Electrochemical metamaterials

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The role of electrochemistry in energy systems such as batteries, fuel cells, capacitors, solar cells and water desalination devices is commonly known to wide public. Many research groups worldwide are involved in a race to make those electrochemically based technologies more efficient, economically viable and environmentally friendly. A lot of progress has already been made, for example fully electric or hybrid cars are seen on our streets, experimental fuel cell buses, supercap-driven trams being recharged at each stop, construction cranes recuperating energy by charging supercaps when laying down their platforms before loading, solar panels in private households, installations for the production of fresh water from sea brine or even from human urine at the international space station—all became reality. For electrochemists, being involved in any part of that research and development make them feel good and needed, contributing to the fight against global warming and making impact to our way of life. Last but not least, it is the safest way to get their research funded.

Much less widely understood is the key research in such areas as electrochemical kinetics and electrocatalysis, underpinning most of those mentioned technologies as well as metal treatment, corrosion science, production of gases by electrolysis, electrochemical synthesis, sensors and many other classical ‘mainstream’ branches of applied electrochemistry.

Generally, the societal priorities of the XXI century in its 3rd decade and expectedly in the next few ones, which determine the priority areas for research funding, can be summarized as shown in a diagram:

Advanced and sustainable manufacturing is not depicted here, as implicit. All of these subjects are deeply inter-related.

Of course, nothing can be done without energy. Public health and well-being are affected by climate, by supply of food and water, which in turn affect and are affected by climate. The whole circle and development of the healthy relationship between its components are boosted by information technologies and artificial intelligence, as well as progress in new materials (although both, as we know, as well as other elements of this scheme if misused, may contribute to ‘pollution’).

As described above, in ‘Energy’ the role of electrochemistry is more or less obvious, and it, of course, contributes to “Climate” through the production of clean energy. There are also other inputs to its related tasks, in the first place through capture or consumption of carbon dioxide, production of hydrogen as a fuel for hydrogen economy via electrolysis or photo-electrolysis, ultra-filtration and treatment of exhaust water, and of course the issues related to ‘Energy’ enriched with such new directions, as ‘blue energy’. In ‘Public health’, the input is less straightforward, underpinned by bioelectrochemistry (protein functioning, biological membranes, nerve signal propagation, pace makers) or molecular electrochemistry for sensing. Achievements in ‘Information Technologies and Artificial Intelligence’ are rather used in electrochemistry, than benefiting...
from the latter. In ‘Food et al.’, this could be, e.g. electrochemical synthesis of ammonia for fertilisers avoiding the production of CO₂ as a side product, and of course the mentioned desalination.

Electrochemistry, indeed, seems to be one of the ‘cleanest’ sciences; it is appropriate to cite here Frumkin’s metaphor from his unforgettable Mendeleev lecture in Alma Aata 50 years ago: ‘Aphrodite came out of sea froth (concentrated electrolyte solution), but not from oil’.

As for ‘Materials’, apart from classical electrochemical synthesis and catalysis of various substances, it is a relatively fresh avenue, and in this short essay, I want to focus on it. Moreover, I will dwell on its least advanced but perhaps hottest direction—design of electro tunable metamaterials.

What are metamaterials? These are materials with unusual, odd, disruptive, funky, ‘crazy’, fancy (you can continue yourself the list of such adjectives) properties. Not all of the discovered and designed metamaterials, may appear useful in the end. But some of them do or at least promise to open entirely novel routes for providing new functions. For me, the best symbol for such materials is as shown below, taken from the world of modern architecture.

This picture symbolises for me the pros and cons of this field. Indeed, it demonstrates the unlimited fantasy of the architect. Its shape puzzles spectators, their—‘wow!’ guaranteed. But, (i) it is a few times more expensive than a conventional building of that size; (ii) the judgement on its functionality can be a matter of taste, and, sadly, (iii) such buildings often get tired faster than ‘classical’ buildings (see the spectacular Gehry’s constructions at MIT campus in Cambridge MA; that equally refers to some ‘meta-buildings’ of his passed away ‘sister-in-arms’, brilliant Zaha Hadid).

The same story is with metamaterials. Not all of them will be able to compete with the conventional ones, some may degrade faster; and will they be cheap enough to create and use them? Still, you cannot stop the human imagination, and this area is extremely attractive for its novelty and the brilliance of ideas advancing it.

The term metamaterials is, however, almost a synonym of optical metamaterials, i.e. those that deliver new optical properties. These are, e.g. quantum metamaterials, nonlinear and amplifying metamaterials, materials for transformation optics, invisible cloaks, designer dispersions for slowing light, chiral metamaterials, sensor metamaterials (e.g. based on SERS), microwave frequency selective materials, all kind of complex optical filters and light trappers [1]. All of these are the key components of what nowadays is called photonics [2, 3], the term widely used to embrace the directions of modern optics that use metamaterials or tricky electrodynamics phenomena to produce novel optical effects.

The materials that deliver interesting mechanical properties and their related exploitation are usually called functional materials [4, 5]; although this term may be questioned, at least I do not like it: what is the point for us to work on and develop materials which are not functional? Except, maybe, for the case when they are purely structural, but even then, they have their clear function! Instead of functional, the term smart materials is widely used [6], underlining the idea that they can do something that other, ‘less-smart’ materials cannot (or that they are designed by smart people?).

One class of such materials which is particularly smart are polymer electrolyte electro actuators for robotics [7], reverse actuators such as harvesters of electricity from walking or any other mechanical motion (see e.g. references on piezo [8, 9], capacitive [10–12] and nanotribological [13, 14] devices, and the literature cited there). To avoid ambiguity, I prefer to call all those materials—mechanical metamaterials. Many of them related with effects that are underpinned by electrochemistry. These materials and physical phenomena they are based upon will require a special coverage. Thus, in these notes, I will concentrate on the perspectives of electrochemistry for optical metamaterials.

The science and engineering of the latter have made great progress due to advances in both theory and in micro and nanoengineering. Theoretical analysis facilitated by rapidly developing novel computational tools resulted in suggestions of new, original scenarios and architectures of such materials for their experimental validation. Grand progress in nanotechnology in the last three decades, allowed for such materials to be constructed. But the majority of them had one common problem. Fabricated based on complex nanotechnology, they were static, performing fixed, given function(s). Already at META—2014 congress in Singapore, Nikolay Zheludev in his opening speech said: ‘The time of metamaterials is…over. It is the time of tuneable metamaterials’. He and his group have been already working in this area, either coming up with standard scenarios of controlling light with light [15] or deforming metamaterials mechanically [16]; there were also a number of other groups who had achieved tunability of optical meta-structures by mechanical stress [17–20].
Electrochemistry is, however, in the unique position here to offer new scenarios for electro tunability, and what is specific for electrochemistry—by tiny voltage variation. By this term, we mean voltage control of the properties of various self-assembling optical systems. The origin of this ‘hope’ goes back to old times.

Two dreams:

A dream of a medieval alchemist was to find a magic composition of a solution under which a homunculus will self-assemble in a test tube. A dream of an electrochemist, already since Faraday times, has been to apply and vary the voltage in an electrochemical cell, and whatever self-assembles or disassembles at the electrochemical interface will depend on the electrode potential. Paraphrasing Dostoyevsky, who said that ‘Beauty will save the World’, a maverick electrochemist would say the same about the potentiotstat.

The first of the two dreams has not been realised literally, homunculus-wise, but after many circles has brought us to modern supramolecular chemistry, polymer chemistry, and colloid science, and it neither escaped the attention of electrochemists [21]. The realisation of the second dream is a continuous process.

We will speak below about the use of both approaches to control first of all the properties of plasmonic metamaterials [22]. But before describing that step—few words on plasmonics. This is a branch of solid-state physics and optics exploring different effects of collective electron plasma excitations (plasmons) in the optical response of interfaces and interfacial films, and exploiting them in various optical, sensing and catalysing processes [23]. Plasmonics took off in the 1960s–1970s, with the pioneering works of Otto [24] and Kretschmar [25, 26] who formulated the principles of techniques for characterisation of surface plasmons and experimental measurements of the spectra of these collective excitations, whereas the concepts of bulk plasmons and detections of surface plasmons in thin films were proposed already in 1950s [27, 28].

Plasmonics entered electrochemistry and was actually boosted by electrochemistry, more than 40 years ago with the discovery of surface-enhanced Raman scattering (SERS) in the 1970s. The famous paper, from the University of Southampton, by Fleischman, Hendra, and McQuillan [29], first reported on the effect of a ‘giant’ enhancement of Raman signal from pyridine molecules adsorbed on a rough silver electrode; but they did not unravel the nature of this effect. ‘Purists’ on the other side of the Atlantic used to say that the Southampton team has observed the effect, whereas Van Duyne [30] at North Western University, with his systematic studies, has unravelled what stands behind it. They attribute the discovery of SERS rather to him, and (justly) acknowledge those who contributed to its scientific understanding (see pioneering theoretical works of Schatz, Moskowitz, and others, as reviewed in [31–33]). Indeed, it had slowly become clear that the giant enhancement of Raman signal from molecules adsorbed on rough surfaces is in the first place related to plasmon resonance due to a possibility of direct excitation of surface plasmons by light incident on a rough surface. First estimates, confirmed later with the development of computational methods of solving Maxwell equations, made it clear that local values of electric component of electromagnetic radiation, $E$, with frequencies close to surface plasmon ones, maybe resonance-enhanced near the tips and in crevices of the rough metal surface by a couple of orders of magnitude. Since the intensity of Raman signal from the analyte molecule scales roughly as $E^4$, one can end up with some $10^5$ times enhanced signals. Although, other factors responsible for the enhancement of the Raman signal (the so-called ‘chemical’ or charge transfer enhancement) have been added to the picture beginning with the paper by Albrecht and Creighton [34], and later dwelled upon by Otto [35], the plasmon-resonance mediated enhancement remains central for SERS. In 1980s, interest to plasmons in electrochemistry beyond SERS coincided with development of electromodulation spectroscopy [36, 37].

The boom in nanotechnology and nanoscience of this century gave rise to the second birth of plasmonics, more generally photonics (the science that including plasmonics as a particular case). Indeed, conventional materials derive their optical properties from atoms and molecules. Nano- and micro-technology could create structures and architectures from larger blocks than simple atoms, e.g. nanoparticles or other components of nanostructures, thus designing larger scale ‘atoms’ to access new functionalities, capitalising on plasma excitations in these complicated structures or more complex effects [38].

Electrochemistry very often rapidly followed discoveries in physics, and there are many examples of it in the history of science. In recent times, some of those were building the in situ STM with atomic resolution which could picture atomistic structure of electrode/electrolyte interface [39, 40], just 2 years after Binning and Rohrer reported their vacuum STM [41, 42], or implementation in a due time of just discovered carbon nanotubes or graphene for building electrode for electrochemical supercapacitors [43]. Electrochemistry had not exploited
the achievement of modern plasmonics in full yet but it can do it, by using nano-engineered electrodes [44–46] or playing with self-assembling nanostructures, and this is what is happening now.

The simplest of such self-assembling nanostructures were arrays of metallic nanoparticles (NPs) at ITIES. The road to these works was opened by pioneering papers of Schiffrin, Girault, Samec and their groups [47–51], who have studied the formation of such layers at electrochemical liquid/liquid interfaces (LLI) [52, 53], i.e. interfaces of immiscible electrolytic solutions (ITIES), including their optical and/or electrical (capacitance) characterisation. Indeed, at LLI properly functionalised NPs, usually dissolved in the aqueous phase, tend to spontaneously adsorb onto the interface to block the unfavourable contact between water and oil. Voltage control over electrosorption of NPs was demonstrated (they were, however, too small, of 1.5 nm in diameter, for any contribution to modification of the linear optical signal) already in Ref. [51] through cyclic voltammetry and capacitance measurements. Second-harmonic generation detecting the presence of NP was first demonstrated in Ref. [50], but again for smallish NPs, 6 nm in diameter.

In 2010, Flatte, Urbakh and I proposed and explored an idea how to influence optical properties of ITIES through voltage controlled assembly disassembly of NP arrays, by tuning concentration of electrolyte and potential drop across the interface [22]. With all the preliminary character of the estimates presented in that Feature Article and some minor inaccuracies, this theoretical paper has set for us a new direction of research: electrotuneable electrochemical plasmonics.

Indeed, the quasi-2d NP arrays of plasmonic MPs (gold, silver, titanium nitride, core-shell composite NPs) adsorbed at the interface of two optically transparent media reflect light in a broad visible range with a maximum reflection centred about the frequency of the localised plasmon modes in nanoparticles. That value depends on the size and composition of NPs; for 20 nm diameter AuNP, it is close to the wavelength of 560–580 nm. The reflection signal depends on the material and size of NPs: the reflection signal is weaker the smaller the size and the stronger is light absorption in the material. Too large sizes of NPs, approaching the wavelength of light, would neither be good for reflection, as this would give rise to diffuse scattering of light. The size between 16 and 40 nm in diameter was found to be most suitable for enhanced reflection. Apart from the size, another critical parameter is the average distance between NPs in the arrays, i.e. the array density. It affects the reflection spectra in two ways: (i) the reflection at maximum and across its whole spectrum increases, the closer to each other are the NPs in the array; (ii) the wavelength of maximum is shifted to the red with increasing the array density.

Thus, the following picture and manipulation scheme was envisaged and later approved by a series of systematic experimental studies:

1. NPs are functionalized with ligands, such as mercaptanoic acid or citrates. Having acidic terminal groups, they dissociate in water, retaining negative charges (other ligands may be designed to deliver a positive charge, but the sign of the charge is not principle rather its value, as long as it is the same for all NPs). The charge thus associated with NPs will protect them against agglomeration that may be caused by van der Waals forces (a standard measure for stabilisation of colloids). A 20 nm NP functionalized at neutral pH by mercaptanoic ligands can bear approximately 900 elementary charges.

2. The electrostatic repulsion of charged NPs can be controlled through the variation of electrolyte concentration that affects Debye screening and through the degree of ionisation of ligands, which can be altered by the variation of pH. More concentrated electrolytes will strengthen screen the electrostatic repulsion in the solution. More acidic pH will reduce the degree of dissociation of acidic terminal groups of functionalised ligands and decrease the overall charge associated with NPs. Both factors will reduce the repulsion of NPs in the array adsorbed at the interface and would favour denser arrays. But obviously, this cannot be overdone—too short screening length in the solutions and too small charges of NPs would lead to their agglomeration in the bulk.

3. The control of inter-NP spacing through electrolyte concentration and pH is, thus, limited. Although each individual NP would wish to settle at the LLI, as we have designed the system in such a way that NPs repel each other strong enough not to fuse with each other in the bulk, the NPs will keep ‘social distance’ between each other when adsorbed at the interface. Thus, the strategy is to reduce repulsion as much as possible, but not that much that they would start agglomerating in the bulk (and then at the interface). Theory may try to estimate the limiting values for electrolyte concentration and pH, but they are best defined for each system experimentally as has been repeatedly done. Indeed, agglomeration of NPs in the bulk can easily be detected through the characteristic features in the light extinction spectra of the solution bulk. All in all, each such system will have some minimal distance for interparticle separation in an adsorbed array. Since nanoparticles were designed to repel each other, for spherical NPs that array will be, on average, hexagonal.

4. If we want NPs to form even denser arrays, a natural idea would be to apply voltage across the ITIES to push the charged NPs to the interface. If the ligands are negatively charged, this will take place polarising aqueous phase negatively. When positively charged ligands are used, this
will be other way around. Polarisating the cell correspondingly will increase the driving force for NPs to settle at the interface. The capillary well and solvation does not let them go into the oil phase completely, so that they settle at the ITIES, piercing it, although when being pushed by electric field they can be somewhat shifted to the oil side of the interface. This would make the potential well for each individual NP deeper, and thereby increase their ‘desire to settle at the interface’, so that NPs will tolerate shorter social distancing between each other.

These ideas were touched in different depths in the 2010 paper, but all their details and complications to overcome were made clear only after a series of follow up theoretical works and brand new experimental investigations, performed for both for LLI and solid/liquid interfaces (SLI).

The theoretical works focussed on the development of a more accurate theory of optical signals from NP arrays settling at optically transparent as well as reflective interfaces, at a later stage tested by full-wave simulations of optical signals. The theory was reduced to precision by Debabrata Sikdar leading to excellent correspondence between theory and full-wave simulations [54, 55]. Systematic realisation of these principles took place at Imperial College, in the laboratory of Joshua Edel—led by him, Anthony Kucernak, and myself, with a crucial work of research associates—Eleanora Velleman, Yunuen Montelongo, and Ye Ma, and few other students and co-workers. This has not only built an experimental basis of electrotuneable electrochemical plasmonics, but also allowed to systematically test the theory on various optical experimental data. I will mention just a few milestones of that research programme.

That period of work was actually preceded by a series of earlier works, performed before achieving electrovariability experimentally, and reviewed in Ref. [56] It has started with experimental investigation of the factors influencing adsorption of NPs onto LLI (electrolyte concentration and the aqueous solution pH) [57] followed by computational analysis of the optical properties of NPs at the interface [58]. The ‘nonelectrotunable’ period culminated in Ref. [59], which has demonstrated an unprecedented level of detection of Raman signals from analyte molecules adsorbed between 40-nm-sized NPs adsorbed at LLI. Again, in that work, control of the density of the adsorbed NP array was only via variation of electrolyte concentration and pH. But it has shown how ‘hot’ can be the spots of enhancement of electromagnetic radiation (near and in between the NPs) probing the vibrational modes of the analyte molecules through their ‘Raman fingerprints’. Although that work did not involve yet an electrochemical setup, and thus no voltage could be applied there to stabilise and densify the adsorbed NP arrays, it was unambiguous evidence of the formation of relatively dense NP arrays at an LLI.

There was then a seminal paper published by the group led by Mark Schlossman at the University of Illinois in Chicago. They have performed very fine investigations of assembly of small (2 nm) NPs functionalized by positively charged ligands [60]. Their localization at the water/DCE ITIES was studied by Grazing Incidence Small Angle Scattering (GISAXS), controlled by the voltage applied across the interface. From Bragg diffraction peaks, they could see a hexagonal arrangement of NPs at the interface, with the decrease of the lattice constant, when NPs are get pushed towards oil by electric field at the interface. Through X-ray reflectivity, they could also assess how deep the NPs can be pushed towards oil, when polarising water positively with respect to oil. The molecular dynamic simulations performed in the same work indicated that when NPs are pressed into the oil side of the interface, their charge seems to be strongly compensate by the counterions binding to the charged groups of the functionalizing ligands.

That work has inspired Joshua Edel to initiate and lead a study, crucial for the whole following work of the Imperial team in this area. The results of this study were published in Ref. [61], which referred to a detailed structural and optical characterisation of NP arrays at the LLI. It was the measurements of spectra of light reflection from NP arrays spontaneously adsorbed at LLI, the density of which was controlled by variation of electrolyte concentration either in aqueous or in organic phase, with simultaneous characterisation of the array structure by X-ray diffraction. The experiments were performed at the Dimond Synchrotron Light Source at Harwell. Importantly, the X-ray diffraction and light reflection measurements were performed at the same time in the same cell.

Again, it was not an electrochemical cell, and the light reflection was not strong, because too dense arrays did not form without assistance of applied voltage, and furthermore the used gold NPs were of intermediate size, 12.8 nm. But the reflection spectra, performed at normal incidence of light to the LLI, were still well detectable. The X-ray diffraction data has shown almost ideal hexagonal structure of the arrays and the expected decrease of the inter-NP distance with the increase of electrolyte concentration. That distance was a key parameter of the theory of light reflection from the array. Thus, for each electrolyte concentration, and thereby inter-NP separation in the array, determined experimentally, it was possible to calculate practically without adjustable parameters the whole reflection spectra and compare them with the experimentally measured ones. The results have shown excellent correspondence of theory [54] and experiments, thus proving that the basic ideas underpinning the whole project were correct.

Let us reiterate that the size of NPs is crucial for the mirror effect. The smaller the NP, the smaller the reflectivity at maximum, and the intensity goes down overall spectrum correspondingly. Girault’s group [62] has studied systematically the size effect, controlling the population of LLI with NPs of
different size by direct addition of NPs to the interface (the method not excluding formation of multilayers of NPs). It was found, not unexpectedly, that already above 60 nm in diameter of NPs, the scattering effects seemingly start to interfere with reflection, as reflection goes down with further increase of the size of NPs (see their Fig. 4). There is thus an optimal NP-size for an ‘ideal’ mirror, as the wave-length must be much larger than the diameter of NPs, to see it as a smooth surface. At the same time, if the NPs are too small, its ‘plasmonic mass’ will also be too small, and the presence of such NPs at the interface will be invisible in linear optics. Interestingly, in this work, the population of the interface was controlled also by measuring the electronic conductivity across the interface (electrons presumably tunnelling along the ligands between the NPs along their quasi-2d percolation network). So, in Ref. [62], they also plotted the reflection signal as a function of estimated NP-population at the interface.

As mentioned, the experiments at Diamond did not and could not yet involve any elements of electrovariableity. It thus remained to determine the latter in an ITIES electrochemical cell. The corresponding opto-electrochemical setup has been built under supervision of Edel and Kucernak. The experiments where performed with 16 nm AuNPs adsorbing and desorbing from aqueous/1.2-DCE interface at optimised pH of the aqueous phase and concentrations of Na\(^+\)Cl\(^-\) electrolyte in it, and TBA\(^+\) TPB\(^-\) organic phase. The experimentally measured reflection spectra at normal incidence, obtained for each given voltage across the ITIES, where treated with the theory by fitting first the wavelength of reflection maxima and their height, thereby extracting the information on the average inter-NP separation (as there were no independent X-ray data available for it), and then calculating the whole spectra. It was full success again: the calculated spectra where practically exactly like the measured ones. This was the first realisation and characterisation of electrovariable mirror at an electrochemical interface [63].

Joshua Edel has shot a video, showing how this mirror looks like, and how it operates: https://www.nature.com/nmat/journal/vaop/ncurrent/full/nmat4969.html The video has been viewed on the web more than 24,000 times, and discussed on many websites, see, e.g. https://www.photonics.com/Articles/Tunable_Nanoparticle_Layer_Switches_Between/a62514. It shows how a tiny, just 0.5 V, variation of voltage can change the interface from a ‘mirror’ to ‘window’ mode. It displays either the reflection of a coin positioned above the interface onto which the light is shine when AuNp form a dense array. When the voltage is switched to push NPs away from the interfaces to the bulk of the aqueous phase, thereby making the interface transparent, you see a £10 banknote underneath.

The video, as spectacular as it is, had, however, to be dramatically sped up, because those experiments deliberately avoided dielectrophoresis [64, 65] that could otherwise drive charged NPs to the interface through the aqueous bulk. The static applied voltage cannot do it. In the absence of electrical current across the interface electric fields in the bulk of each phase are screened and both aqueous and oil bulks stay electroneutral. The electric field is concentrated only within the electrical double layers on either side of the interface and at the electrodes. Thus, when the aqueous phase was polarised more negatively, increasing thereby the potential wells that trap NPs at the interface, NPs in the bulk could not know anything about it. They would feel the difference, only when they get close to the LLI. But they could reach the interface to get trapped there only via random diffusion from the bulk, which is a slow process. Compare yourself reaching the point of interest in a historical town with a guide or by random walk.

A simple estimate shows that the diffusion time for NPs to reach the interface to form a monolayer is inversely proportional to the square of the bulk concentration, \(c\), of NPs [63]. Increasing their concentration through the increase of the number of NPs would not, however, be a good idea, because the aqueous solution in which such NPs will be dissolved would get heavily coloured—it will look like Bordeaux wine. But one can reduce the thickness of the aqueous phase, maintaining a minimal number of NPs (just enough to cover the monolayer at the interface). This has been tested experimentally, and the \(1/c^2\) law for the switching time was approved [63]. This opens a door to building electrochemical microcells, without invoking dielectrophoresis (involvement of the latter is possible, but it would have made practical implementations of such cells more complicated). Estimates show that properly designed microcells could deliver millisecond switching time between window and mirror modes.

With a dream to use such optical switches in various applications, and particularly in the building industry—switching windows to mirrors in skyscrapers, saving energy on air conditioning when offices are not occupied, as first spelled out in Ref. [22], which may require vertical orientation of the interface—ITIES do not look too practical. Electrochemical cells with vertical ITIES have been created from the early days of ITIES in laboratory (see papers cited here), but no one tried to upscale them. So our thought was to create similar optical switches on the basis of adsorbing/desorbing NPs from electrolytes onto/from transparent electrodes, such as ITO, ZnO, or graphene on a glass-substrate. Corresponding calculations have been performed showing very promising results. But this idea has not yet been experimentally realised. In the meantime, a more conventional (for electrochemistry) prototype of such system was considered.

Indeed, a purely theoretical analysis, confirmed also by full wave simulations, has shown another interesting effect, when NPs get electrosorbed or electrodeposited on a mirror metal electrode [55]. Without NPs adsorbed on the surface of such electrodes, it should provide classical wide band reflection
spectrum of that bare surface. But as predicted by the theory, with a dense monolayer of plasmonic NPs of sufficiently large size adsorbed on it, another effect should take place: a broadband quenching of reflection. The wavelength of the reflection dip is centred again at the wavelength of plasmon resonance of individual NPs, moving slightly to the red with densification of the array.

This effect was not broadly known. All aspects of it were clearly described in Ref. [55], but it was also predicted within an earlier version of the effective medium theory [66] (substantially improved in Ref. [55]), and by using another theoretical method by Truong and de Dormale [67]. In another configuration (adsorption of AuNP on thin gold film that allowed studying light reflection from the adsorbed NP-array side as well as shining light onto the other side of the film, thus probing plasmon resonances by evanescent wave excitation), an effect of similar nature was experimentally studied by the group of David Smith [68]. Moreover, in an overlooked early paper of the Liverpool group led by David Schiffrin signatures of a similar effect have been seemingly observed in the electroreflectance signal from the bi-layer of 5nm AuNps on functionalized gold electrode [69].

Somehow, predictions of the theory [55] have been systematically tested experimentally by the Imperial team [70]. Silver electrode, protected by nano-thin layer of titanium nitride (thin enough not to affect optical properties of the silver surface) in aqueous solution of AuNps functionalized by negatively charged ligands, was shown to experience the described quenching of reflection with electrosorption of NPs, returning the optical reflectivity of bare silver in water with their electrosorption. Within the electrochemical window, adsorption of negatively charged NPs was stimulated by continuous variation of electrode potential from PZC to +0.6 V relative to the latter: each electrode potential resulted in different density of the NP array, and correspondingly, the more positive the electrode potential resulted in denser arrays of adsorbed NPs and the stronger reflection quenching. Within the acquired potential window, the array was never denser than the one corresponding to 5 nm inter-NP surface-to-surface separation. But even then, as was found in a later paper [71], for large NPs (40 nm in diameter), almost 100% quenching of reflectance was observed at the wavelength of reflection minimum. All the experimentally obtained spectra were fitted to the theory, through the wavelength of the reflection dip and its depth, with one fitting parameter—the average distance between NPs. Thus, the average inter-NP separation was retrieved for each applied electrode potential. Then, again, using the values of this key parameter of the theory, the whole spectra were calculated, reproducing amazingly well the experimental curves.

Encouraged by this success, the team went back to NP-array-based SERS measurements, now not on LLI (work in progress), but on a metal substrate. Physically, the space between electrosorbed NPs (still protected against direct contact with the metal substrate by their functionalizing ligands) and the electrode deliver hot spots for electromagnetic radiation, even when the NPs are far apart. But when NPs come closer to each other, the additional hot spots emerge in the space between NPs, as well as those under the NPs. Thus, the Raman signals from any analytes that could get into the mentioned regions will get enhanced. Those effects have been demonstrated in Ref. [71]. Again, varying electrode potential altered the average distance between the NPs with their negatively charged ligands; more positive electrode polarisation created a denser array of NPs, and amplified the SERS signal.

The Raman signal from the analytes was increasing proportionally to the increase of the number of NPs on the surface but only when NPs were on average still far from each other. Further densification of NP array was making this dependence nonlinear, thus demonstrating clear emergence of hot spots not only underneath NPs but also in between NPs, when they come closer to each other, and, plasmon resonance wise, start ‘talking with each other’! In the same work [71], measurements of the Raman signals were paralleled by simple reflectivity measurements. The latter allowed to retrieve the average distances between NPs, control the proximity of the measured and calculated reflections spectra (with full success, again), and with thus evaluated distances between NPs in the monolayer to calculate the enhancement of Raman intensity.

Note that the idea of building metasurface for enhanced Raman signal by designed arrays of plasmonic nano-objects has long history [read, e.g. 72–74], although the funky term ‘metasurfaces’ was not used then. What was shown in Ref. 71 is that one can manipulate the structure of such arrays, ‘electroadsorbing/electrodesorbing’ them onto/from solid electrodes, at our will and in real time, and how it thereby affects the SERS signal, demonstrating a crucial role of electromagnetic enhancement mechanism [75].

Correspondence between theory and experiments was perfect.

Few other configurations have been considered, based on similar principles.

This is, e.g. a theory of functioning of Fabry-Perot cell (cavity) that would ‘trap’ light between two semi-transparent electrode plates, and filter the transmission of light through the cell, by controlling the density of NP arrays on the plates through NP electrosorbtion/desorption from/to electrolyte solution filling the cavity [76, 77]. It is an interesting alternative to the standard mechanical adjustment of the cavity length, as it shows opportunities of sharp light transmission filtering by tiny voltage variation.

Another example goes third dimension: electro-feeding with NPs of the void space between transparent electronically conducting ZnO nano-columns, subject to the applied voltage [78]. The electrodeposition of metals into well-defined porous structures and dimensions is well-known, extending as far as to growing ultrathin nanowires in metal-organic frameworks.
(MOFs) that can function as versatile templates for the growth of metallic nanostructures with precisely controlled shapes and sizes (see e.g. [79, 80]). We know how to electrosorb large ionic liquid ions into MOF pores [81]. But, to my knowledge, building columns of individual NPs by physical electrodeposition in pores of large size has not yet been tried.

Interesting opportunity could be offered by anchored systems: changing orientation with respect to the electrode of the metallic (plasmonic) ‘flat’ nano-cuboids linked on one narrow edge to a transparent electrode, with variation of electrode polarisation. The calculation shows a remarkable effect of the reorientation of cuboids with respect to the electrode on the overall optical response of the interface [82], with fast dynamics, as not depending on translational diffusion but only reorientation of cuboids.

None of these three last examples have been yet experimentally tested, so they remain hypothetic systems. But all the previous experience with their sister systems suggests that they would work as predicted by the theory.

It was furthermore shown that interfacial optical properties of NP arrays can also be controlled not through applied electrode potentials, but temperature variation in a system that keeps the array together by thermo-auxetic ligands. Such thermally controlled optical switch was proposed by Joshua Edel. It was based on the self-assembly of poly(N-isopropyl-acrylamide)-functionalized gold nanoparticles on a planar macroscale gold substrate, as been built and explored by Ye Ma, Deb Sikdar, et al in Ref. [83] Shrinking of these ligands with temperature increase and expansion with temperature decrease affects inter-NPs separation; this effect can be calibrated theoretically and experimentally, in a similar way as in electrosorption-controlled arrays. Again, this particular system was shown also to operate in harmony with the theory [83].

One should also mention a series of works by the Dryfe’s group in Manchester related to different aspects of localisation of NPs at ITIES, including voltage control over the density of adsorbed NP arrays. For instance, they have published a paper on electrrotuning SERS at the ITIES by voltage-controlled population of silver NPs at the interface [84]. I will not go into any further details of the works of this group, referring the reader to a comprehensive review article by Booth and Dryfe [85].

There were a number of works exploring electrocatalytic properties of self-assembled layers of NPs at ITIES, such as reported, e.g. in Ref. [86–88]. This subject is very interesting itself, but it lies beyond the central theme of this article; moreover, intriguing from the fundamental point of view, it is not clear yet how practical such systems could be. Indeed, liquid/liquid interfaces are flat, but not volume-filling as required for industrial applications of electrocatalysis. Creating, however, decorated novel electrode materials by controlled self-assembly at such interfaces is an interesting direction to mention [89]. Preparation of solid-state nanomaterials using electrodeposition of larger size colloidal particles onto templated electrodes in ordinary electrochemical cells is a separate direction for electrochemical nanotechnology [90].

All in all, the thus far obtained results have demonstrated that the physical picture of the effects of electrochemical plasmonics has been properly understood, and it can be laid into basis of the corresponding electrochemically controlled optical devices, to be built resting on the obtained fundamental knowledge. As mentioned, this will require miniaturisation of electrochemical cells, moving to micro-cells, and micro-compartmentalised cells. This is a long road. It will take time, efforts and money, to demonstrate the proof of concept of these novel devices. But the general direction has been made clear.

Now, how about electrochemical photonics, i.e. the effects beyond simple plasmonic resonances, such as reflectivity, SERS, and alike? I have listed the main directions of photonics above, many of which are based on much more sophisticated theories and experimental setups. But what electrochemistry could do is to make them electrotunable. This will refer preferably to super structures with the characteristic dimensions larger than wavelength of light.

A simple idea, proposed to us by one of the photonics champions—N. Zheludev, is to lay on the surface of an electrode (transparent or reflecting one, subject to the task of interest), an optically and electrochemically inert mask with any figures of interest cut out of the mask, thereby opening those sections to electrosorption of NPs. The population of those sections with NPs can then be controlled by polarising the electrode, leading to the variation of the corresponding effects of the optical properties of those sections. The underlying electrode could further be compartmentalised, so you could control the properties of each section independently. Options here are unlimited, subject to the researchers’ imagination, ideally working in cooperation with cutting-edge gurus from the corresponding branches of photonics.

Electrochemistry is an amazing interdisciplinary science that had already ‘donated its blood’ to many branches of sciences and engineering, from energy storage to biology and medicine. I have outlined above a new, emerging avenue for it. But let’s be honest, solid-state scientists and engineers (and photonics is based on solid-state physics and electronics) are afraid of anything that is ‘wet’. Would ‘wet photonics’ ever take off? For fundamental investigations—why not, but for practical applications?

One way (absolutely not tried and not tested yet) is to use ionic liquids as electrolytes, perhaps slightly doped with organic solvent to reduce their viscosity (the so called “solvent-in-salt” systems). They are not volatile, behave almost like gels, even more than the aqueous electrolyte in the Dead Sea, and most of them are optically transparent. The question is whether they could let NPs move easily and fast away from or to the surface, and generally form dense electrosorbed monolayers at the interface. If they do, when implemented in micro-engineered electrochemical cells, the latter would not
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