrochemical deposited 4-methoxybenzenediazonium tetrafluoroborate (oab) and 4-cyanobenzenediazonium tetrafluoroborate (cab). Reprinted with permission from ref. 39, copyright 2005 American Chemical Society.

was shown that alumina coating improves considerably the electron lifetime by an almost complete passivation of surface trap states.^{23,38} Absorption of molecules with different dipole moment modifies the energy level of a nanostructured semiconductor immersed in solution, as shown in Fig. 3 and discussed later.³⁹ Besides shifting the conduction band of the semiconductor, adsorbed molecules also are able to reduce recombination and increase the efficiency of DSC.^{40,41}

Finally some architectures require the combination of different elements with specific photophysical, physical or electrochemical functions. For example nanostructured solar cells are envisioned with a matrix of TiO_2 nanoparticulate film sensitized with surface-attached quantum dots⁴² in order to exploit the opportunities arising from discretization of energy levels in quantum dots, which could greatly enhance the photocurrent by multiple exciton generation from a single high energy photon.^{43–46} Devices such as photocapacitor⁴⁷ and self-driven photoelectrochromic window⁴⁸ have been formed by combining in direct contact several nanoparticulate layers with different functions.

1.2 Basic features of nanostructured electrochemical devices

Prompted by key applications such as the Li battery and DSC, the widespread investigation of electroactive materials started with electrochemical configurations. The basic structure of such devices is shown in Fig. 4. It is composed of a nanoparticulate electroactive electrode, a counterelectrode and an ionic (liquid or solid electrolyte) or hole conductor that joins both films. The first electrode is "active" in the sense that it realizes the main steps in the function in the device, such as sunlight conversion to electricity, or energy storage. Usually in realizing such function the material undergoes a huge change in the concentration of some species, either electrons or ions, or both, with the consequent change of the electrochemical potential.

The process of electrochemical charging is shown in more detail for the DSC in Fig. 5. In this system the change in electron density in the TiO_2 nanoparticles by photoinjection from surface-adsorbed dye molecules is facilitated by positive but inert ions at the nanoparticles surface, in order to maintain electrical charge neutrality. Electron density may vary from nearly zero, in the insulator state, to 100 electrons per 10 nm



Fig. 4 Scheme of an electroactive device formed by a nanostructured film and counterelectrode joined by an electrolyte (or hole transport material).

diameter nanoparticle. Here a main point of contrast with solid-state semiconductor devices must be emphasized. In semiconductor devices many device properties are ruled by electrical fields in space–charge regions at the interfaces. In general, whether the internal field can be built or not into an object of nanometric dimensions, depends on the relationship of the object size to the Debye screening length. In the case of the DSC indicated in Fig. 5, long range electrical fields do not govern the properties of the device. In fact if the particle is low-doped and not too large there is no room to build internal electrical fields in the nanoparticles⁴⁹ and the conduction band remains homogeneous in the major part of the nanostructure (the opposite situation will be treated in section 2.4).



Fig. 5 Schematics of a dye-sensitized solar cell, consisting on dye molecules adsorbed on nanoparticulate TiO₂ that is deposited over a transparent conducting oxide (TCO). Photoinjection increases the chemical potential (concentration) of electrons in the TiO₂ phase (A). The electron Fermi level, $E_{\rm Fn}$, is displaced with respect to the lower edge of the conduction band, $E_{\rm c}$. The electrode potential, V, is given by the difference between $E_{\rm Fn}$ and the redox level $E_{\rm redox}$. The increasing negative charge in the semiconductor nanoparticles is compensated by positive ionic charge at the surface (B). With the change of $E_{\rm Fn}$ also changes the electrostatic potential of the Helmholtz layer and semiconductor bandbending at the interface between the exposed surface of the transparent conducting oxide substrate and the electrolyte (C).



Fig. 6 Basic energy diagram for the systems of Fig. 4 and 5, showing the applied potential V, the electrochemical potentials η , the chemical potential μ of electrons in phase A, and the energy level E_A in the phase A. (a) Equilibrium situation without applied potential. (b) Equilibrium situation under an applied potential.

When the TiO₂ electrode in Fig. 4 contains a significant concentration of electrons, it is in an out-of-equilibrium situation and is able to do work as it returns to equilibrium. In fact the device is arranged in such a way that the only path for electrons is the external circuit (except for recombination due to interfacial charge transfer), because the electrolyte only conducts ions in batteries, and holes in solar cells. The amount of work that this system can do is described by the free energy per carrier, which is the electrochemical potential (η), also known as the quasi-Fermi level ($E_{\rm F}$), in the case of electrons in a solid, and as the redox potential ($E_{\rm redox}$), in the case of ionic species in solution. These different denominations refer to the same thermodynamic quantity.⁵⁰

Considering the energy diagram in Fig. 6, we assume that the species injected in phase A with number density n_A , is distributed in equivalent sites with total density N_A , hence the fractional occupancy of sites is $x_A = n_A/N_A$. The electrochemical potential, $\eta_A = \eta_A(x_A)$ has two basic components

$$\eta_{\rm A} = E_{\rm A} + \mu_{\rm A}.\tag{1}$$

The first one, E_A , is the energy level, it is the energy gained when the system acquires one carrier. For noninteracting species, the energy level E_A is independent of the concentration, and $\eta_A(x_A)$ is equivalent to the Langmuir isotherm. The energy level for electrons can be associated with ϕ_A , the Galvani (electrostatic) potential with respect to some suitable reference level, as follows

$$E_{\rm A} = -q\phi_{\rm A} \tag{2}$$

where q is the positive elementary charge.

The second component in eqn (1), μ_A , the chemical potential, is an entropic contribution that accounts for the dispersion of the carriers over all the available sites. If the species is distributed randomly in the available sites the following expression holds:

$$\mu_{\rm A} = k_{\rm B} T \ln \frac{x_{\rm A}}{1 - x_{\rm A}} \tag{3}$$

where $k_{\rm B}$ is Boltzmann's constant and T the absolute temperature.

If the semiconductor nanocrystallites are wide enough to neglect confinement effects, they will have a transport band similar to that for carriers in macroscopic materials. Let us assume that electrons have been injected to the conduction band with density n_c . The lower edge of the conduction band E_c is related to the potential in the semiconductor phase with respect to the electron at rest in vacuum, as $E_c = -q\phi$.⁵¹ N_c is the effective density of conduction band states. Usually the Fermi level remains below the conduction band level $E_c - E_{Fn} \gg k_B T$, hence $n_c \ll N_c$ and the electrochemical potential or Fermi level, can be expressed as

$$\eta_{\rm n} \equiv E_{\rm Fn} = E_{\rm c} + \mu_{\rm n} = E_{\rm c} + k_{\rm B} T \ln(n_{\rm c}/N_{\rm c}).$$
 (4)

The cell potential in Fig. 5 corresponds to the difference of electrochemical potential of electrons between the contacts,⁵² as indicated in the scheme of Fig. 6

$$-qV = \eta_{\rm A} - \eta_{\rm C}.\tag{5}$$

In equilibrium the electrochemical potential is homogeneous through the device, $\eta_A = \eta_C$ and V = 0 as shown in Fig. 6(a). The quasi-equilibrium situation under a nonzero potential is represented in Fig. 6(b). Note that the chemical potential in A $(\mu_A = \eta_A - E_A)$, and hence the concentration of electrons, increases with respect to the situation V = 0, while the electrochemical potential in the electrolyte, η_c , remains stationary.

The distinction between the chemical and electrostatic potential of species in a given phase, stated in eqn (1), is entirely conceptual. A priori the contributions cannot be separated experimentally without making non-thermodynamic assumptions.⁵³ Furthermore, this distinction is lost for carriers strongly interacting with their neighbors, as in examples discussed further on. Nonetheless, the statistics of independent particles is broadly used and applies in a great variety of electrochemical systems (e.g., low-doped nanostructured semiconductors and electronically conducting polymers). When the DOS at the Fermi level is low, the potential has the effect of displacing the Fermi level η_A with respect to the energy level $E_{\rm A}$. In these situations there is a significant change in the concentration of carriers that can be interpreted as a change in the chemical potential (see for example ref. 54). Additionally changes may occur in the local electrostatic level associated with self-charging of the phase or interaction of the carriers.⁵⁵ In contrast, for metals and highly doped semiconductors, the DOS at the Fermi level is very high. Significant changes in the density of carriers, as an effect of an applied potential, are not possible, because the Fermi level is pinned at a fixed energy level.

When the Fermi level of electrons, determining the cell potential V, changes, not only the chemical potential is modified, but variation of *electrostatic* potential between the electrodes occurs as well *inside* the cell. However it may be confined by shielding to a short region at the substrate surface, as indicated by the step of the conduction band level in Fig. 5.⁴⁹ Therefore in many situations the diagrams of Fig. 6 constitute a good approximation to describe the device.



Fig. 7 Scheme of an electrochemical cell for measuring a nanoparticulate film as working electrode (WE), indicating the reference electrode (RE) and counterelectrode (CE). (a) Normal electrochemical cell. (b) The RE is formed of the same material as the WE by introducing an insulating gap in the conducting substrate.

For electrochemical measurements, the potential of the active electrode (the working electrode WE in the electrochemical cell, Fig. 7(a)) is recorded with respect to a reference electrode RE, while the current flows between working and counterelectrode CE.⁵⁶ In devices where WE and CE need to be close to each other special precautions are needed for using a RE. For example in DSC an internal photo-reference electrode is proposed with the cell structured as indicated in Fig. 7(b).⁵⁷

Instead of photoinjecting the electrons in a DSC, as indicated in Fig. 5, the change of electron Fermi level can be induced by applied voltage using a potentiostat in the configuration of Fig. 7(a). It is important to recognize that the manipulation of the Fermi level induces a huge change in an array of physical properties related to the electron density in the nanostructure. Therefore, the electrochemical configuration constitutes an excellent tool for studying the physical properties of nanoparticles and nanoparticulate films. One can monitor properties such as the electronic conductivity over many orders of magnitude in a very simple way.58,59 In addition, interesting devices are formed, based on the manipulation of electronic density in the nanostructure by potential control. For instance molecules attached at the metal-oxide surface as in Fig. 5 can be oxidized and reduced, changing the optical transmission of the film as a whole, which yields an electrochromic device.⁶⁰ It is even possible to electronically address from the substrate the molecular layer absorbed in the surface of the mesoporous structure.^{61,62}

Since the changes in electrode potential modify the properties of the electrode, very often we are interested in characterizing the nanostructured electrode as a function of the steadystate Fermi level or electrochemical potential. To this end one uses a measurement based on a general principle that involves two levels of perturbation. A large one establishes a steady state in the semiconductor, for instance by a certain level of photon irradiation or steady-state bias potential, while a small perturbation probes the properties of that particular steady state, usually by time transient or frequently resolved means. The small perturbation produces differential quantities such as a resistance or capacitance. Studies of macroscopic assemblies of semiconductor nanoparticles often utilize such techniques, including electrochemical impedance spectroscopy (EIS),⁶²⁻⁶⁷ intensity-modulated photocurrent spectroscopy (IMPS).68-72 intensity-modulated and photovoltage spectroscopy (IMVS).⁷³ Recently, many studies have determined the properties of DSC using EIS.^{40,64,66,74–78} This technique has the advantage of identifying different types of operational elements, consisting either in electronic or ionic processes, in photoelectrochemical devices.



Fig. 8 (a) Electrochemical transistor measurement configuration. The conducting substrate over which the film is deposited is divided into two regions, separated by an insulating gap. The separated regions serve as two working electrodes WE₁ and WE₂. (b) Equivalent circuit of the electrochemical transistor configuration. When WE₁ and WE₂ are shorted the film can be operated as in normal electrochemical cell with potential U_{bias} . The two working electrodes can also be operated independently with potentials U_1 and U_2 with respect to the reference electrode (RE).

An important measurement of this class is the determination of electronic conductivity as a function of electrode potential and it uses a conducting substrate that is divided in two separated regions, bridged by the nanostructured film, Fig. 8(a).7,58,79 These regions can be controlled as independent working electrodes. Then it is possible to govern the Fermi level in the film with a bias $U_1 \cong U_2$ as in Fig. 8(b), while maintaining a small potential difference between the two sides, $\Delta U = U_2 - U_1$, which causes a current flow ΔI between WE₁ and WE₂ that enables to measure the electronic conductivity, σ , related to $\Delta I / \Delta U$ by the geometry of the film.^{58,59,80,81} An example of the application of this technique is shown in Fig. 3 for measurements of the conductivity of nanostructured TiO₂ with different surface treatments. Changes in the conductivity are obtained by a shift of the conduction band of TiO₂ due to the different dipole moments of the molecular absorbed species.39

Other electrochemical techniques scan the properties of the nanostructured electrode using a single large perturbation. For example in cyclic voltammetry (CV) the current is recorded while the electrode potential is varied at a constant rate. CV has the advantage of providing a global view of the properties of the electrode in a swift measurement.⁸² When a thin film is voltage-scanned close to equilibrium, CV provides the DOS of the film material, as discussed later.^{82,83}

In electrochemical devices, the counterelectrode may require special properties, for example for storing a large amount of ions expelled from the active film. High-energy density of the Li battery necessitates, in addition to high specific capacity, that the difference in equilibrium potentials between positive and negative electrodes be large, so that both anode and cathode must be optimized. For anode, Li metal can be replaced by some appropriate hosts capable of Li-ion insertion at a potential not far from the equilibrium potential of Li electrode, *e.g.* graphite and some other carbonaceous materials. Therefore two active films may be coupled in a device, and electrochemical potentials will change in both.

Liquid electrolytes have the benefit of realizing a perfect junction whatever the morphology of the active film, so that a huge active area, addressable from the macroscopic contacts, is readily obtained. In addition, liquid electrolytes provide a very large ionic (or hole) conductivity and a wide potential window of operation. But liquids are not generally convenient for all devices, due to technical problems of sealing and volatility. Consequently, a major trend of current research in DSC aims at solid or quasi-solid electrolytes (such as conducting polymers and room-temperature ionic liquids) with good ionic or hole conducting properties for replacing the liquid electrolytes.^{84,85} Conversion efficiencies exceeding 8% have been achieved with non-volatile electrolytes in combination with specially designed dyes, standing 1000 h accelerated test at 80 °C without decay of performance.⁸⁶

It is appreciated that devices, which can be solar cells, supercapacitors, batteries, and others, formed with nanostructured semiconductor films filled with a convenient "soft" conducting medium (able to fill the solid nanostructure), rely on the separation of Fermi levels or electrochemical potentials in at least two phases that are separately connected to the outer electrodes, as indicated in Fig. 6. The voltage generated by the device is determined by the ability of the different phases to maintain the excess carriers. In batteries, the electrolyte must be a perfect electronic insulator for maintaining the stored energy over extended periods of time. In solar cells an electronic connection exists necessarily between the two phases, otherwise the process of photogeneration would be impossible.³ So the kinetics of the reciprocal process (recombination) must be very slow. The surface of nanoparticles and quantum dots is therefore a critical aspect for most applications, it is where carriers in separate phases meet each other and eventually recombine. If the electrons are lost across the particle's surface the voltage in the solar cell device will be very low. Furthermore, often at the surface bandgap electronic states (surface states) are formed that modify the expected physical properties of quantum dots.

During operation, charge carriers in each phase must be well connected to the respective electrode, and therefore efficient electron or hole transport in each medium is a necessary requirement. Indeed, the work per carrier described by the electrochemical potentials as commented above, is restricted to a near equilibrium situation, *i.e.* when the current is close to zero. Devices such as solar cells and supercapacitors need to supply a significant power, and this is determined both by the voltage and current that the device generates. A solar cell with a very high open-circuit voltage may be a useless device if it cannot supply the electrons due to a low conductivity in the semiconductor nanostructure.

Summarizing, in the nanostructured electrochemical devices the classical, metal/solution interface of electrochemistry has been largely extended in several ways. Microporous or nanoporous films up to 10 μ m and larger are used on top of the metal contact. Besides the interfacial processes at the contact of the nanoparticles and the solution, a variety of electronic and ionic processes occur in the material itself. It is necessary to use an array of chemical, electrochemical and physical methods and techniques for understanding the behavior of such materials and improving the performance of devices.

2. The density of states

One important question for the analysis of electronic properties of a given nanostructured film is the relationship between the average electron density and the electrochemical potential. Theoretically this is a problem amenable to the methods of thermodynamics and statistical mechanics. If the electrons can be treated as non interacting entities, the question is related to finding the one-particle DOS. When there are interactions between the electrons and with surrounding media, for example through polarization effects, the question becomes a manybody problem that is much more complex from a theoretical point of view. Examples of systems with interacting carriers will be presented in section 3.

2.1 The electrochemical capacitance

Let us consider first the case in which the energy of a state does not depend on electrochemical potential, *i.e.* we neglect many particle effects. One of the methods for determination of DOS is based on the measurement of the variation of the density of electrons (holes) caused by a variation of Fermi energy.⁸ As an example in Fig. 9 a possible DOS for the nanostructured TiO₂ electrode discussed before in Fig. 5 is indicated. As already commented on before, as a result of ionic charge compensation the electronic density can be varied as a univocal function of a homogeneous Fermi level $E_{\rm Fn}$.

We consider one specific electronic state characterized by the energy E. This energy is defined to be increasingly negative for states deeper in the gap. The average occupancy is described by the Fermi–Dirac distribution function

$$f(E - E_{\rm Fn}) = \frac{1}{1 + \exp[(E - E_{\rm Fn})/k_{\rm B}T]}.$$
 (6)



Fig. 9 Electron energy diagram illustrating the behavior of a nanostructured TiO₂ electrode (shown in the top scheme) when a variation dV of the electrochemical potential of electrons η_n (Fermi level) is applied, assuming that conduction band energy (E_c) remains stationary with respect to the redox level, E_{redox} . Changes of occupancy both of conduction band, dn_c , and trapped electrons in localized levels, dn_L (shaded region of the bandgap), are indicated.



Fig. 10 Schematic representation of (a) electrostatic capacitor and (b) chemical capacitor.

A displacement of the Fermi level causes a variation of the state occupancy in the following way

$$\frac{\mathrm{d}f}{\mathrm{d}E_{\mathrm{Fn}}} = \frac{1}{k_{\mathrm{B}}T}f(1-f). \tag{7}$$

Let us introduce the concept of the electrochemical capacitance. We consider in Fig. 10 two basic mechanisms of accumulating charge with respect to voltage in electrochemical systems. The first one, Fig. 10(a), is a standard dielectric capacitor where energy is stored in the electrical field, related to spatial charge separation. This capacitance is ubiquitous at interfaces with space charge such as Schottky barriers and Helmholtz layers. In general the dielectric capacitance is associated with charging the spatially separated plates of the capacitor with a displacement current. The second one, Fig. 10(b), consists of increasing the chemical potential of the species in a bulk material by increasing their number. Here the "plate" of the capacitor is charged by conduction current. This second case can be called the chemical capacitor^{85,87} and it is specially relevant for intercalation batteries and solar cells.85

As commented before, it may not always be possible to separate electrostatic and chemical potential contributions in the electrochemical potential. In the general case one defines the electrochemical capacitance, that relates the change of concentration to the change of electrochemical potential

$$C_{\eta} = Nq^2 \frac{\mathrm{d}f}{\mathrm{d}E_{\mathrm{Fn}}} \tag{8}$$

where the capacitance is given per unit volume using the volume density of the state, N.

In the case of Fig. 9, it has already been discussed that the voltage variation is absorbed at the TCO/TiO₂ electrolyte interface, hence the relationship $-qdV = dE_{Fn}$. The Fermi level inside the TiO₂ nanostructure is displaced towards the conduction band, *i.e.*, the change of electrochemical potential implies a change of the chemical potential of electrons: $dE_{Fn} = d\mu_n$. Eqn (8) gives a purely chemical capacitance

$$C_{\mu} = \frac{Nq^2}{k_{\rm B}T} f(1-f).$$
(9)

For a single state the capacitance peaks at the energy level of that state, see below (Fig. 17(b)).

In Fig. 9 the nanostructured electrode is treated as a continuum in which the DOS, constitutes a probability of

finding a number of electronic states g(E)dE in the energy interval dE. An exponential distribution of localized states in the bandgap is described by the expression

$$g(E) = \frac{N_{\rm L}}{k_{\rm B}T_0} \exp[(E - E_{\rm c})/k_{\rm B}T_0].$$
 (10)

Here $N_{\rm L}$ is the total density and T_0 is a parameter with temperature units that determines the depth of the distribution. Also a band of transport states at the energy level $E_{\rm c}$ is indicated in Fig. 9.

When we consider the distribution of bandgap states g(E), the chemical capacitance is obtained integrating all the contributions through the bandgap

$$C_{\mu} = q^2 \int_{-\infty}^{+\infty} g(E) \frac{\mathrm{d}f}{\mathrm{d}E_{\mathrm{Fn}}} \mathrm{d}E. \tag{11}$$

Using $df(E - E_{Fn})/dE_{Fn} = -df(E - E_{Fn})/dE$ and integrating eqn (11) by parts, we arrive at

$$C_{\mu} = q^2 \int_{-\infty}^{+\infty} \frac{\mathrm{d}g}{\mathrm{d}E} f(E - E_{\mathrm{Fn}})/\mathrm{d}E.$$
(12)

A simple solution to eqn (12) is obtained by the zero-temperature limit of the Fermi function, *i.e.* a step function at $E = E_{\text{Fn}}$ separating occupied from unoccupied states. Then it follows that

$$C_{\mu} = q^2 \int_{-\infty}^{E_{\rm Fn}} \frac{\mathrm{d}g}{\mathrm{d}E} \mathrm{d}E = q^2 g(E_{\rm Fn}). \tag{13}$$

In this approximation, eqn (13), the charging related to the perturbation dV corresponds to filling a slice of traps at the Fermi level. Note that in eqn (13) the thermal spread of the distribution function is completely neglected. Near room temperature this approximation is justified to some extent if the DOS is smoothly varying as a function of the energy. However, detailed features of the DOS are rounded off, so for a more exact determination the thermal effect of the distribution should be taken into account. An iterative method to derive the DOS from charging data has been developed.^{88,89}

On the other hand, for the transport states at the energy level E_c , and for potentials such that $E_{Fn} \ll E_c$ ($f \ll 1$), the chemical capacitance of the conduction band (cb) states is given by the tail of the Boltzmann distribution indicated in eqn (4)

$$C_{\mu}^{\rm cb} = \frac{N_{\rm c}q^2}{k_{\rm B}T}f = \frac{N_{\rm c}q^2}{k_{\rm B}T}\exp[-(E - E_{\rm Fn})/k_{\rm B}T].$$
 (14)

Let us mention that the total capacitance can take negative values in nanostructured devices, provided that the system is far from equilibrium by applied bias.^{67,90}

2.2 Factors affecting the chemical capacitance

According to eqn (13) in a first approximation the chemical capacitance is simply proportional to the DOS. This method has been applied in many works for investigation of DOS in conducting polymers,^{54,91} quantum dots,⁵⁹ and porous semi-conductors.^{13,82,92–94} The electrochemical determination of the

DOS in metal-oxide nanostructures and organic conductors has been reviewed recently,⁸ and are only briefly commented on here.

For the case of nanostructured TiO_2 the capacitancevoltage relationship, according to the traps distribution in eqn (10), is exponential

$$C_{\mu}^{\text{traps}} = \frac{N_{\rm L}q^2}{k_{\rm B}T_0} \exp[(E - E_{\rm c})/k_{\rm B}T_0]$$
(15)

with a slope $1/k_{\rm B}T_0$ in log–linear representation, a fact that has been observed many times in the literature, using EIS, $\rm CV^{13,40,66,82}$ and electron extraction as a function of opencircuit potential.^{92,95,96}

In organic conductors, disorder or structural correlations with correlation lengths of a few intermolecular distances (fluctuation in the local conjugation length) lead to a dispersion of energies. It is widely agreed that the distribution of electronic states in disordered organic conductors has a Gaussian shape, and this is observed by capacitance measurements.^{83,91} When a conducting polymer is oxidized electrochemically, the excess carriers (holes) in the polymer chains usually lead to a molecular deformation, which causes a lowering in energy for the excess carrier. Such a carrier together with its produced molecular deformation is called a polaron (P). If two charge carriers share the same molecular deformation a bipolaron (B) is formed; the energy gained by forming only one deformation may outweigh the increased Coulomb repulsion energy.⁹⁷ It has been pointed out that the relative stability of one B versus two P may depend on the experimental conditions such as solvent and type of counterions.98

The statistics of P and B in conjugated polymers are well known,^{99,100} and from these expressions the chemical capacitance can be calculated. In addition, the effect of the Gaussian disorder on the statistics of carriers has been described.⁵⁴ Fig. 11 shows the predictions of the model⁵⁴ for the chemical capacitance with the successive formation of P and B, in the presence of Gaussian disorder. Initially, at low oxidation potentials, only P are formed, but since the B energy is lower than that of two P (in this example), in the simulation the P readily recombine to form B. Therefore, the P peak in the chemical capacitance, Fig. 11(b), is quite small, and the CV is dominated by the B peak. Experimental examples are found in ref. 8. Note also the negative capacitance of polarons. This is due to the decrease in their number, when the B formation is dominant, despite the increase in the thermodynamic driving force.

In ion-intercalation materials the cell voltage variation with the amount of inserted ions (fractional occupancy of lattice sites x) is an essential characteristic for applications such as Li ion batteries. For example modification of composition in metal oxides Li_x MO determines the chemical potential variation, $\mu(x)$, with respect to the Li/Li^+ potential (μ_{ref}). The observed capacitance is dominated by chemical capacitance, eqn (8), which can be written, with respect to N, the number density of intercalation sites, as

$$C_{\mu} = q^2 N \frac{\partial x}{\partial \mu}.$$
 (16)

Obviously, the chemical capacitance can be determined as the inverse derivative of the equilibrium voltage-composition



Fig. 11 Simulation of the charging experiment for a polymer with site density $N_{\rm s} = 10^{20} \,{\rm cm}^{-3}$. The polaron levels are centered at $\varepsilon_{\rm P} = 1.5 \,{\rm eV}$ with dispersion $\sigma_{\rm P} = 0.2 \,{\rm eV}$, and the bipolaron levels are centered at $\varepsilon_{\rm B} = 2.8 \,{\rm eV}$ with dispersion $\varepsilon_{\rm B} = 0.15 \,{\rm eV}$. (a) Number density of polarons and bipolarons. (b) Chemical capacitance of polarons, $C_{\mu}^{\rm P}$, bipolarons, $C_{\mu}^{\rm B}$, and total capacitance, as a function of Fermi level.

curve. This derivative parameter is widely used in insertion studies because it provides a detailed view of the features of the thermodynamic function. Many studies of *crystalline* intercalation systems used the mean-field (Bragg–Williams¹⁰¹) approximation, corresponding to the Frumkin isotherm,¹⁰² to describe the voltage–composition and equilibrium capacitance curves.^{102–106} We should also remark that in diffusion control the CV departs from the capacitive behavior. Numerical methods are required for explaining detailed features of voltammetry in the presence of the phase transitions that are characteristic of Li intercalation.¹⁰⁷

In contrast, in *amorphous* intercalation materials, in particular in $\text{Li}_x WO_3$, phase transitions are absent and the voltage-composition curve shows a monotonic characteristic that was first described in terms of a Gaussian distribution of states related to energetic disorder.¹⁰⁸ Recently, it was shown that contributions to the chemical potential of Li^+ inside the amorphous film caused by host distortions play a central role both in the voltage–composition curve and in the ion diffusion effects.^{109–112} In this approach the electrochemical potential takes the form

$$\eta = E_0 + (1+\gamma)Gx^{\gamma} + \frac{1}{k_{\rm B}T}\ln\frac{x}{1-x}.$$
 (17)

The first term E_0 in eqn (17) corresponds to a constant energy level, the second one $(1 + \gamma)Gx^{\gamma}$ relates to the dependence of the intercalant–host interaction with the insertion level. For G > 0 additional energy is needed to deform the host. A power-law dependence with composition with $\gamma < 1$ was reported.¹⁰⁹ The last summand accounts for the entropic term. Chemical capacitance results in

$$C_{\mu} = C_0 \left[\frac{\gamma(1+\gamma)}{k_{\rm B}T} G x^{\gamma-1} + \frac{1}{x(1-x)} \right]^{-1}.$$
 (18)

where $C_0 = q^2 N / k_B T$ is a constant of the order 20 kF cm⁻³ depending on N.

Measurements of the chemical capacitance in a-WO₃ films of different thickness agree well with eqn (18), as shown in Fig. 12. Remarkably, the effect of elastic interactions is a strong function of the film thickness. For the thinnest film (100 nm), the ideal statistics of a dilute solution (corresponding to non-interacting lattice gas model), given in eqn (3), is observed, while the Gx^{γ} term becomes increasingly dominant for larger thickness.¹¹¹ Further studies¹¹² related the intercalation thermodynamics with the models for elastic interactions due to film expansion.^{113,114}

2.3 Depletion capacitance in nanowires

Nanowires and nanotubes are drawing a tremendous attention due to their potential applications in various nanoscale devices. In particular ZnO can be easily induced to grow in the form of rods with high aspect ratio and nanometric dimen-



Fig. 12 Equilibrium (chemical) film capacitance C_{μ} as a function of the composition (molar fraction) x for a-Li_xWO₃ film thickness (in nm, from bottom to top): 100, 200, 300, and 400. Fine solid lines correspond to fits. Capacitance is multiplied by 2, 4 and 8 for thicknesses 200, 300 and 400 nm, respectively, to improve the clarity of the graph. A reference line of slope 1 is shown. Reprinted with permission from ref. 112, copyright 2006 American Chemical Society.

sions. Recently, some papers have addressed the electrochemical properties of electrodes formed by an array of ZnO rods.^{115–117} So far we have considered in the previous section capacitance models where both the Fermi level and conduction band level are homogeneous inside the semiconductor nanostructure. However, when the semiconductor is highly doped and/or the minimal size increases, the Debye screening length shrinks well bellow the nano-object dimensions, and an internal field distribution, associated with band bending, is obtained. As in classical photoelectrochemistry,¹¹⁸ the Fermi level in the semiconductor equilibrates with the redox level in solution, resulting in surface depletion layer that is manifest in the Mott–Schottky (MS) formula of the capacitance

$$\frac{1}{C^2} = \frac{2}{N_{\rm D}q\varepsilon} (V - V_{\rm fb}). \tag{19}$$

Here N_D is the donor density in the n-type semiconductor with dielectric constant ε , V is the voltage and V_{fb} is the flatband potential, that may be displaced due to the Helmholtz layer at the solution side of the interface.¹¹⁹

When an array of semiconductor nanorods as that shown in Fig. 2 is immersed in solution, the outer surface of the rods is depleted of carriers, forming surface bandbending in the radial direction, while the central region of the rods is a conducting (quasi-neutral) tubular region connected to the substrate, as indicated in Fig. 13. Note that the structure shown in Fig. 13 is ideal for channeling electrons towards the collecting contact, avoiding recombination at the surface, in DSC-type solar cells. Since the whole surface of the rods is an equipotential, the surface barrier can be manipulated by modifying the voltage of the substrate with respect to solution. Further, the depletion layer has a circular shape, Fig. 13, and this introduces a strong modification to the standard MS relationship, eqn (19).¹¹⁵

In order to calculate the capacitance of the depletion layer in the geometry of Fig. 13, we consider the charge outside a cylinder of radius r

$$Q = qN_{\rm D}\pi (R^2 - r^2)L.$$
 (20)



Fig. 13 Electrical field and carrier distribution in a semiconductor rod of radius R in contact with electrolyte. E_c is the conduction band energy, E_F is the Fermi level and V_s the electric potential at the surface. The surface depletion layer and the central conducting region (radius r_n) are indicated.

Applying Gauss law, the electrical field is

$$F = \frac{Q}{2\pi r L\varepsilon} = \frac{q N_{\rm D} (R^2 - r^2)}{2r\varepsilon}.$$
 (21)

Let r_n be the radius of the quasi-neutral region. The potential across the depletion layer $V = V_n - V_s = V_{app} - V_{fb}$ is

$$V = \int_{r_n}^{R} F \, \mathrm{d}r = -\frac{qN_{\rm D}}{2\varepsilon} \left[\frac{1}{2} (R^2 - r_{\rm n}^2) + R^2 \ln \frac{r_n}{R} \right].$$
(22)

Now we take the derivatives in eqn (20) and (22)

$$\frac{\mathrm{d}Q}{\mathrm{d}r_n} = -2N_\mathrm{D}\pi r_n L \tag{23}$$

$$\frac{\mathrm{d}V}{\mathrm{d}r_n} = -\frac{qN_\mathrm{D}}{2\varepsilon r_n} (R^2 - r_n^2) \tag{24}$$

therefore the capacitance of the rod wall is given by

$$C = \frac{\mathrm{d}Q}{\mathrm{d}V_{\mathrm{s}}} = \frac{4\varepsilon\pi r_n^2 L}{(R^2 - r_n^2)} \tag{25}$$

and the specific capacitance per unit surface is

$$c = \frac{2\varepsilon r_n^2}{R(R^2 - r_n^2)}.$$
 (26)

Now eqn (22) and (26) when combined provide the voltagedependence of the capacitance, which is illustrated in Fig. 14 for different levels of doping. For the less-doped sample, Fig. 14(a), the MS plot shows a large curvature due to the restriction of the depletion layer in the tubular geometry. As the doping increases, Fig. 14(b), the depletion layer is confined in a short region close to the surface and the straight MS line, eqn (19), is recovered. Experimentally, it has been found that



Fig. 14 (a) and (b) Simulation of a Mott–Schottky plot for the specific surface capacitance of a semiconductor rod with R = 50 nm, $\varepsilon = 10 \varepsilon_0$, and different donor densities as indicated. (c) Experimental data of electrodeposited ZnO rods array on TCO, after annealing 30 min in air, rod dimensions R = 105 nm, L = 950 nm, density 2.40×10^9 cm⁻². The line is a fit to the model described in text that gives $N_{\rm D} = 1.78 \times 10^{18}$ cm⁻³. Data courtesy of I. Mora-Seró.

electrochemically grown ZnO rod arrays show a very high doping of the order 10²⁰ cm⁻³, which decreases considerably by thermal annealing.¹¹⁵ Fig. 14(c) shows a characteristic MS plot of the capacitance measured in the ZnO nanorods array. Note that the good fit to the model is expected in highly monodisperse samples. Carrier densities of 10¹⁹ cm⁻³ have also been found for the ZnO nanorods prepared by chemical bath.¹¹⁷

3. Transport properties

The electronic transport properties of assemblies of colloidal particles vary strongly depending on the individual properties of the nanoparticles, the extent of contacting or electronic coupling between the particles, and the overall geometrical configuration of the assembly.

In the nanostructures formed by sintering of low-doped colloidal nanoparticles, electronic continuity is formed at the contact between adjacent particles. The assembly could be treated in principle as a conventional semiconductor with a band of extended states. The salient difference with respect to monolithic semiconductors, is that the electron density can be varied easily in the whole nanostructure by potential control from the substrate, or photoinjection, as already explained in previous sections, while in compact semiconductors, only the surface density can be varied. In addition, in nanostructured semiconductors the large ratio of surface to volume usually produces a large amount of surface states in the bandgap. The presence of impurities and other factors not totally clear as yet, also cause bulk trap states in the inner part of the nanoparticles, as described above in relation to the DOS in different systems. Therefore the influence of traps on the transport must be considered.

The simplest approach to take trapping into account is the classical multiple trapping (MT) framework.^{120–123} In this approach transport through extended states is slowed down by trapping–detrapping events, while direct hopping between localized states is neglected. The mobility decreases rapidly below a certain value of energy defining the transport states, so that the motion of a bound electron is limited by the rate of thermal excitations to $E \ge E_c$.

Detailed information on the physical parameters related to transport in DSC has been obtained using small perturbation techniques at a fixed steady state such as IMPS^{72,73} and EIS.^{38,65,66,78} Short range electron displacement in nanoporous semiconductors and DSC has been described in terms of the continuous time random walk (CTRW) formalism,124 but the most widely used approach to long range electronic motion, involving macroscopic transport equations, is the MT model, which has been described in a number of papers.^{68,72,125–127} It was found that both the effective electron diffusion coefficient, D_n , and the effective electron lifetime, τ_n ,^{128,129} that are measured become a function of the steady state.^{13,68,72,129–135} Using quasi-equilibrium arguments, the variations of both diffusion coefficient and lifetime were attributed to the statistics of electrons in the material, which deviates from dilution, as described by thermodynamic factors.¹³⁶ The varying D_n was recognized as a chemical diffusion coefficient, 55,127 and the correlation between variations of D_n

and τ_n ,¹³⁵ was explained by a common origin of their variations in an exponential distribution in the bandgap.^{95,136} The application of the multiple trapping model in DSC will be described in detail in section 3.4.

There are many situations in which it is necessary to consider the transitions between localized levels as the main transport mechanism. Wave functions of localized states decay with the increase of the distance from localization center r. This decay occurs at some characteristic distance a and can be often approximated by:

$$\psi(r) = \psi_0 \exp(-r/a). \tag{27}$$

Therefore, the tunnelling probability decreases exponentially with increasing distance. On the other hand the probability of phonon-activated transitions towards higher energy levels also depends exponentially on the energy difference. When the distribution is very wide in the energy scale, it is necessary to consider the combination of probabilities for hopping at different distances and hopping at different levels, as is well established in the theories of hopping conductivity for bulk amorphous semiconductors,^{137,138} and this will be summarily discussed below.

The extent of disorder is an important aspect for characterizing the electronic properties of nanostructured materials, first because of the dispersion of energy of charge carrier within quantum dots, but also due to the influence on long range paths for electron transport. Nanoparticulate film architectures vary from spatially regular to random nanoparticulate networks that form highly disordered structures. Geometrical disorder may become increasingly important for more open structures, due to the existence of highly branched particle structures that influence electron transport dynamics. Studies of electron percolation^{139,140} show that for compact TiO₂ films (40% porosity) used in DSC, the average coordination number is about 6.6, whereas for open-structured films (80% porosity), the average number of particle interconnections is as low as 2.8. Therefore, increasing the porosity has the effect of increasing the average number of particles visited by electrons by 10-fold, from 10⁶ to 10⁷ during their respective transit through 50 and 75% porous 10 µm thick films.140 Fig. 15 shows the reported¹⁴¹ evolution of the diffusion coefficient of electrons in nanostructured TiO₂, as a function of the excess porosity above the critical value for transport. Samples of different porosity are achieved by consecutively pressing the TiO₂ film, from 7 to 4 um thickness. Therefore energy distribution and trapping factors are not significantly changed for different porosity samples. The results show a change of the diffusion coefficient due to the geometrical effect of the coordination between nanoparticles.

In the following we will consider in more detail some basic concepts that are important for the interpretation of measurements of transport in a connected array of nanoparticles. First the influence of non-ideal statistics on the measured quantities such as the diffusion coefficient will be described, and also the deviation from standard Einstein relationship due to thermodynamic factors. Then, a number of specific models will be reviewed in detail, in order to show the effects of the DOS and interactions between carriers, on the measurable macroscopic transport coefficients, when the Fermi level is displaced in the



Fig. 15 Plot of the effective (chemical) diffusion coefficient of electrons, obtained from transient photocurrents, as a function of the porosity of TiO_2 layers. The critical porosity was set to 0.76. Reprinted with permission from ref. 141. Copyright 2006, American Institute of Physics.

nanostructure by potentiostatic control from a substrate. The range of models investigated is summarized in Fig. 16. Free (non-interacting) carriers are considered in a first instance and then the effects of interactions are explored. The hopping conductivity in a broad DOS will be also commented.

3.1 Diffusion and chemical diffusion coefficient

The diffusion of carriers in disordered and nanostructured media is affected by several factors: the available states for hopping, the interactions between carriers, the extent of shielding by surrounding media, etc. Therefore, the relationship between the microscopic hopping mechanisms and the macroscopic transport coefficients is not always straightforward and must be carefully defined. In the following we present a useful framework that dates back to the statement by Onsager and Fuoss of the thermodynamic component of the diffusion coefficient in non-ideal solutions.¹⁴² These ideas have been vastly applied in surface diffusion¹⁴³ and in the simulation of model systems consisting of interacting particles diffusing on the lattice.^{144,145} Recently, the chemical diffusion coefficient has been found quite useful for rationalizing the transport of electrons in metal-oxide nanostructures, 55,127 and it allows also the formulation of the generalized Einstein relation in a very clear way, as discussed later. All these concepts have been well known for a long time in the field of electrochemistry of inorganic solids.⁵¹ It was pointed out¹⁴⁶ that thermodynamic factors play an important role in Li diffusion in intercalation materials, and a correlation of chemical capacitance and the Li chemical diffusion coefficient is often observed in such materials.¹⁴⁷ In nanostructured semiconductor and organic conductors, a broad DOS may cause dominant effects on transport coefficients. It is therefore important to adapt these concepts to electron transport in the presence of energy disorder, and to show the application with examples.

When an electronic or ionic species with concentration n diffuses in a material, the true driving force for diffusion is the gradient of its chemical potential. In the Onsager form of the diffusion law, a linear relationship is assumed¹⁴² between the



Fig. 16 Schematic representation of several transport models in an array of quantum dots or in a distribution of electronic states. (a) Hopping transport through a unique level. (b) Transport in a single level affected by trapping in a deeper level. (c) Hopping transport in a two-level system. (d) Transport between discrete energy levels in an array of quantum dots. (e) Conduction band transport affected by trapping in a wide distribution of states in the bandgap. (f) Hopping transport in a wide distribution of states in the bandgap. $E_{\rm Fn}$ is the Fermi level, $E_{\rm c}$ the conduction band energy, and E_i are the energies of discrete levels.

diffusive flux and the gradient of the chemical potential

$$J_n = -\frac{nu_n}{q} \frac{\partial \mu_n}{\partial x}.$$
 (28)

The prefactor $L_n = nu_n/q$ is known as the Onsager coefficient. On another hand, diffusion is also formulated in terms of the *concentration* gradient in Fick's form,

$$J_n = -D_n \frac{\partial n}{\partial x}.$$
 (29)

The coefficient D_n in eqn (29) is called the *chemical diffusion* coefficient.^{144,148} It is given by

$$D_n = \left(\frac{k_{\rm B}T}{q}u_n\right) \left(\frac{n}{k_{\rm B}T}\frac{\partial\mu_n}{\partial n}\right) \tag{30}$$

 D_n contains two components: (i) a phenomenological coefficient u_n (the mobility) and (ii) the term $n\partial \mu_n / \partial n$, that accounts for the difference between a gradient in concentration, and a gradient in chemical potential. This last term is expressed in dimensionless form as the *thermodynamic factor* introduced by Darken¹⁴⁹

$$\chi_n = \frac{n}{k_{\rm B}T} \frac{\partial \mu_n}{\partial n}.$$
 (31)

Note that $\chi_n = 1$ in the case of Maxwell–Boltzmann distribution of eqn (4). Let us introduce the jump (or kinetic) diffusion coefficient,¹⁴³ which is simply proportional to mobility

$$D_J = \frac{k_{\rm B}T}{q} u_n. \tag{32}$$

Now we can write the chemical diffusion coefficient as the product

$$D_n = \chi_n D_J. \tag{33}$$

Alternatively, we have

$$D_n = \frac{k_{\rm B}T}{q} \chi_n u_n. \tag{34}$$

The phenomenological quantities introduced so far can be given more precise meaning in terms of statistical concepts. The thermodynamic factor can be expressed with respect to the chemical capacitance as

$$\chi_n = \frac{q^2 n}{k_{\rm B} T} \frac{1}{C_\mu} \tag{35}$$

and the chemical capacitance is related to the mean-square fluctuation of the particle number N in a volume V by the formula¹⁵⁰

$$\langle (\delta N)^2 \rangle = \frac{k_{\rm B} T V}{q^2} C_{\mu} \tag{36}$$

where $\langle\rangle$ denotes a statistical average. Therefore the thermo-dynamic factor is

$$\chi_n = \frac{\langle N \rangle}{\langle (\delta N)^2 \rangle}.$$
 (37)

The chemical diffusion coefficient D_n in eqn (33) can be derived from a microscopic approach using Green–Kubo theory.^{144,145} The jump diffusion coefficient has the form

$$D_J = \frac{1}{6t} \left\langle \left(\frac{1}{N} \sum_{i=1}^N \Delta r_i \right)^2 \right\rangle$$
(38)

where Δr_i is the displacement of the *i*th particle at time *t*. In a regular lattice of spacing *l*, D_J can often be expressed as

$$D_J = \frac{1}{6} \Gamma(n) l^2 \tag{39}$$

in terms of a mean effective jump frequency $\Gamma(n)$, that varies with the occupancy of the lattice.^{143,148}

It is important to remark that routine electrochemical methods, based on a step of the voltage (either in time or frequency domain), measure the chemical diffusion coefficient, see *e.g.* ref. 151 and 152. In contrast, in numerical simulation methods, the quantities χ_n and D_J are evaluated separately, and then combined to form D_n by the so-called indirect method, ^{153,154} see *e.g.* ref. 155–158. The term χ_n can be determined in grand canonical Monte Carlo simulations by use of the fluctuation formula in eqn (37). The kinetic factor is obtained by monitoring the random walk in the lattice¹⁴⁹

$$D^* = \lim_{t \to \infty} \frac{1}{6Nt} \left\langle \sum_{i=1}^{N} \left(\Delta r_i \right)^2 \right\rangle.$$
(40)

More precisely, eqn (40) defines the tracer diffusion coefficient, D^* , that reflects random walks of a particle, while the jump (or kinetic) diffusion coefficient defined by eqn (38) reflects diffusion of the centre of mass of N particles. The difference between these two coefficients (the Haven ratio) is small, usually less than a factor of 2.

3.2 The Einstein relation

The ratio of the mobility and the diffusion coefficient is known as the Einstein relation. The application of this relation in systems with a broad energy disorder is not always transparent, therefore we discuss in the following the statement of the Einstein relation for systems in a quasi-equilibrium regime in terms of the transport coefficients that have been introduced.

We consider the motion of electrons in an organic or inorganic semiconductor material, with concentration n(x)and electric field $F(x) = -\partial \phi / \partial x$ at position x. The electrical current is given by the sum of conduction and diffusion currents

$$j_n = qnu_n F + qD_n \frac{\partial n}{\partial x}.$$
(41)

The electrochemical potential of electrons is $\eta_n = -q\phi + \mu_n$. In equilibrium we have $\partial \eta_n / \partial x = 0$ and consequently

$$q\frac{\partial\phi}{\partial x} = \frac{\partial\mu_n}{\partial x}.$$
(42)

Therefore eqn (41) gives

$$j_n = \left(-nu_n + qD_n\frac{\partial n}{\partial \mu_n}\right)\frac{\partial \mu_n}{\partial x}.$$
(43)

Since the current in equilibrium is $j_n = 0$, the parenthesis in eqn (43) is zero. This imposes a relationship (Einstein's) between the mobility and the chemical diffusion coefficient. Such a relation can be stated in two equivalent forms. Eqn (34) is the generalized Einstein relation,¹⁵⁹ and it differs from the classical Einstein relationship by the thermodynamic factor, so that for classical statistics, with $\chi_n = 1$, the traditional relation, but it includes the jump diffusion coefficient. These relations were already obtained with hindsight in the previous section, by including the mobility as the transport coefficient in the diffusion law, eqn (28). Now eqn (41) can be written in a

similar form, as follows:

$$j_n = -nu_n \frac{\partial \eta_n}{\partial x}.$$
(44)

This linear relationship between the electrical current and the gradient of the electrochemical potential, η_n , is usually employed in electronic device modeling, and it is also used in nanostructured photoelectrochemical solar cells.^{160,161}

It is possible to adopt an alternative expression for the mobility that includes the factor χ_n .^{91,162} In this procedure, it is assumed that in the quasi-equilibrium situation only electrons within $k_{\rm B}T$ of the Fermi level contribute significantly to the conductivity.^{163,164} The effective density of carriers \hat{n} is given by¹⁶⁴

$$\hat{n} = k_{\rm B} T \frac{\mathrm{d}n}{\mathrm{d}\mu_n} = \frac{n}{\chi_n}.$$
(45)

In terms of the normal mobility, the electron conductivity is

$$\sigma_n = nqu_n. \tag{46}$$

Then one can define an effective carrier mobility $\hat{u}_n = \sigma_n/q\hat{n}$,

$$\hat{u}_n = \frac{q}{k_{\rm B}T} D_n. \tag{47}$$

It is clear that \hat{u}_n expresses the chemical diffusion coefficient in units of mobility, while the standard mobility relates to the jump diffusion coefficient, as stated in eqn (32). When discussing experimental data, it should be born in mind that both mobilities are related as $u_n = \chi_n \hat{u}_n$, so they can have very different values. For example in assemblies of quantum dots the thermodynamic factor can take large values,^{165,166} as discussed below.

Using eqn (34) and (35), the conductivity can be expressed in terms of chemical diffusion coefficient and chemical capacitance as

$$\sigma_n = C_\mu D_n. \tag{48}$$

This last expression can also be written as

$$\sigma_n = q^2 D_n \frac{\mathrm{d}n}{\mathrm{d}\mu_n} \tag{49}$$

which is the standard form in studies of metal-insulator transition, for example.¹⁶⁷ However, it appears that eqn $(48)^{55}$ could be suitable for the analysis of electrochemistry experiments, since the three quantities appearing in eqn (48) can be measured directly and separately by electrochemical methods. In contrast, the mobility u_n , is usually obtained from the conductivity and the total carrier number, eqn (46).^{168,169} In a system with a peak-shaped DOS, there are site-saturation effects affecting the mobility. When the number of carriers *n* is very large the conductivity decreases and eventually becomes null, which implies that $u_n \rightarrow 0$, according to eqn (46), see Fig. 17(c). This is in fact observed in the electrochemistry of conducting polymers due to the finite (Gaussian) DOS.^{170,171}

In recent years, the relevance of eqn (34) (in different formulations) has been increasingly realized in relation with disordered materials with a broad distribution of states in the energy space. For example non-ideal Einstein relation was observed in hydrogenated amorphous silicon¹⁷² and this was

explained in terms of the thermodynamic factor of the exponential distribution.¹⁷³ The same thermodynamic factor, which is given below in eqn (55), is observed for electron diffusion in nanoporous TiO_2 in ref. 127. Several works have also discussed the Einstein relation in organic semiconductors with a Gaussian distribution of electronic or polaronic states.^{174–179}

3.3 Transport along discrete energy levels

In this section we consider a number of models for electron transport along discrete energy levels. The models can be applied to the hopping of electrons in arrays of quantum dots, or electron transport between localized sites in a homogeneous semiconductor. The main aim of these models is to obtain the parameters of macroscopic diffusion (chemical capacitance, chemical diffusion coefficient, *etc.*) as a function of the Fermi level, in relation to the microscopic properties determining electron accumulation and hops. We will analyze step by step models of increasing complexity. We start with the assumption of independent carrier transport in a single energy level, then in several energy levels and thereafter discuss the effects of short-range interactions.

We first consider a simple model that introduces some important concepts related to the finite occupation of electron sites.⁵⁵ It is a 3-dimensional solid composed of a cubic lattice of weakly interacting quantum dots of radius *R* and separation *d*, with a single electronic state at the energy level E_0 , that can be occupied at most by one carrier, as indicated in Fig. 16(a). This model neglects all interactions and also stands, with obvious modifications, for electron hopping in a single energy level in a nanostructured material. The model, with addition of mean-field interactions, is also applied for Li-ion diffusion in battery materials such as Li_{1-x}CoO₂.¹⁰⁶

Since the electrons do not interact with each other, the random statistical distribution is valid, and the probability of occupancy is determined by Fermi statistics, eqn (6). The single electronic state is filled when the Fermi level crosses the site energy, as seen in Fig. 17(a). The chemical capacitance takes the form of eqn (9), where $N = l^{-3}$ is the volume density of electronic states. The thermodynamic factor is

$$\chi_n = \frac{1}{1 - f}.\tag{50}$$

When $E_{\rm Fn} \approx E_0$ the chemical capacitance peaks, Fig. 17(b). Thereafter the statistics departs strongly from dilution and the thermodynamic factor increases by orders of magnitude because of the departure from ideality by the exclusion "interaction", Fig. 17(b): a carrier in a localized site exerts an infinite repulsion over others attempting to hop to that site. Ref. 180 discusses the Einstein relation in this model.

The average distance travelled by an electron in one hop between dots is l = 2R + d. The mean effective jump frequency is $\Gamma(f) = \nu(1 - f)$, which gives the number of jumps into empty sites per unit time. The hopping rate constant ν for electron hopping between neighboring dots has the form $\nu =$ $\nu_0 \exp(-2d/a)$, in terms of the decay length *a* of the wave function in the localized state, and the attempt-to-hop frequency ν_0 .¹⁸¹

In the case $f \ll 1$ (*i.e.*, $E_{\rm Fn} \ll E_0$) one obtains the diluted limit corresponding to Maxwell–Boltzmann statistics. The occupancy restriction is lifted and $\chi_n = 1$. Let us define the diffusion coefficient in the dilute limit, which is a constant $D_0 = l^2 \nu/6$. For arbitrary f the jump diffusion coefficient can be written $D_J = (1 - f)D_0$ in terms of the diluted value. When χ_n increases the jump diffusion coefficient decreases considerably due to scarcity of vacant neighbors to accept hopping electrons, Fig. 17(c). The occupancy dependencies of χ_n and D_J compensate exactly giving the *constant* chemical diffusion coefficient $D_n = D_0$,⁵⁵ see Fig. 17(c). In general this model is a good first approximation to the materials with a peak-shaped chemical capacitance. As



Fig. 17 Representation of several quantities for electron accumulation and diffusion in a cubic array of QDs 3 nm diameter and 1 nm interdot distance. (a) Isotherm (number of electrons *vs.* potential). (b) Chemical capacitance and thermodynamic factor. (c) Jump (D_J) and chemical (D_n) diffusion coefficients. Parameters: T = 300 K, $\nu_0 = 10^{12}$ s⁻¹, $a = 0.2 \times 10^{-7}$ cm.

mentioned above, it should be observed that the mobility u_n (proportional to D_J) decreases when the carrier number saturates at high Fermi level, producing a decrease of the conductivity. It should also be remarked that D_n and u_n behave in very different ways.

We now consider the effects of energy distribution in the electronic states. The simplest approach is to assume a static distribution of energy levels. The sites have different energies that are independent of occupation of neighboring sites. The probability of occupancy is determined by Fermi statistics with a common chemical potential (Fermi level) for the different levels. A two-level system is indicated in Fig. 16(b), and it describes well certain ion–intercalation systems.^{183–185} This is a model in which carrier transport occurs only through the shallow level, while the deep state captures and releases carriers, which affects the transient behaviour of the system.^{182,186} This assumption is the basis of all multiple trapping models.

Main features of the two-level model with trapping transport of Fig. 16(b) are shown in Fig. 18.¹⁸² Thermodynamic quantities in this model are composed of a combination of the features of the separate states, which were discussed in Fig. 17. First Fig. 18(a) shows the consecutive charging of the two levels as the Fermi level increases, and Fig. 18(b) shows the corresponding peaks of the chemical capacitance. The thermodynamic factor shows a peak at the intermediate values of energy between the two levels, and then increases when the two-level system approaches full occupancy. Since the transport along the shallow level is delayed by the trapping in the deep level, when the later is filled there is a strong increase of the chemical diffusion coefficient, as shown in Fig. 18(d). However, the conductivity, represented by diffusion resistance¹⁸² in Fig. 18(c), is not affected by the occupancy of the deep level. This is because the conductivity is a steady-state property, that reflects only the hopping along the transport (shallow) level.

Let us consider the effect of interactions between particles in nearest neighbor sites (or quantum dots). A calculation of Monte Carlo simulations¹⁸⁷ of a two-dimensional two-level lattice gas with repulsive interactions between nearest neighbors is shown in Fig. 19. $J = E_2 - E_1$ is the interaction energy between carriers on nearest neighbor sites, where E_1 and E_2 are site energies ($N_1 = 1/3N$, $N_2 = 2/3N$). In this model system, the transport goes through the lines of Fig. 19(a), as suggested in the scheme of Fig. 16(c). The critical temperature of this system is given by $J/k_BT_c = 3.064$,¹⁸⁸ and the results of Fig. 19 correspond to the temperature $T = 1.2T_c$, above the critical value. Therefore the separation between the two energy levels is of the order of the thermal energy, $E_2 - E_1 = 2.55k_BT$.

With respect to the non-interacting two-level system discussed before in Fig. 18, in which the states are occupied sequentially when the Fermi level crosses the state energy, an entirely new feature in Fig. 19(b) is a strong decrease of the occupancy of the deep site when the Fermi level lies between the energies of the two levels, Fig. 19(b). This is reflected in a new peak of the chemical capacitance, Fig. 19(b), or equivalently, a dip in the thermodynamic factor. The inversion of the population of the deep level near c = 1/2 is due to interactions between carriers. At c = 1/3 some particles appear at the shallow sublattice, however, their interaction with particles on the deep sublattice is weak. At c = 1/2 particles on the shallow sublattice force the other particles to move on the shallow sublattice because there are many sites on this sublattice and particles do not interact with each other. It follows, that such a behavior is energetically and combinatorially preferred. This process happens without strong external influence what leads to high chemical capacitance at c = 1/2. The jump diffusion coefficient is characterized by minimum value at c = 1/3because at this concentration almost all the deep sites are occupied. When the concentration is approaching this value and a particle jumps to a nearest shallow site it is repelled back



Fig. 18 Representation of a two state system with $E_1 = E_0 - 10k_BT$ and sites concentration $N_0 = 1/3N$, $N_2 = 2/3N$. (a) Occupancy dependence on voltage. (b) Chemical capacitance and thermodynamic factor. (c) Diffusion resistance. (d) Chemical diffusion coefficient.¹⁸²



Fig. 19 (a) Arrangement of sites in the two-level systems. The lines indicate the transport pathways. Results of Monte Carlo simulations for this system at $T = 1.2T_c$: (b) Fractional occupancies of the two levels and total number of particles; (c) chemical capacitance and thermodynamic factor and (d) jump diffusion coefficient, as a function of the electrochemical potential. Data courtesy of V. S. Vikhrenko, adapted from ref. 187.

by its neighbors on the nearest deep sites. At c > 1/3 some particles must be on the shallow sublattice and thus the mobility sharply increases due to "exclusion" mechanisms.¹⁸⁷ A complete calculation of one-dimensional diffusion in two-level systems with interactions is given in ref. 154.

Next we consider a study of electron transport in a cubic array of quantum dots with discrete energy levels, see Fig. 16(d). The energy for addition of electrons to a quantum dot is considered in the scheme of Franceschestti and Zunger, 189,190 taking into account the self-energy of the electron interacting with its image charge at the surface of the particle, a weak electrostatic interaction of electrons in a dot, due to screening by the external medium, and the exchange energy.¹⁶⁶ The rate of transfer between energy levels in different dots consists of the Miller-Abrahams hopping rate¹⁸¹ considering only nearest neighbors. Disorder is incorporated in the array of quantum dots via a log-normal distribution of sizes with relative standard deviation σ_R that causes dispersion from the size dependence of the singleparticle energy levels and that of the electrostatic interaction and polarization energies. The conductivity is calculated from Monte Carlo simulations of the electron flux maintaining a small difference of electrochemical potential at the contacts. The mobility is determined from eqn (46) and the chemical diffusion coefficient is obtained by determining the electron gradient with eqn (29). Then the thermodynamic factor is obtained from eqn (34).

In Fig. 20 several quantities are represented as a function of the electrochemical potential of the injected electrons, both for low and high disorder. Fig. 20(a)–(c) show the thermodynamic features corresponding to electron charging and the associated thermodynamic factor and electrochemical capacitance. It was shown¹⁶⁶ that these features can be successfully modelled using a mean-field approximation and the equilibrium definition of the thermodynamic factor, eqn (31), with respect to the

electrochemical potential. A salient characteristic of the results is the presence of strong Coulomb blockade effect for the lowdisorder sample when completing the 1S₀ shell at n = 2 and the 1P₀ shell (n = 8) at $\eta_n = 3.3$ eV. This gives rise to a large increase in the thermodynamic factor, which increases by a factor of 1000 at n = 2 and a factor of 10 at n = 8. Additional minor peaks are obtained when the average occupation number is an integer. However, for the array with $\sigma_R = 0.1$ these features are almost completely smoothed. These results indicate that the apparition of Coulomb blockades necessitates both small size dispersion and a Coulomb interaction energy that is larger than k_BT .

A main result obtained in this study¹⁶⁶ is the considerable difference between Fermi level dependencies of the mobility and chemical diffusion coefficient in the low-disorder array, as a result of the large thermodynamic factor. Thus while the diffusion coefficient only decreases by a factor of 10 in the $1S_0$ shell-filling region, the mobility shows a strong minimum with a decay of three orders of magnitude.¹⁹¹ It is also noteworthy that the conductivity, Fig. 20(f), when averaged over local peaks remains nearly constant during the filling of the P shell, due to compensation of the increase in carrier density with the decrease in the mobility.

3.4 Transport in a continuous density of states

Multiple trapping in a broad distribution of states. We have already mentioned that results of electron transport in DSC have been largely interpreted in terms of the MT model in a continuous distribution of states. Let us analyze this model, represented in Fig. 16(e), in more detail. Electronic states are composed of a transport state (usually identified with the lower edge of the conduction band) at the energy level E_c , with a diffusion coefficient D_0 , and a density of localized states g(E) distributed in the bandgap. The total electron density is



Fig. 20 Representation of several quantities for electron accumulation and diffusion in a cubic array of QDs 3 nm mean radius and 2 nm interdot distance, parametric on size dispersion. (a) Isotherm (number of electrons *vs.* potential). (b) Thermodynamic factor. (c) Chemical capacitance. (d) Mobility. (e) Chemical diffusion coefficient. (f) Conductivity. Parameters: T = 300 K, $\nu_0 = 10^{12}$ s⁻¹, $a = 0.5 \times 10^{-7}$ cm. Data courtesy of J. van de Lagemaat, adapted from ref. 166.

 $n = n_{\rm c} + n_{\rm L}$, *i.e.* the sum of electron densities in conduction band and localized states, Fig. 9. In general for the multiple trapping model we obtain the thermodynamic factor,⁵⁵

$$\chi_n = \frac{n}{n_c} \left(1 + \frac{\partial n_L}{\partial n_c} \right)^{-1}.$$
 (51)

The mean effective jump frequency is obtained from the MT condition that only free electrons in transport states contribute to the diffusion process, hence the jump diffusion coefficient is

$$D_J = \frac{n_{\rm c}}{n} D_0. \tag{52}$$

Using eqn (32) and (46), the electron conductivity can be written as

$$\sigma_n = \frac{q^2 n}{k_{\rm B} T} D_J. \tag{53}$$

Applying eqn (52), it is found that

$$\sigma_n = \frac{q^2 n_{\rm c}}{k_{\rm B} T} D_0. \tag{54}$$

The only Fermi-level dependent quantity in eqn (54) is the free electron density, and this is observed experimentally in nano-structured TiO_2 .⁵⁸ As already commented in connection with

the two-level system, the steady-state conduction is not affected by the trapping process, because the traps remain in equilibrium. Alternatively, one can view conduction as the result of the displacement of the whole electron density, eqn (53), but with the jump diffusion coefficient, which is not constant.

The chemical diffusion coefficient of electrons, D_n , is given by eqn (30). The effect of trapping in the chemical diffusion coefficient is dominant when $\partial n_{\rm L}/\partial n_{\rm c} \gg 1$. In this case the result is

$$D_n = \left(\frac{\partial n_{\rm c}}{\partial n_{\rm L}}\right) D_0. \tag{55}$$

The prefactor in eqn (55) is the relationship of free to trapped number of electrons for a small variation of the Fermi level. This prefactor describes the delay of response of the chemical diffusion coefficient, with respect to the free electrons diffusion coefficient, by the trapping and detrapping process.^{136,192} Such a delay is unavoidable when measuring the chemical diffusion coefficient by any transient technique. In the quasistatic approximation¹³⁶ the effect of electrons densities. This is related to application of the principle of detailed balance, that links the kinetic constants for trapping and detrapping to the

equilibrium occupancies.¹⁹³ Using the zero temperature limit of the Fermi function, it is obtained that the reciprocal of the prefactor in eqn (55) is given by¹³⁶

$$\frac{\partial n_{\rm L}}{\partial n_{\rm c}} = \frac{C_{\mu}^{\rm traps}}{C_{\mu}^{\rm cb}} = \frac{k_{\rm B}T}{n_{\rm c}}g(E_{\rm Fn}) \tag{56}$$

see eqn (14) and (15).

Multiple trapping in exponential distribution. Let us consider in particular the exponential distribution of localized states, with a characteristic energy $k_{\rm B}T_0$, given in eqn (10). As mentioned before it is well established that nanostructured TiO₂ used in DSC (anatase) shows this type of distribution of states in the bandgap,^{13,40,66,82,95} and even the direct monitoring of Fermi level has been realized.^{194,195} The main features of the MT model for this distribution⁵⁵ are illustrated in simulation in Fig. 21 with realistic parameter values.^{82,125} We first discuss the regime of electrochemical potentials in which the Fermi level is well below the conduction band. Hence the free electron density is much lower than trapped electrons number, $n_c/n_{\rm L} \ll 1$.

Fig. 21(a) shows the extent of electron charging, and Fig. 21(b) shows the chemical capacitance of the localized states, given in eqn (15), with a slope $(k_B T_0)^{-1}$. The thermodynamic factor is constant⁵⁵

$$\chi_n = \frac{n}{n_c} \frac{\partial n_c}{\partial n_L} = \alpha^{-1} \tag{57}$$

where

$$\alpha = T/T_0. \tag{58}$$

For the typical values of T_0 , $\chi_n \approx 5$ at room temperature, as shown in Fig. 21(b). The calculation of the jump and chemical

diffusion coefficient, eqn (30), gives, respectively⁵⁵

$$D_J = \alpha D_n \tag{59}$$

$$D_n = \frac{N_c T_0}{N_L T} \exp\left[(E_{\rm Fn} - E_c) \left(\frac{1}{k_{\rm B} T} - \frac{1}{k_{\rm B} T_0} \right) \right] D_0.$$
(60)

Results of the diffusion coefficients are shown in Fig. 21(c).

Fig. 21 also shows the evolution of parameters when Fermi level approaches the conduction band and $n_c \approx n$. The chemical capacitance of the conduction band states is given in eqn (14) and its slope corresponds the thermal energy value $(k_BT)^{-1}$, reflecting the Maxwell–Boltzmann distribution of the dominant free carriers. The thermodynamic factor takes the value of 1 when approaching the conduction band from below, due to the onset of ideal statistics. In this domain MT trapping ceases to hold, because the different kinetic phenomena occur directly through the conduction band states with no interference of the traps (which are nearly full of electrons) and both D_J and D_n approach D_0 .

Application of multiple trapping in dye-sensitized solar cells. The MT model just discussed has been very useful for rationalizing the experimental results on electron transport in DSC, for example the observation by IMVS of a chemical diffusion coefficient that varies with the Fermi level in the same way as shown in Fig. 21.^{72,134} The MT model, in combination with the quasistatic approximation¹³⁶ also provides an explanation for the observed compensation between chemical diffusion coefficient and lifetime¹²⁹ dependence on Fermi level, to give a nearly constant electron diffusion length as reported by L. M. Peter and other authors.^{6,15,72,95,135,196}

Characteristic experimental results of the main parameters, obtained by EIS in several DSCs,⁶⁶ are shown in Fig. 22. An exponential dependence of the total film capacitance is



Fig. 21 Representation of several quantities for charge accumulation and diffusion by multiple trapping in a nanostructured semiconductor electrode of thickness $L = 10 \,\mu\text{m}$ with an exponential distribution of bandgap states. (a) Isotherm (total charge vs. potential). (b) Chemical capacitance and thermodynamic factor. (c) Free electrons diffusion coefficient (D_0) , and jump (D_J) and chemical (D_n) diffusion coefficient. Parameters: $N_c = 6.8 \times 10^{20} \,\text{cm}^{-3}$, $N_L = 2.0 \times 10^{19} \,\text{cm}^{-3}$, $T = 300 \,\text{K}$, $T_0 = 1400 \,\text{K}$, $D_0 = 10^{-2} \,\text{cm}^2/\text{s}$.



Fig. 22 Parameters resulting from the fit of experimental EIS spectra of three dye-sensitized solar cells at different bias potentials in the dark. The electrolyte composition in the reference cell is 0.5 M LiI, 0.05 M I₂ and 0.5 M 1-methylbenzimidazole (MBI) in 3-methoxypropionitrile (3-MPN). The second cell has 0.5 M NaI instead of LiI, and the third cell has no MBI. (a) Capacitance of the cell without the contribution of the Pt capacitance. Also indicated is the estimated open-circuit voltage of the cell at 1 sun illumination (measured value is 0.58 V at 0.1 sun). (b) Chemical diffusion coefficient of electrons. (c) Transport resistance in TiO₂. The lines are fits to eqn (61)–(63), and the parameters resulting from fits are indicated. *T* is the actual temperature of the cell, assumed 300 K, *T'* is the effective temperature obtained from the slopes of the plots as indicated, and T_0 is the characteristic temperature of the exponential distribution of localized states in the bandgap. Data courtesy of F. Fabregat-Santiago, adapted from ref. 66.

observed in an intermediate domain of potentials, which is interpreted as C_{μ}^{traps} . At the more *positive* potentials, the metal oxide is insulating and the capacitance observed corresponds to the conducting substrate that is exposed to the solution.¹¹⁹ At the more *negative* potentials the Fermi level should approach the conduction band. As shown in Fig. 21(b) the slope of the capacitance should increase towards the thermal value $(k_{\rm B}T)^{-1}$, but in contrast to this, the experiments in Fig. 22(a) show that the slope decreases at the more negative potentials. This is interpreted as a shift of the conduction band potential due to increasing potential drop at the Helmholtz layer, at the high electron densities considered.⁸² The conduction band is displaced upwards as it is approached by the Fermi level, which makes it fairly difficult to detect the extended transport states of the conduction band.⁵⁵

Let us make an analysis of experimental results on chemical capacitance, chemical diffusion coefficient and diffusion resistance, in order to check the predictions of the MT model with the exponential distribution of states. The potential-dependence of these elements, according to the model, are

$$C_{\mu}^{\rm traps} \propto e^{-qV/k_{\rm B}T_0} \tag{61}$$

$$D_n \propto e^{-qV(1-\alpha)/k_{\rm B}T} \tag{62}$$

$$R_{\rm diffusion} \propto e^{qV/k_{\rm B}T}$$
. (63)

In Fig. 22 the results of these parameters are shown for three DSCs containing different species in the electrolyte. In each

case the absorption of the indicated species in the surface of TiO₂ determines the surface dipole and provokes a global shift in the potential scale (in the same way as in Fig. 3, above). The three elements, C_{μ}^{traps} , D_n and $R_{\text{diffusion}}$, show exponential dependencies in the potential and shift consistently in the voltage scale. We examine the reference cell, containing LiI electrolyte, in more detail. First, in the intermediate domain where the traps capacitance is observed separately, we fit the capacitance to eqn (61) and obtain $k_{\rm B}T_0 = 62 \text{ mV}$ corresponding to $\alpha = 0.42$ at T = 300 K. This implies an exponent $k_{\rm B}T/(1 - \alpha) = 44$ mV for the chemical diffusion coefficient, and this in fact is the value obtained in Fig. 22(b). The diffusion resistance is the reciprocal of the conductivity in eqn (54), therefore the dependence on potential is given by the thermal energy $k_{\rm B}T = 26$ mV at room temperature, eqn (63), and this prediction is also well realized by the data in Fig. 22(c). Recent measurements of the chemical diffusion coefficient in high efficiency DSC at different temperatures further confirmed the predictions of MT transport,⁷⁸ and activation energies for electron transport have been discussed as well using this model.197

The MT model describes well the diffusion coefficient and conductivity of nanostructured TiO₂ when the Fermi level lies deep in the exponential trap distribution in the bandgap. However, the free electrons diffusion coefficient D_0 implicit in the MT model has not been separately identified. The Hall mobility of electrons in large single crystals of anatase-TiO₂ at room temperature has the value $u_{\rm H} = 20 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$,¹⁹⁸ corresponding to the diffusion coefficient $D_0 = 0.5 \text{ cm}^2 \text{ s}^{-1}$.

But such a large value of the electron diffusion coefficient has not been approached in DSC. Measurements of electron diffusion coefficient in nanoporous TiO₂ electrodes, in different electrolytes and determined by different techniques, remain below the value of 10^{-3} cm² s⁻¹.⁷⁸ The upper limitation to the photovoltage in DSC is given when the Fermi level meets the lower edge of the conduction band of TiO2.^{3,199} In Fig. 22(b) we see that the extrapolation of D_n to V_{oc} (at 1 sun) remains lower than 10^{-3} cm² s⁻¹, which is two orders of magnitude below the value expected for bulk crystals. However, some estimates put the open circuit Fermi level at least 150 meV below the conduction band even at 1 sun. Furthermore, electron transport is normally measured under short circuit conditions where the Fermi level is almost 500 meV below the conduction band. Therefore, it is not yet clearly established whether the free electrons diffusion coefficient in DSC is consistent with the value for bulk anatase.

Hopping conduction. The theory of hopping conduction^{137,138} describes the transport by carrier jumps via localized states, as indicated in Fig. 16(f). A general description of hopping conduction in disordered systems with a broad distribution of localized states is a complex problem. Given a distribution of hopping sites, the possible set of transitions are in principle known. However, averaging over all possible spatial and energy configurations in order to find the average jump rate and hopping distance that relate to the observable transport coefficients, is not feasible. The analysis is partially simplified in a system with a steep distribution of localized states, where the hopping process is well described with the concept of transport energy, originally formulated for systems with an exponential DOS.²⁰⁰⁻²⁰⁴ For carriers situated deep enough energetically, a particular level, called the transport energy, $E_{\rm t}$, determines each single hopping event. Therefore, when the Fermi level is well below the transport level, hopping systems behave very similar to a MT system, with the level E_t playing the role of the conduction band edge in the above description. The concept of transport energy has been extended to systems with a Gaussian DOS.²⁰⁵⁻²⁰⁷

Another limiting case of hopping models, first suggested by Mott,²⁰⁸ considers the competition between hopping at different entergies and at different distances, and is generally termed variable-range hopping (VRH). This approach optimizes the hopping rate of a single hop from one site to another under the constraint that at least one such hop is possible. Later, more systematic treatments were initiated that were based on percolation arguments, the so-called critical path analysis.^{137,138} VRH has been observed in a wide variety of systems such as Si- and Ge-based inorganic semiconductors,²⁰⁹ conducting polymers and assemblies of quantum dots.²¹⁰

The hopping conductivity has been amply studied for bulk amorphous semiconductors^{137,138} and organic semiconductors with a Gaussian DOS,^{206,207,211–217} but these developments have not been generally applied in electrochemistry experiments. Arkhipov, Bässler and coworkers have reported some models of the electronic conductivity of electrochemically doped polymers.^{218,219}

In an array of quantum dots with strong spatial confinement and good electronic coupling between dots, a narrow band,

providing coherent transport, can be formed at each discrete energy level. However, there are several factors that lead to localization of carriers in the quantum dots,²²⁰ as discussed in the example of Fig. 20. Colloidal nanoparticles prepared by wet chemical methods fluctuate in size and chemical composition. This introduces some inherent disorder in an array of quantum dots, that produces a dispersion of the energy levels.²²⁰ The difference between energy levels in different quantum dots makes tunneling of electrons between quantum dots difficult. Moreover such dispersion leads to the random scattering of electronic waves. Thus dispersion of quantum dot sizes and fluctuation of the chemical composition contribute to localization.²²¹ The Coulomb repulsion between two electrons (holes) sitting on the same quantum dot introduces an energy gap between occupied and empty states (Hubbard insulator). A number of characteristic phenomena for transport in disordered semiconductors have been reported in arrays of quantum dots, such as metal-insulator transitions^{16,222} and Mott-like conduction gaps,^{59,221} the VRH transport,²¹⁰ and percolation thresholds depending on disorder.²²³

4. Conclusions and outlook

Extensive work over the last decade and a half on nanocrystalline systems and organic conductors in electrochemical configuration has brought a great deal of understanding on these systems. The knowledge initiated from a mixture of the classical concepts of electrochemistry and semiconductor physics. The main novelty came from the fact that the semiconductor Fermi level in electrolyte-surrounded nanocrystalline networks could be homogeneously moved with a potentiostat. But some general features complicate the study of these systems. First, heterogeneity at the nanoscale is always present. Second, there is no attempt at purity or ideality of materials in this field, on the contrary, materials development are driven by the need of simple preparation methods that provide effective outcomes when tested as part of complex devices. Therefore these materials usually present a great extent of disorder and variability depending on preparation methods, properties of the solution, etc. This is why the great deal of interest on nanocrystalline TiO2 for dye solar cells has been a very positive asset. A great number of laboratories and scientists focused their work on this widely accessible system, and in consequence, a large body of experimental results revealed the regularities of the physical electrochemistry (and photoelectrochemistry) of this system. As a result, a number of specific experimental methods in combination with a wellestablished battery of conceptual tools, which we have attempted to review here, has become available for testing the properties of similar nanocrystalline systems.

In my view, it is remarkable that the electronic properties of nanocrystalline semiconductors can now be reliably extracted and described with relatively simple concepts and one-dimensional models, taking into account the intrinsic variability, disorder and heterogeneity of these systems. These methods are now routinely applied to investigate different types of metal-oxide nanocrystalline electrodes. The techniques and concepts mentioned provide an identification of the density of electronic states, diffusion coefficient, lifetime, *etc.*, which constitutes highly valuable information for characterizing the materials prepared and for designing new ones.

However, there are still important challenges in this field. First of all, the electronic behaviour of the nanocrystalline semiconductors consistently changes as a function of the medium (liquid or solid) filling the pores. In part this change is due to modifications of electron lifetimes, due to the variation of interfacial charge transfer properties, however the transport properties are also affected, meaning that some interaction takes place between the electronic carriers in the semiconductor and the surrounding medium. While it has been possible to control the shift of energy levels by absorption of molecular species in the surface, as discussed in the main text, additional effects of interactions are very poorly understood, and there are no swift methods to predict or analyze the behaviour of new electrolytes or organic hole conductors. Another aspect that needs more work is the high accumulation regime; the free electrons' diffusion coefficient has not yet been clearly identified; the transport mechanism at high carrier densities is not really known, and in addition, the electrochemical effects of packing a large number of carriers in small nanoparticles have been little investigated. Finally, an important route of research is the properties of ordered nanostructures such as nanotubes, which are reaching perfection. By removing the uncertainties introduced by spatial disorder, these last systems may provide in the near future considerable information on the physical electrochemistry of nanoscale semiconductors.

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