Tunable Surface Structuration of Silicon by Metal Assisted Chemical Etching with Pt Nanoparticles under Electrochemical Bias

Encarnación Torralba,†,‡ Sylvain Le Gall,†,‡ Raphaël Lachaume,‡ Vincent Magnin,§ Joseph Harari,§ Mathieu Hallwax,§ Jean-Pierre Vilcot,§ Christine Cachet-Vivier,† and Stéphane Bastide*†

†Institut de Chimie et des Matériaux Paris-Est, CNRS, Université Paris-Est, 2-8 rue Henri Dunant, 94320 Thiais, France
‡Group of electrical engineering – Paris, UMR CNRS 8507, Centrale Supélec, Univ. Paris-Sud, Université Paris-Saclay, Sorbonne Universités, UPMC Université Paris 06, 3 & 11 rue Joliot-Curie, Plateau de Moulon, 91192 Gif-sur-Yvette CEDEX, France
§Institut d’Électronique, de Microélectronique et de Nanotechnologie, UMR CNRS 8520, Université de Lille 1 - Sciences et Technologies, Avenue Henri Poincaré, CS 60069, 59652 Villeneuve d’Ascq cedex, France

Supporting Information

ABSTRACT: An in-depth study of metal assisted chemical etching (MACE) of p-type c-Si in HF/H₂O₂ aqueous solutions using Pt nanoparticles as catalysts is presented. Combination of cyclic voltammetry, open circuit measurements, chronoamperometry, impedance spectroscopy, and 2D band bending modeling of the metal/semiconductor/electrolyte interfaces at the nanoscale and under different etching conditions allows gaining physical insights into this system. Additionally, in an attempt to mimic the etching conditions, the modeling has been performed with a positively biased nanoparticle buried in the Si substrate. Following these findings, the application of an external polarization during etching is introduced as a novel efficient approach for achieving straightforward control of the pore morphology by acting upon the band bending at the Si/electrolyte junction. In this way, nanostructures ranging from straight mesopores to cone-shaped macropores are obtained as the Si sample is biased from negative to positive potentials. Remarkably, macroscopic cone-shaped pores in the 1–5 μm size range with a high aspect ratio (L/W: 1–1.6) are obtained by this method. This morphology leads to a reduction of the surface reflectance below 5% over the entire VIS-NIR domain, which outperforms macrostructures made by state of the art texturization techniques for Si solar cells.

KEYWORDS: metal assisted chemical etching, c-Si, platinum, cyclic voltammetry, impedance spectroscopy, 2D band bending modeling, Si texturization, cone-shaped macropores

INTRODUCTION

During the past decade, metal assisted chemical etching (MACE) of Si has attracted considerable attention as an efficient method to produce Si nanostructures with high aspect ratio. Nanowires, nanogratings, and nanopores have been synthesized by MACE and their potential applications have been tested in fields ranging from solar energy conversion to chemical and biological sensing.

Noble metals, including Ag, Au, Pt, Pd, and Rh are known to be effective catalysts for MACE. Using single nanoparticles (NPs), etching with Ag results in the formation of straight mesopores with smooth walls whose diameter matches the particle size. Ag is therefore the metal of choice for highly localized etching and formation of well-defined nanostructures (e.g., Si nanowires). On the contrary, Pt leads to the formation of cone-shaped pores with walls fully covered by porous Si. The behavior of Au seems to be intermediate between that of Ag and Pt. The etch rate increases in the following order: Ag < Au < Pt. Although well-known, these differences have started to be addressed only recently. Yae has established that the etch rates depend on the difference in catalytic activities of the metal for O₂ reduction in the HF/O₂ system. Kolasinki has recently brought to attention the importance of the Si/metal Schottky junction properties on the (de)localized character of etching. Depending on the metal work function, the junction can be rectifying (Ag) or not (Au, Pt) and thus can modify hole (h⁺) injection into Si.

Despite the fact that Pt has been associated with MACE of Si since the beginning, the mechanisms for the formation of cone-shaped pores by Pt particles and the dependence of their morphology on the etching conditions have not been studied in great detail so far (contrarily to Ag). A better understanding of Pt-based MACE, and its peculiarities compared to other metal etching systems, is therefore needed. To the best of our knowledge, MACE was first reported by Zhang et al. in 1993. In their work, n-type Si samples in contact with an Hg pool were immersed in HF solutions with dissolved O₂ and under illumination. Strong etching occurred when a Pt wire was
placed in contact with the Hg pool and the electrolyte. At the end of the 1990s, electroleless Si etching obtained by Pt deposition in HF/PtIV salts leading to porous Si formation was reported by Gorostiza et al. and by Li and Bohn who coined the term MACE with their work on porous Si formation with platinized Si surfaces. At the same time, galvanic etching of Si with Pt in HF/H2O2 was proposed by the group of Kelly as an electroleless technique to form porous Si on large scale samples for application in microelectronics.

The discovery that metal NPs actually dig pores into Si substrates was eventually made by Tsujino and Matsumura with Ag2 and Pt NPs and by Peng et al. with Ag for the elaboration of Si nanowire networks. In the case of Pt, etching in HF/H2O2 with NPs (50 nm in diameter) resulted in the formation of pores consisting of an empty core of the same diameter as the Pt NP and a large surrounding porous Si region exhibiting a cone-shaped morphology. Using Pt beads of micrometer size and a longer etching time, cone-shaped macropores with a thin layer of porous Si on the walls were formed. To explain these results and the difference with the straight and smooth mesopores obtained by MACE with Ag NPs, the authors suggested that an easy transport of h+ across the Pt/Si interface together with a strong catalytic activity of Pt was leading to porous Si formation at the pore walls via diffusion of injected h+. The cone-shaped morphology was attributed to the fact that the porous Si layer formed at the walls becomes thicker as a function of time, i.e., with a thickness gradient from the top to the tip of the pore. In more recent reports, injection and diffusion of h+ from the metal catalysts have been considered to be key parameters to explain the etched structure morphologies obtained by MACE.

In this work, MACE of p-type c-Si using Pt NPs as catalyst is investigated both experimentally and theoretically. Special emphasis is paid to the study and interpretation of the electrochemical processes governing the mechanisms of pore formation and their dependence on the etching conditions. Combination of electrochemical characterization (cyclic voltammetry, open circuit measurements, chronoamperometry, and impedance spectroscopy) and 2D band bending modeling of the metal/semiconductor/electrolyte interfaces demonstrates that the particular features of Pt etching can be explained on the basis of the ohmic nature of the Pt/Si contact which induces an anodic polarization of the Si bulk. To the best of our knowledge, this is the first time in the field of MACE that a rigorous modeling of the band modulation under etching conditions (electrostatically biased and buried NP) is presented.

In line with this mechanism, the application of an external potential during MACE is investigated as a simple way to achieve a straightforward control of the pore morphology, by modifying the band bending at the Si/electrolyte junction. A similar approach has been reported by Chourou et al. where MACE in HF/H2O2 with Ag, Pd, and Pt has been studied on lightly doped p-type Si under small anodic polarization. In the case of Pt, straight macropores were formed under these etching conditions. We show here that with p-type Si and at higher anodic potential, high aspect ratio cone-shaped pores can be obtained, with potential applications in photovoltaics due to the very low reflectance of the resulting surface morphology (R ~ 3%).

**EXPERIMENTAL SECTION**

**Reagents.** Polished, single crystalline (100) p-type (boron) and n-type (phosphorus) Si wafers, 350–450 μm thick, with a resistivity of 1–3 Ωcm were used. 30% H2O2, 96–97% H2SO4, 40% HF, and 65% HNO3 reagents of analytical grade (VWR chemicals) were employed as received for cleaning and etching Si and for porous Si dissolution. H2PtCl6·6 H2O (99.9%, Sigma-Aldrich) was used for Si electroleless platinization. Dilution and cleaning were performed with 18.2 MΩcm Ultrapure water (Millipore).

**Electroleless Metallization of Si Samples.** Si wafers (p or n type) were cleaved into 1 × 1 cm² pieces and cleaned for 20 min in piranha solution (3:1 v/v concentrated H2SO4/30% H2O2) to eliminate organic residues. After copious rinsing with ultrapure water, the samples were immersed for 5 min in HF 1 M to remove surface oxides and to hydrogenate the Si surface. Pt NPs were then deposited on the cleaned Si substrate by immersion in an aqueous solution of 1 mM H2PtCl6 and 90 mM HF for 30 to 300 s.

**Metal Assisted Etching.** Etching was performed just after platinization in an aqueous solution of HF and H2O2. In all cases, a ratio ρ = [HF]/([HF]+[H2O2]) was set to 0.75.

**Electrochemical Characterization.** All the electrochemical experiments were performed in a homemade O-ring three-electrode cell. p or n type Si (bare or platinized) were used as working electrode, a KCl saturated calomel electrode (SCE), protected from the HF solution by a KCl saturated agar–agar bridge, was used as reference and a Pt wire as counter electrode.

The geometrical area of Si exposed to the solution in the electrochemical cell was 0.38 cm². The electrical contact at the back side of the Si sample was made by rubbing InGa alloy onto it. Owing to the HF concentrations used (typically 0.5 M or above) no supporting electrolyte was added. Approximately 20 mL of solution were used for the electrochemical experiments. All potentials are referred to SCE.

For the open circuit potential (OCP) determinations, the initial solution of 0.5 M HF was degassed for 10 min under Ar prior to the experiment to remove dissolved oxygen. Air atmosphere was maintained throughout the experiment. Given the negligible oxidizing effect of O2 in the presence of H2O2, the working solutions were not degassed for the MACE experiments.

For impedance spectroscopy measurements, the amplitude of the ac signal was 10 mV rms and several frequencies ranging from 10 to 100 kHz were used. The differential capacity C was determined from the imaginary part of the complex impedance (C = −1/(2πf Z′′)) in the frequency range for which the system behaves primarily as a combination of capacitive circuit elements (phase angle greater than 80°). All impedance measurements were carried out in the dark.

**Instrumentation.** Cyclic voltammetry, chronoamperometry, and impedance spectroscopy were performed with a PGSTAT 20 potentiostat/galvanostat Metrohm Autolab, equipped with GPES and FRA software. Scanning electron microscopy (SEM) images were obtained with a Merlin FEG microscope from Zeiss. The optical reflectivity measurements were performed in the 350–1500 nm range using a Hitachi UV–vis-NIR 4001 spectrophotometer fitted with an integrating sphere.

**RESULTS AND DISCUSSION**

**Platinization of Si Samples.** Electroleless metal deposition on Si is a well-known procedure based on a localized galvanic
displacement reaction. In the case of Pt, the reduction of a metal complex, such as PtCl$_6^{2-}$ (from H$_2$PtCl$_6$), coupled to the simultaneous dissolution (in the divalent state) of nearby Si atoms, leads to metal deposition:\(^1\)

\[
\text{PtCl}_6^{2-} + 4e^- \rightarrow \text{Pt} + 6\text{Cl}^-; \quad E^0 = 0.74\text{V/SHE}
\]  

(1)

at cathodic sites:\(^2\)

\[
\text{Si} + 4\text{HF} \rightarrow \text{SiF}_4^{2-} + 2\text{HF} + \text{H}_2 + 2e^-; \quad E^0 = -1.20\text{V/SHE}
\]  

(2)

In order to find the conditions leading to the deposition of well-defined Pt NPs, different metallization times were tested. Figure 1 shows SEM plane views of p-type (100) Si samples with Pt NPs deposited by immersion in a metallization bath containing 1 mM of H$_2$PtCl$_6$ and 90 mM of HF, for three different times (cf. Experimental Section for details).

![Figure 1](image1.png)

Figure 1. SEM images of platinized p-type (100) Si surfaces obtained by immersion in an aqueous solution of 1 mM H$_2$PtCl$_6$ and 90 mM HF for 30 s (a), 120 s (b), and 300 s (c).

MACE and Pore Morphology. MACE uses noble metals to induce local oxidation and reduction reactions under open circuit. The metal deposited on the Si surface serves as a local cathode to catalyze the reduction of oxidants (usually H$_2$O$_2$). The produced h$^+$ are then injected into the Si valence band, which initiates the oxidation of surface atoms that eventually become soluble as SiF$_6^{2-}$ complexes in the acidic HF solution. This results in the removal of Si without net consumption of the metal.\(^24\)

In the particular case of Pt and by analogy with the mechanism proposed for Ag,\(^22\) the overall reaction of MACE in a solution of H$_2$O$_2$ and HF can be written as

\[
\text{Si} + 6\text{HF} + \frac{n}{2} \text{H}_2\text{O}_2 \rightarrow \text{H}_2\text{SiF}_6 + n\text{H}_2\text{O} + \left[\frac{4-n}{2}\right]\text{H}_2
\]

(3)

where $n$ varies from 2 to 4 as the Si dissolution regime evolves from porous Si formation to electropolishing.\(^25\)

Cross sectional SEM images of typical cone-shaped macro-pores obtained by MACE of p-type c-Si with Pt NPs are given in Figure 2. To obtain these structures, the platinized p-type Si samples were immersed for 10 min in a solution of 3.63 M HF and 1.2 M H$_2$O$_2$ ($\rho = 0.75$).

![Figure 2](image2.png)

Figure 2. SEM images in cross section of p-type (100) Si samples after MACE with Pt NPs (increased magnification from a to c).

As already described by Tsujino and Matsumura,\(^6\) a thick porous Si layer covers both the surface and the surrounding of the inner core etched by the NP (Figure 2a). After 10 min etching, the depth reached by the Pt NPs is around 6 \(\mu\)m (with respect to the top porous Si surface), corresponding to an etch rate of \(\sim 0.6 \mu\text{m/min}\). Figure 2b focuses on a single macropore, where the Pt NP is distinguishable slightly below the pore tip (since it was not cleaved right in the middle).
The cone angle $\alpha$ defined by the porous Si/c-Si interface is \(\sim 20^\circ\). Interestingly, the high magnification image of another pore tip in Figure 2c shows that the empty core has the exact diameter of the NP and reveals that there is no porous Si beneath it.

This is in contrast to MACE experiments with Au networks dedicated to form Si nanowires, for which etching involves the formation of a porous Si layer beneath the Au film.\(^{26}\) The lack of underlying porous Si in the present case, as well as for MACE with Ag NPs, seems to indicate that electropolishing governs dissolution at the Pt/Si interface. However, for a 3.63 M HF solution, the critical current density ($j_{fs}$) is \(\sim 75\) mA/cm\(^2\)\(^{25}\) whereas the obtained etch rate of 0.6 $\mu$m/min corresponds to a current of \(\sim 21\) mA/cm\(^2\) (assuming 3 $\ell^/$/dissolved Si atom), i.e., etching at the level of the NP should be in the porous Si formation regime. This discrepancy could be due to the actual presence of a very thin porous Si layer that is constantly etched by the NP as it sinks into the bulk or more likely to a lower HF concentration at the tip of the growing pore than in the bulk of the electrolyte due to mass transport limitation. In addition, the metal is an obstacle for the access of the electrolyte to the etching front and may also lower the effective HF concentration.\(^{27}\) This could imply a much lower $j_{fs}$ value and thus explain the lack of porous Si at the pore tip. As a matter of fact, experiments performed on (111) oriented and multicrystalline p-type Si show that the Pt NPs follow $<100>$ directions [unpublished results, cf. S5 in SI] which is in an indication of etching at $j_{fs}$.\(^{25}\)

When comparing Figures 1b and 2a, it is interesting to note that the density of macropores at the surface is much lower than the initial density of Pt NPs (by a factor of \(\sim 20\)). Only a fraction of the initial Pt NPs are found at the etching front, the others being embedded in the top porous Si layer due to the overlapping of macropores. In line with these findings, it is assumed that an increase of the Pt coverage (as in Figure 1c) leads to the same macropore density after etching. Conversely, it has been observed that decreasing it eventually leads to the formation of one macropore per Pt NP (i.e., isolated macropores, results not shown here).

**Electrochemical Characterization.** Cyclic voltammetry and open circuit potential (OCP, or rest potential) measurements were performed for “in situ” electrochemical characterization of the Pt/Si/electrolyte system. Figure 3 shows the evolution of OCP in the dark for bare and platinized Si (dotted blue and solid red lines, respectively), in contact with a solution of HF (0.5 M) and upon addition of $\text{H}_2\text{O}_2$ (0.17 M, $\rho = 0.75$). To test the effect of illumination on the electrochemical response, light was sequentially switched on and off during the experiment.

In presence of HF only, the OCPs in the dark of bare and platinized Si are close (\(\sim 0.36\) against \(-0.30\) V vs SCE, respectively). The OCP can be identified as a redox mixed potential resulting from the equilibration of positive and negative charge transfers at anodic and cathodic surface sites of the Si electrode. As pointed out in several studies from Chemla’s group, in the absence of a redox couple and dissolved O$_2$ in solution, this charge transfer corresponds to proton discharge (H$^+$ + e$^-$ + Si $\rightarrow$ Si-$\cdot$H, $E_{\text{HF}} = -0.06$ pH V vs SHE) and Si oxidation (Si + F$^-$ $\rightarrow$ Si-F + e$, E_{\text{Si}} = -1.2$ V vs SHE).\(^{28}\) Under these conditions, the OCP can be described by

$$E_{\text{OCP}} = \frac{1}{2}(E_{\text{Si}} + E_{\text{HF}}) + \left(\frac{kT}{e}\right) \ln \left(\frac{j_{OC}}{j_{OA}}\right)$$

where $j_{OC}$ and $j_{OA}$ are the cathodic and anodic exchange current densities, characteristic of the charge transfer rate on Si surface cathodic and anodic sites (cf. [22] for a detailed explanation of eq 4). The slightly more positive potential of platinized Si can be then be ascribed to a higher $j_{OC}$ resulting from the catalytic activity of Pt toward proton discharge, or more likely in our case toward the reduction of O$_2$ traces in the reaction media (O$_2$ + H$_2$O $\rightarrow$ H$_2$O + O$_2$, $E^\circ$ = +0.68 V vs SHE) since simple Ar bubbling was used without further precautions.\(^{29}\)

Upon addition of $\text{H}_2\text{O}_2$, both bare and platinized Si experience a shift in the OCP toward positive values coinciding with the beginning of electroless Si dissolution. This shift can also be ascribed to an increase in $j_{OC}$ due to the strong oxidizing character of H$_2$O$_2$ (H$_2$O$_2$ + 2H$^+$ + 2e$^-$ $\rightarrow$ H$_2$O + O$_2$, $E^\circ$ = +1.77 V vs SHE). The OCP is considerably more positive in the presence of Pt (+0.05 against \(-0.16\) V vs SCE for bare Si at the end of the experiment) which reflects the catalytic activity of Pt for H$_2$O$_2$ reduction during MACE.

In presence of HF, the effect of light is to shift the OCP toward positive potentials by 0.20 V (bare Si) and 0.14 V (platinized Si), while after $\text{H}_2\text{O}_2$ addition, scarce or no light influence is observed. This absence of photoresponse in the presence of $\text{H}_2\text{O}_2$ can be attributed to the high level of H$^+$ injection at the surface, such that the change in carrier concentrations by photogeneration becomes negligible.\(^{30}\) This effect is strongly accentuated for the platinized sample.

The analysis of the potential dependence of the MACE process was carried out by means of cyclic voltammetry (CV) measurements in the dark. Figure 4 shows the CVs corresponding to bare and platinized Si (dotted blue and solid red lines, respectively) in a solution of 0.5 M HF and 0.17 M H$_2$O$_2$ ($\rho = 0.75$), together with an enlarged view of the OCP zone as inset.

The current/potential curve of bare Si in HF/H$_2$O$_2$ is surprisingly identical to the response of p-type Si in HF.\(^{31}\) Under anodic polarization above OCP (\(-0.16\) V vs SCE), an exponential increase of the current density corresponding to the porous Si formation regime (h$^+$ diffusion control) leads to
the attainment of $I_{PS}$ ($\sim$9 mA/cm$^2$, forward scan), followed by the electropolishing regime (HF diffusion control).

Under cathodic polarization below OCP, the current density is close to zero. On the other hand, the current/potential response of platinized Si is fairly different: $I_{PS}$ is higher (12 mA/cm$^2$), probably because of the higher surface area rapidly induced by MACE, and a noticeable cathodic current is recorded, with a plateau value attained below $-0.5$ V vs SCE ($\sim-3.3$ mA/cm$^2$). As reported by Kooji et al. in the case of HF/HNO$_3$ etching, this cathodic current obtained for a p-type Si sample in the dark can be attributed to the collection of h$^+$ injected by the oxidizing agent, the plateau value being determined by kinetics. In the present case, a high level of injection is obtained when the reduction of H$_2$O$_2$ is catalyzed by Pt. The lack of cathodic current in the absence of Pt is in agreement with the very low etch rate of Si in HF/H$_2$O$_2$ solutions (nm/h).33

The inset of Figure 4 shows that the OCP of platinized Si (+0.03 V vs SCE, circle), an anodic current ($\sim-0.5$ mA/cm$^2$) is obtained for bare Si (arrow). This implies that under MACE conditions with Pt, porous Si formation will occur at any bare Si/HF area and not only in the close surroundings of the Pt NPs. This situation is similar to that reported by Kelly et al. for porous Si formation with galvanic cells in HF/H$_2$O$_2$ where the back surface of a Si wafer is fully metallized. Although MACE differs in the nanometric size of the metal contacts, the etching mechanisms and overall results are the same as for macroscopic galvanic cells. They imply however some interesting band modulation effects due to the embedding of the nanoscaled metal/Si junction within the much wider Si/electrolyte junction.

Local modifications of the band structures induced by the presence of metal NPs on the electrochemical properties of Si have already been reported by Nakato et al. for hydrogen photoevolution. They showed qualitatively that, depending on the NP size, specific electrochemical behavior can be expected. In the next section, we present a band bending modeling based on numerical simulations that gives a precise spatial description of the band structures of Pt/Si/electrolyte junctions at the nanoscale which helps to further rationalize the mechanisms of MACE with Pt NPs. This kind of simulation studies has been reported by Rossi and Lewis for nanometer-scale Ni arrays on Si electrodes and by Huang et al. in the case of metal assisted electrochemical etching with Ag NPs on p-type Si under anodic polarization in HF solutions. The latter authors have shown that the modulation of the band bending results in the collection of h$^+$ around the Ag NPs and there promotes a localized Si dissolution with the formation of mesopores.

**Band Bending Modulation.** Numerical simulations in 2D of the valence band modulation at the Pt/p-Si/electrolyte interfaces were carried out using SILVACO© ATLAS, a TCAD simulation software (cf. section S3.1 in SI for more details). The modeled structure consists of a p-type Si substrate with a thickness of 100 μm and a width of 0.1 μm, covered by a 12 nm large Pt contact in the middle, surrounded by two electrolyte contacts (a scheme is provided in SI, Figure S3).

Experimental Mott–Schottky (MS) plots of bare and platinized Si/HF junctions were first recorded in order to extract the experimental parameters required for the simulations, i.e., the doping density and flat band potential values (Figure S2 in SI). The experimental data and the detailed procedure for flat band calculation can be found in SI (sections S2 and S3.2). The Si/electrolyte and Pt/Si interfaces were both modeled as metal/Si contacts with two different work functions. The fitting between simulated and experimental MS plots was optimal when setting the work function of the electrolyte ($W_{el}$) and Pt ($W_{Pt}$) at 4.5 and 5.6 eV, respectively. These values are consistent with those found in the literature for p-type Si in HF solutions (on the basis of OCP values) and for polycrystalline Pt. For macroscopic contacts on p-type Si, work function values of 4.5 eV and 5.6 V give a rectifying (Schottky) contact and an ohmic contact, respectively.

*Figure 5a* shows a 3D plot of the Si conduction band (CB) and valence band (VB) modulations around the Pt NP for the system at equilibrium. A 2D projection of the VB energy in the (x,y) plane is displayed on the bottom plane of this graph, with the zero energy reference corresponding to e$^-$ and h$^+$ Fermi level (FL).

In the bulk of Si, the equilibrium CB and VB energies are found at 0.93 and $-0.19$ eV, respectively. The large difference between $W_{el}$ (4.5 eV) and $W_{Pt}$ (5.6 eV) induces a modification of the band bending at the surface close to the NP. Away from the Pt NP, i.e., at the p-Si/electrolyte Schottky junction, the surface VB energy is $-0.67$ eV, implying an energy barrier height of 0.48 eV for h$^+$ of the bulk. On the contrary, beneath the Pt NP, the surface VB energy is positive (by up to +0.43 eV), i.e., the VB is above the FL, indicating the presence of a strong accumulation layer of h$^+$. This accumulation layer extends up to ~4 nm in Si, where the VB crosses the FL. Therefore, the contact made by a Pt NP of 12 nm to p-type Si remains ohmic, as it is the case at the macroscopic scale. Note that this accumulation layer of h$^+$ is consistent with etching in the electropolishing regime beneath Pt (core of the mesopore).

The 2D modulation of the VB at equilibrium is shown in Figure 5b. Here, the energy is referenced with respect to the bulk VB energy, where the VB is flat. Most of the modulation actually occurs in a region of the order of the NP size, the space charge region (SCR) of the Si/electrolyte junction (violet areas) being fully recovered after ~25 nm from the NP borders.

In order to simulate the MACE process, the oxidizing power of H$_2$O$_2$ was mimicked by a positive polarization applied to Pt (against the electrolyte, cf. section S3 in SI for details). *Figure 5c* gives the 2D modulation of the VB under +0.2 V and Figure...
Under positive bias, the major effect originating from the ohmicity of the Pt contact is that the entire voltage drop actually falls at the p-Si/electrolyte interface. It is clear from Figure 6 (and also by comparing Figure 5b and c) that the VB in the bulk follows the applied polarization by the same extent. Consequently, the applied voltage (or the presence of H$_2$O$_2$) reduces the barrier height of the Si/electrolyte junction.

A more accurate “in situ” picture of MACE is given in Figure 7a, which represents the simulated 2D profiles of both the VB

Figure 6. 1D profiles of the VB energy for different Pt polarization (a) under the Pt NP ($x = 50$ nm) and (b) under the electrolyte ($x = 0$ nm).

and the space charge region (SCR) in Si and around the Pt NP once it has sunk into the substrate (cf. Figure S4 in SI for a closer view). For that, we have modeled a Pt NP with a size of 50 nm, buried at the bottom of a 1 μm deep mesopore without porous Si, and polarized at +0.2 V. Figure 7a shows that the width of the SCR surrounding the NP is approximately 400 nm (this value depends on the doping concentration of Si) and is constant almost from the pore tip to the surface. This should lead to a homogeneous supply of h$^+$ (hence porous Si formation rate) along the mesopore.

On the basis of the experimental results and numerical simulations, the scheme of Figure 7b is proposed to rationalize the formation of cone-shaped pores by MACE with Pt NPs. According to this scheme, injected h$^+$ from Pt NPs (reduction of H$_2$O$_2$) spread in the bulk due to the ohmic nature of the Pt/Si contact. This injection is split into two contributions: core etching in the electropolishing regime confined at the Si/Pt interface and anodic polarization of the bulk leading to porous
Si formation at the pore wall and at any p-Si/HF interface of the sample (e.g., front and back side). The porous Si layer thickness will be proportional to the etching time, i.e., large at the pore top and almost zero at the pore tip. Since the c-Si/porous Si interface is relatively straight (cf. Figure 2) we can assume that (after a certain time) both etching processes occur at constant rates, i.e., there is a stable supply of HF/H₂O₂ and a constant polarization of the bulk. The cone angle depends on the balance of etch rates such that an angle of 45° would actually imply that the two etching process have equal rates.

On n-type Si, the use of Pt NPs for MACE in HF/O₂ has been studied by Yae et al. Compared to the above results for p-type Si, a significant difference is that less porous Si is formed around the pores and at the sample surface. We have also tested n-type Si MACE with Pt under our experimental conditions for p-type Si. In agreement with Yae et al., the quantity of porous Si formed at the surface and around the pores for the n-type sample was significantly lower than that for p-type. Moreover, no porous Si was found at the back side of the samples, whereas the p-type Si sample exhibits a few hundred nm thick layer of porous Si at the back side. These results agree well with a positive polarization of bulk Si under MACE with Pt which results in an anodic current only when majority carriers are h⁺.

Several reports on MACE with different metal catalysts have already considered h⁺ injection and diffusion as key parameters to explain the (de)localized character of etching. For instance, a similar mechanism to the one exposed above, comprising vertical and sidewall etching, has been recently proposed by Li. et al. in an attempt to correlate the 3D profile obtained with p- and n-type Si samples during micro-MACE with μm-sized straight-line patterns of Au as catalyst. In complement to these works, our results (i) demonstrate the ohmic nature of the nanocontacts made with Pt NPs (that should also apply in the case of Au), (ii) rationalize the effect of h⁺ injection in the case of p-type Si as a polarization of the bulk, and thus (iii) explain the various morphologies obtained by MACE (cone-shaped pores, tapered nanowires, etc.).

Application of an External Potential. In line with the mechanism proposed above, the application of an external potential during MACE can be envisaged as a way to counterbalance or enhance the polarization of the bulk induced by Pt nanocontacts. This means controlling the band bending at the Si/electrolyte junction, hence the formation of porous Si at the pore wall. MACE under external polarization was performed under the same conditions than regular MACE (3.63 M HF and 1.2 M H₂O₂, ρ = 0.75). Figure 8 shows the current/potential response obtained for platinized Si in HF/H₂O₂, together with SEM images of samples etched for 10 min at different cathodic or anodic potentials (highlighted by circles).

SEM observations show that upon increasing the potential from cathodic to anodic values, the resulting Si nanostructures evolve from a collection of straight mesopores to an array of well-defined cone-shaped pores. At −1 V vs SCE (a), the porous Si layer around the core is thin and parallel to the wall (except at the tip) and becomes thicker as the potential is made more positive. The cone angle varies very significantly, from nearly 0° at −1 V up to 70° at +0.6 V vs SCE. The porous Si layer thickness at the surface follows the same trend, increasing from 2 to 6 μm (b–f).

These results can be readily understood taking into account that injected h⁺ are preferentially collected in the bulk under negative bias, whereas an additional h⁺ current from the bulk is supplied to the Si/electrolyte interfaces under positive bias. The formation of porous Si at the pore walls and at the surface is then reduced in the first case and enhanced in the second.

Recently, Li et al. have reported on analogous results obtained with an electric bias-attenuated MACE method that could be applicable for manufacturing microelectronic devices. With p-Si and Au catalysts, they found that a
negative electric polarization applied from the back side of Si substrates could avoid excessive etching occurring in traditional MACE. This is consistent with our experimental results and modeling because Au has a WF almost as large as Pt (5.47 vs 5.60 eV, respectively) and consequently Si/Au contacts are ohmic as well (as confirmed by simulations made with Si/Au NPs/electrolyte models but not shown here).

Chourou et al. have also reported on electrochemically assisted MACE of p-type Si (10–30 ohm.cm) in HF 4.5 M and H2O2 0.9 M (ρ = 0.83) with Pt NPs and a superimposed anodic current of 1 to 2 mA/cm2.21 In this case, straight pores with a cone angle of ∼5° are formed. In contrast, etching at very small anodic current close to OCP leads to much more opened cone shaped pores (25°) in our case. We assume that the main reason for these differences is related to the doping level of Si. With p-type Si, a large space charge layer at the bottom of the pores and a depletion of carriers between them can be expected. Hence, the collection of h+ should occur preferentially at the pore tip rather than at the pore walls and favor an in-depth growth.

Surface Texturization. The ability to vary the pore angle without constraint is interesting in the frame of Si surface texturization. This process aims at reducing the reflectance to improve light absorption in photodetectors and solar cells by creating surface morphologies that force the incident light to bounce back onto the device surface, and hence to interact with the material, rather than be directly reflected. A well-known example is the elaboration of a network of inverted square-based pyramids for monocrystalline (100) oriented Si solar cells. The walls and the angle of the pyramid (54.7°) are defined by the intersection of (111) planes revealed during an alkaline etching subsequent to a lithographic process. A lot of work is devoted to find surface treatments leading to macrostructures with higher aspect ratio (i.e., lower opening angles) to improve light coupling by increasing the number of light bounces.

In this context, a polarization of +0.4 V vs SCE was found to be optimal for the reduction of reflectance. SEM images of sample surface morphology after MACE under +0.4 V vs SCE and porous Si removal in HNO3/HF 99:1 V:V (5 min), are given in Figure 9a,b. Well-defined cone-shaped pores with an angle of ∼35°, corresponding to an aspect ratio (D/W) of 1.6, are covering the whole sample surface. The spread in size is relatively large, with the diameter (depth) ranging from 0.5 to 5 μm (0.8 to 8 μm). Because such morphologies are difficult to obtain even with advanced plasma etching techniques, MACE with Pt has therefore a strong potential for Si surface structuration. These structures advantageously reduce the surface reflectance down to 3% in the 500–1000 nm range, as can be seen on the reflectance spectrum of Figure 9d (circles). This reflectance value compares favorably with state of the art texturization techniques for Si, such as inverted square-based pyramids arrays (R ∼ 9%) used in record efficiency solar cells.46

The reflectance of this surface was modeled by using a ray tracing program presented in a previous publication.47 In this model, the surface is represented using a regular honeycomb formed of inverted hexagonal pyramids with 35° opening angles (Figure 9c), which is obviously an idealized description of the real sample. Figure 9d presents the comparison between modeled and measured reflectance for this particular surface, together with that corresponding to polished Si, taken as reference. For the polished Si surface, the agreement between simulation and experimental measurements is very good up to λ ≥ 1000 nm, since the model assumes that all the light penetrating Si is absorbed independently of the wavelength. However, in the infrared, the absorption coefficient of Si falls as the wavelength increases, and so, an increasing part of the incident light is reflected on the rear side of the substrate and exits by the front side, resulting in a large increase of the measured reflectance, as depicted by the experimental spectra. This limitation of the model for λ ≥ 1000 is not detrimental for its reliability at shorter λ.

Concerning the textured surface, as only perfect and regular hexagonal pyramids are considered to mimic the cone-shaped pore morphology, the values of reflectance provided by the model are lower than those exhibited by the real surface (0.34 vs 3.7% at 700 nm, respectively). Taking into consideration that the surface obtained experimentally is not homogeneously etched, with some distribution of etching angles and a small fraction which can be considered as nearly planar (cf. Figure 9a,b), the measured reflectance is actually a weighted sum of the reflectance of different parts of the surface. For example,
setting 10% of planar surface in the model would be sufficient to obtain an identical value of reflectance for simulation and experiments in the visible part of the spectrum. These results suggest that a better control of the experimental conditions during MACE, for example of the density and spatial distribution of the Pt NPs, could yield textures with even lower reflectance than 3%. Current efforts are being developed in our laboratory to pursue this goal with the aim to reach a reflectance below 1%.

CONCLUSION

A comprehensive detailed study of MACE with Pt NPs has been presented, paying particular emphasis on the understanding of the electrochemical mechanisms governing the etching and the resulting pore morphology. For that purpose, a combination of electrochemical techniques and 2D band bending modeling, with a positively biased NP at the top and buried in the Si substrate has been used. This novel kind of simulation mimics the “real” etching conditions and it has proved to be a powerful tool to describe the band modulation at the nanoscale. In our opinion, the approach followed in this work should provide new insights on MACE with other metal nanocatalysts in the future. The results obtained experimentally and from simulations allow concluding that h+ injection during MACE can be split into two contributions: core formation by direct dissolution of Si in contact with Pt and anodic polarization of the bulk with porous Si formation at the pore wall. These two simultaneous processes, and the resulting typical cone-shaped pore morphology, are a consequence of the ohmic nature of the Pt nanocontacts. Experiments in which an external polarization is applied during MACE have further confirmed this results and proved to be a remarkably simple and efficient method to achieve structures ranging from straight mesopores to cone-shaped macropores of controlled opening angle as the Si sample is biased from negative to positive potentials. High aspect ratio macropores obtained under moderate anodic bias leads to a reduction of the surface reflectance down to 3% in the 500–1000 nm wavelength range. This value is significantly lower than that obtained by state of the art texturization techniques (e.g., inverted pyramids with R ~ 9%). A 3D ray tracing simulation indicates that even lower reflectance could be achieved with cone-shaped pore arrays. Under optimized metallization, we believe that this tunable structuration of Si by MACE with Pt NPs has the potential to provide Si reflectance below 1%.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.6b09036.

Nanoparticle size distribution on p-type (100) Si surfaces, experimental Mott–Schottky plots, design of the Pt/Si/electrolyte structure used for modeling, simulated Mott–Schottky plots and MACE of (111) oriented Si (PDF)

AUTHOR INFORMATION

Corresponding Author
*Phone: +33 1 49 78 13 30; E-mail: bastide@icmpe.cnrs.fr.

Author Contributions
E.T. and S.L.G. made equal contribution to the manuscript

Notes

The authors declare no competing financial interest.

ACKNOWLEDGMENTS

The authors acknowledge the support of the French Agence Nationale de la Recherche (ANR), under grant ANR-14-CHE07-0005-01 (project PATTERN).

REFERENCES


