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¹ Low-Load Metal-Assisted Catalytic Etching Produces Scalable ² Porosity in Si Powders

³ Konstantin Tamarov, Riku Kiviluoto, Joseph D. Swanson, Bret A. Unger, Alexis T. Ernst, Mark Aindow, ⁴ Joakim Riikonen, Vesa-Pekka Lehto, and Kurt W. Kolasinski*



9 produces porous Si instead of Si nanowires. In this work, we demonstrate 10 that some of the features of LL-MACE cannot be explained by the present 11 understanding of MACE. With mechanistic insight derived from extensive 12 experimentation, it is demonstrated that (1) the method allows the use of 13 not only Ag, Pd, Pt, and Au as metal catalysts but also Cu and (2) judicious 14 combinations of process parameters such as the type of metal, Si doping 15 levels, and etching temperatures facilitate control over yield (0.065–88%),



¹⁶ pore size (3-100 nm), specific surface area $(20-310 \text{ m}^2 \cdot \text{g}^{-1})$, and specific pore volume $(0.05-1.05 \text{ cm}^3 \cdot \text{g}^{-1})$. The porous structure ¹⁷ of the product depends on the space-charge layer, which is controlled by the Si doping and the chemical identity of the deposited ¹⁸ metal. The porous structure was also dependent on the dynamic structure of the deposited metal. A distinctive comet-like structure ¹⁹ of metal nanoparticles was observed after etching with Cu, Ag, Pd, and, in some cases, Pt; this structure consisted of 10–50 nm main ²⁰ particles surrounded by smaller (<5 nm) nanoparticles. With good scalability and precise control of structural properties, LL-MACE ²¹ facilitates Si applications in photovoltaics, energy storage, biomedicine, and water purification.

22 KEYWORDS: low-load metal-assisted catalytic etching (MACE), porous silicon, powder, porosity, nanoparticles, silicon nanostructures

23 INTRODUCTION

²⁴ Metal-assisted catalytic etching (also known as metal-assisted ²⁵ etching or metal-assisted chemical etching), MACE, is a simple ²⁶ and versatile method of Si micropatterning. Nanostructuring of ²⁷ Si with MACE has attracted considerable attention in the past ²⁸ few years to fabricate nonreflecting surfaces for photo-²⁹ voltaics,¹⁻³ to prepare patterned structures for sensing,⁴⁻⁶ and ³⁰ to overcome the huge volume expansion of Si-based Li-ion ³¹ battery anodes.^{7–9} Porous Si nanowires (NWs) and nano-³² needles formed by MACE have been used in a large number of ³³ different biological applications including sensing,¹⁰ studies of ³⁴ cell–Si interfaces,^{11,12} and therapeutic delivery.^{13–15}

Typically, MACE is performed in a two-step process (metal deposition and etching are performed separately)¹⁶ although one-step MACE (combined metal deposition and etching)^{17,18} has also been demonstrated successfully. In the two-step process, a piece of Si is first immersed into a metal plating solution consisting of hydrofluoric acid (HF), water, and dissolved metal salt, e.g., AgNO₃. Ag⁺ ions deposit as particles, dendrites, or layers on the Si surface, and this process can be further controlled by applying masks or even more sophisticated methods such as lithography, thermal decomposition, or sputtering. In the second step, the Si is immersed in a solution de containing HF and oxidant, usually H_2O_3 . The metal catalyst facilitates hole injection into the Si valence band, which controls 47 the rate of the etching reaction as long as the concentration of 48 HF is sufficiently high. 49

Due to the high concentrations of holes near the metal/Si 50 interface and generally high metal coverage, the most common 51 outcome of MACE is the formation of 80–150 nm etch track 52 pores bored by the cooperative motion of metal particles, which 53 are formed dynamically at the beginning of etching.^{19,20} The 54 walls of the pores are often called Si NWs, even though etching 55 first forms interconnected walls and ridge-like structures that are 56 cleaved readily by, e.g., ultrasound or capillary forces to form 57 stand-alone Si NWs.

In an attempt to improve the cost-efficiency of the MACE 59 process, several recent studies focused on etching inexpensive Si 60 grains, chunks, and powders instead of wafers. It is especially 61 interesting that even in powders, metal particles still move 62 cooperatively, descend into Si perpendicular to particular 63

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⁶⁴ facets,¹⁹ and produce etch track pores and Si NWs similar to ⁶⁵ MACE of wafers.

Thus, MACE was applied to inexpensive and relatively impure metallurgical-grade (MG) powders. Preferential etching of the impurities led to purification of Si,^{21,22} which made it possible to apply the etched particles for Li-ion battery anodes^{23,24} and photovoltaics²⁵ and to produce biocompatible nanostructures r1 for nanomedicine.²⁶

Scaling up MACE of powders requires control of the etching 72 73 process. The rate of this highly exothermic reaction is 74 proportional to the surface area. The high surface area of 75 powders compared to that of Si wafers leads to considerable heat 76 production during MACE. Thermal management during the 77 reaction is crucial to improve the uniformity and reproducibility 78 of the product. Similar to regenerative electroless etching 79 (ReEtching),²⁷ we exploited a syringe pump to inject oxidant at 80 a constant rate, giving better control over MACE and avoiding 81 excessive heating. In addition, we demonstrated the use of a 82 syringe pump to deposit Ag controllably. Combined with 83 vigorous stirring of Si powder and addition of acetic acid to HF, ⁸⁴ we achieved uniform Ag deposition over a wide range of n(Ag). ss A_{Si}^{-1} , from 10 to 0.002 mmol·m⁻², where n(Ag) is the amount of ⁸⁶ silver and A_{Si} is the surface area of the Si powder subjected to 87 MACE.²⁰

Varying the Ag coverage, we demonstrated that large etch so track pores produced by the correlated motion of metal catalyst nanoparticles and Si NWs are not the only outcome of Ag MACE.²⁰ When the Ag amount is low in the depositing solution $(n(Ag) \cdot A_{Si}^{-1} \le 0.06 \text{ mmol} \cdot \text{m}^{-2})$, Ag deposits as individual 10– 320 nm nanoparticles instead of a thick layer of interconnected 4 clusters and dendrites. During etching, these small nanoparticles 55 move in an uncorrelated manner producing random pores with 6 10–20 nm diameter and, if the p-type doping level is high 97 enough, 4–6 nm tortuous pores. Thus, a mesoporous layer is 98 formed in contrast to Si NWs. We call this regime low-load 99 MACE or LL-MACE to emphasize its difference from 100 conventional high-load MACE or HL-MACE.

For a comprehensive discussion of HL-MACE, and structures 101 102 accessible through this process, we refer to the recent review by 103 Alhmoud et al.²⁸ Here, we summarize a few salient points. 104 Depending on the metal catalyst, Si doping level, and oxidant amount, the porosity of etch track pore walls and Si NWs in HL-105 106 MACE can be varied. Etching highly doped p-type Si ($\rho = 0.01 -$ 107 0.02 Ω ·cm) with Ag generally resulted in NWs with porous walls, while lower doping levels led to solid Si NWs.^{29,30} Etching 108 with Au produced porous Si NWs for a Si resistivity of up to 1 Ω · 109 110 cm, depending on the H_2O_2 concentration.³¹ Pt formed both a 111 porous layer and helical etch track pores on 10 Ω ·cm p-type 112 wafers; the latter was attributed to the multifaceted crystalline 113 shape of the Pt particles. Pd was found to be more convoluted ¹¹⁴ than others as it catalyzed etching even without an oxidant and ¹¹⁵ resulted in electropolishing of Si. $^{32-34}$ HL-MACE with Cu never 116 produced defined etch track pores^{35,36} and Si NWs, except in the 117 case of electrochemical etching under applied bias.³⁷ This is due to the low electronegativity and reduction potential of Cu 118 compared to those of other metals, which prevents efficient 119 120 deposition and leads to enhanced dissolution during etching. 121 Overall, depending on the chemical identity of the metals, their ¹²² catalytic activity toward the reduction of H_2O_2 , and induced ¹²³ band bending,^{33,38,39} different morphologies were produced by 124 HL-MACE.

Herein, we present a comprehensive study of LL-MACE of Si powders, where we use low amounts of Cu, Ag, Au, Pt (12μ mol-

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m⁻²), and Pd (1.2 μ mol·m⁻²) to etch mesopores. Various Si 127 grades were used including powders prepared from P++, P+, P, 128 N+, N++, and undoped wafers (UW) as well as metallurgical- 129 grade powders with either 99.6% (MG) or 99.997% (MC10) 130 purity. We demonstrate that by choosing Si grade, metal, and 131 temperature, one can vary over a wide range the yield (0.065– 132 88%), pore size (3–100 nm), surface area (20–310 m²·g⁻¹), and 133 pore volume (0.05–1.05 cm³·g⁻¹).

Overall, we found a great variety of porosity after LL-MACE. 135 This can be applied to optimize porosity and surface area for 136 applications such as Si-based Li-ion battery anodes, antireflection coatings for photovoltaics, and to load drugs or 138 biomolecules of a specific size. Furthermore, LL-MACE is 139 easy to scale up, uses minimal amounts of metal, and works well 140 for certain Si grades with Cu, opening a pathway to significantly 141 less expensive processing. 142

EXPERIMENTAL SECTION

Samples were prepared and characterized in line with previously 144 published methods.^{20,32,40} Briefly, Si powders of known purity/doping 145 levels were milled and cleaned prior to the deposition and nucleation of 146 a metal catalyst. Subsequently, etching was initiated by the addition of 147 oxidant (H_2O_2) . Both nucleation and etching were aided by the use of 148 acetic acid as a surfactant and steady metered injection of dissolved 149 metal ions or H₂O₂ with a syringe pump. The sample temperature 150 during reaction was regulated by immersing the reaction vessel in either 151 an ice/water or temperature-controlled water bath. After well-defined 152 rinsing and drying procedures, the samples were characterized by plain- 153 view and cross-sectional scanning electron microscopy (SEM) with 154 focused ion beam (FIB) cutting. FIB sectioning was also used to 155 prepare samples for transmission electron microscopy (TEM). Further 156 characterization was performed with N2 sorption followed by analysis 157 with the Brunauer-Emmett-Teller (BET) model and Barrett- 158 Joyner-Halenda (BJH) theory. The detailed methods can be found 159 in the Supporting Information. 160

RESULTS

Controlled injection together with acetic acid as a surfactant 162 resulted in the uniform coverage of Si surfaces with individual 163 5-30 nm metal nanoparticles separated by similarly sized 164 gaps.^{20,32} Specifically for Ag,²⁰ we found deposited nanoparticles 165 with a mean size of 19.0 \pm 9.5 nm and a center-to-center 166 separation of 25.6 \pm 9.0 nm (uncertainty given as 1 standard 167 deviation) to be responsible for catalyzing efficient etching in the 168 LL-MACE regime. These values were calculated under the 169 assumption that the particle surfaces were perfectly planar. As 170 shown previously,²⁰ these well-spaced nanoparticles move in a 171 completely uncorrelated manner during etching, following paths 172 determined by the minimum energy required to etch out one 173 atomic plane of Si.¹⁹

In the etching step of LL-MACE, the H_2O_2 solution was also 175 injected in a controlled manner using a syringe pump; this 176 provided a minimal steady-state supply of the oxidant that 177 significantly reduced the rate of heat generation. In most cases, 178 the temperature of the etching solution did not exceed the 179 temperature of cooling the bath by more than 12 °C. It should be 180 stressed that, while Ag, Pd, Pt, and Au are commonly applied in 181 HL-MACE, ¹⁶ Cu is never used because it fails to deposit on Si in 182 a sufficient quantity and it dissolves efficiently in the presence of 183 excess H_2O_2 . However, the use of a syringe pump for the dosing 184 of H_2O_2 enables us to reduce its steady-state concentration to a 185 level that does not remove Cu nanoparticles from the etch front. 186

Application of different metals in LL-MACE resulted in 187 distinctive morphologies of etched pores, yet they also had some 188 fi

f1

f2

189 common features. Top-view SEM images (Figure 1 and Sections
190 S3-S6, Supporting Information) show that Cu and Pt keep the



Figure 1. General view of $11-25 \,\mu\text{m}$ MG Si particles after LL-MACE with Cu, Ag, Pd, Pt, and Au. The scale bar represents 2 μm .

191 Si particle surfaces flat and edges sharp. A unique feature of Pt
192 particles is that they tend to move laterally across the Si surface
193 producing many, sometimes helical channels before descending
194 into Si. Ag, Pd, and Au, on the other hand, smoothen Si edges
195 while roughening initially flat facets, which produces a fuzzy
196 appearance that is distinctly different from the texture of
197 particles produced by HL-MACE.

Cross-sectional images are presented in Figure 2. An enlarged 198 199 version of Figure 2 with additional annotation can be found in 200 the Supporting Information (Figure S2.1). These images reveal 201 that the etched Si for all metals consisted of two distinct regions: 202 (1) etch track pores in the direction of the metal nanoparticle 203 movement and (2) a layer of tortuous pores surrounding the 204 tracks or extending deep into Si. The tortuous pores were similar 205 to those observed in ReEtching of Si powders, where no metal 206 catalyst was deposited on P++ and MG Si.²⁷ For Cu, Ag, and Pd, 207 tortuous pore regions tended to surround etch track pores 208 making the boundary between etched and bulk Si uneven, 209 composed of connected spherical regions. Cu nanoparticles 210 sometimes were not surrounded by tortuous pores. In contrast, 211 the boundary between tortuous pores and bulk Si for Pt and Au 212 was remarkably flat despite different Pt and Au particle 213 morphologies (Figure 2). The observation of both etch track 214 and tortuous pores is consistent with previous results such as 215 those of Hochbaum et al.,⁴¹ who were the first to observe porous 216 nanowire formation, Chiappini et al.,³¹ and Patil et al.⁴²

The dynamic nature of the metal nanoparticle structure 217 218 caused by the etch process is illustrated by a comet-like structure 219 reported here for the first time (Figures 2 and S2.1 and Sections 220 S3-S6, Supporting Information). Cu and Pd nanoparticles had 221 rather smooth although not round surfaces, and they were 222 always followed by a halo of small <5 nm nanoparticles. The 223 large metal nanoparticles form the cores of these comet-like 224 structures; they were 50-80 nm for Cu and 30-50 nm for Pd. 225 Interestingly, Cu "comets" have small nanoparticle "tails" behind 226 their etch track, while Pd "comet" cores were surrounded by 227 small nanoparticles on all sides. Pt particles possessed rich 228 irregular structures, which appear to form by aggregation at the 229 beginning of etching when particles moved extensively along the 230 Si surfaces. These aggregates expose different Pt crystalline 231 planes, which, we suggest, make the change of etch direction 232 easy and, in some cases, produce helical cavities. Despite the 233 previously considered stability of Pt particles during MACE, 234 they sometimes, although not always, showed comet-like 235 structures and were surrounded by small nanoparticle satellites. 236 Au nanoparticles were the most stable and round; they never showed a halo of small nanoparticles and kept their sizes in the 237 238 range of 10-40 nm during etching.

The outcomes of LL-MACE changed dramatically with the 240 metal catalyst, Si grade (Table 1), and temperature (Figure 3 241 and Sections S3–S12, Supporting Information). In SEM images, 242 Cu displays the most prominent change in structure with the



Figure 2. Cross-sectional bright-field (BF) and high-angle annular dark-field (HAADF) scanning transmission electron microscopy (STEM) images after LL-MACE of 11–25 μ m P++ (for Cu) and 2– 44 μ m MG (for other metals) powders in an ice bath. The left column (HAADF, metal appears light) shows the border between the bulk and etched Si: Cu and Pd produce an irregular border and tortuous pores surrounding metal particles; Pt and Au present a sharp and uniform border between the porous layer and solid core. The middle column (HAADF) takes a closer look at the porous layer showing both etch track pores produced by the metal particle movement and tortuous mesopores produced by remote etching. The right column (BF, metal appears dark) depicts the structure of metal particles: Cu and Pd always present a halo of small particles following big ones; Pt sometimes presents a halo of small particles and the large particles have an irregular structure; Au never shows the halo. An enlarged, annotated version of this figure and more images of metal particles are shown in the Supporting Information.

increasing temperature. For low-temperature (LT) etching in an 243 ice bath, Cu particles descend into Si and produce pores. At high 244 temperature (HT, water bath at 43 °C), most of the Si becomes 245 electropolished: P++, MG, and MC10 present broken Si pieces 246 with smooth surfaces, while UW and N++ have macropores with 247 crystallographically defined edges. N++ powder additionally 248 presents numerous uniformly distributed openings of tortuous 249 pores at HT (Figures 3 and S3.10, Supporting Information). 250 Other metals demonstrate less pronounced differences between 251 LT and HT. One can notice similarity for powders etched with 252 Ag and Au. Pd and Pt show distinctive morphologies, which, 253 however, look similar for all Si grades. Pt is the only case that 254

abbreviation	$\stackrel{\rm resistivity}{(\Omega \cdot cm)}$	doping type	doping density (cm ⁻³)	particle size (µm)
$P++^{b}$	0.0186	boron	4×10^{18}	11-25
$P+^{b}$	0.105	boron	3×10^{17}	11-25
P^{b}	25	boron	5×10^{14}	11-25
UW ^b	>5000	boron	$<1 \times 10^{12}$	11-25
$N+^{b}$	1.15	phosphorus	3×10^{15}	11-25
$N++^{b}$	0.0144	phosphorus	3×10^{18}	11-25
$MG^{c,d}$		Fe/Ti/Ca/Al		11-25
MC10 ^{c,e}		Fe/Ti/Ca/Al		1.5-10

^{*a*}Dopant concentrations were calculated using the Caughey–Thomas expression⁴³ for electron and hole mobilities. ^{*b*}Ground single-crystal (100) wafers from Okmetic Oy, Finland. ^{*c*}Metallurgical-grade Si from Elkem Silicon Materials, Norway. ^{*d*}Purity: 99.8%. ^{*e*}Purity: 99.997%.

255 allows easy observation of the openings of tortuous pores on 256 SEM images and the increase of their diameters with the increase 257 of temperature.

Measured pore size distributions for P++, UW, and N++ 259 powders (Figure 4) and for MG and MC10 powders (Figure 260 S13.1, Supporting Information) after LL-MACE at three 261 temperatures reveal quantitative insights into the structure of 262 etched Si because the presence or absence of tortuous pores 263 (<10 nm) can be easily recognized. It should be noted here that 264 the Barrett–Joyner–Halenda (BJH) analysis applied to N₂ 265 desorption isotherms is not sensitive for large pores (>75 266 nm). Therefore, etch track pores, for example, produced by Pt 267 particles, are not visible in the pore size distributions.

Cu is the metal with the lowest reduction potential, and it readily dissolves especially at HT, when holes are injected into it readily dissolves especially at HT, when holes are injected into it rate than at LT and room temperature (RT). Consequently, it may be anticipated that Cu LL-MACE either rate (1) partially follows a ReEtching-like mechanism where Cu ions rate tholes into Si directly and then get regenerated by H_2O_2 to repeat the process and/or (2) preferentially adsorb at step and defect sites on the Si surfaces akin to underpotential deposition, which in turn causes step-flow etching rather than porosification. The latter observation is consistent with the observation of 277 crystallographically defined etched features noted above. The 278 combination of these effects results in complete electropolishing 279 of Si at HT. At LT, Cu is capable of producing tortuous pores for 280 highly doped P++ and N++ powders (Figure 4) and MG 281 (Section S3, Supporting Information). The increase of the 282 etching temperature to RT and HT for P++ Si results in the 283 formation of mostly etch track pores by comet-like particles. In 284 the case of N++ powder, tortuous pores were observed at all 285 temperatures. Their volume was the highest in N++ among all Si 286 grades and their average width increased with the temperature. 287 We attribute this to the partial recombination of injected holes, 288 which prevented electropolishing. For UW wafer, a minor 289 component of etching is the formation of pores below 100 nm, 290 while macropore formation represents the majority (Figure 3). 291

Use of Ag significantly changes the outcome of LL-MACE 292 compared to that of Cu-catalyzed etching (Figure 4 and Section 293 S4, Supporting Information). The volume of tortuous pores for 294 P++ powder at LT ($0.037 \text{ cm}^3 \cdot \text{g}^{-1}$) is much smaller than for Cu 295 ($0.18 \text{ cm}^3 \cdot \text{g}^{-1}$; Tables S3.2 and S4.2, Supporting Information) 296 and increases with increasing temperature (note the different *y*- 297 axis scales in the graphs). Etch track pores produced by 10–30 298 nm Ag particles are clearly visible. They become wider at HT, 299 possibly due to high hole injection and Si polarization, which 300 results in an additional etching around Ag particles and thus pore 301 widening. In UW powder, Ag is only able to make etch track 302 pores, while in N++ tortuous pores were observed again and in 303 higher volume than in P++. 304

Pd is the most peculiar among the metals. Pd loading had to 305 be decreased by a factor of 10 to avoid complete Si dissolution. 306 Pd was found to create a moderate number of tortuous pores in 307 P++ and a high number in N++ powder, while UW powder is 308 mostly electropolished (Figure 4). The average pore width 309 increased for the former cases with the increase of temperature 310 along with the decrease of volume. We attribute this to the 311 enhanced electropolishing triggered by a higher rate of hole 312 injection into Si at RT and HT. N++ Si at HT was completely 313 dissolved (yield was 0.06%). Etch track pores were hardly 314



Figure 3. SEM images of the outer surfaces of Si particles after LL-MACE of P++, UW, and N++ $11-25 \mu$ m powders with different metals. Low-temperature (LT) and high-temperature (HT) columns show particles etched in an ice bath and water bath at 43 °C, respectively. The scale bar represents 200 nm.

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Figure 4. BJH pore size distributions of P++, UW, and N++ powders after Cu, Ag, Pd, Pt, and Au LL-MACE at three different temperatures: low temperature (LT, ice bath), room temperature (RT, water bath), and high temperature (HT, water bath at 43 $^{\circ}$ C).

315 observed (small peak around 40–50 nm) in the few Si particles 316 that remain.

Pt and Au are the two metals that could etch tortuous pores in 317 318 all Si grades (Figure 4). Notably, Pt did not distinguish between 319 P++ and N++ powders. Pore sizes and volumes are very similar 320 in both cases; the average pore widths and volumes increased with the increase of etching temperature. In Au LL-MACE, 321 322 tortuous pore volumes for N++ powder were 1.6 \pm 0.1 times 323 higher than for P++. Pore volumes, just as for Pt, grew with the increase in temperature. However, for UW powder, the pore size 324 and volume change as a function of temperature are opposite for 325 326 Pt and Au. While for Pt the volume of tortuous pores decreases $_{327}$ from 0.17 to 0.04 cm³·g⁻¹, for Au it grows from 0.02 to 0.11 cm³· g^{-1} with the change from LT to HT. Note here again the clear 328 329 presence of etch track pores produced by 10-40 nm Au 330 nanoparticles similar to that of the Ag ones. This explains the 331 resemblance between SEM images in Figure 2 for Ag and Au. 332 However, that is the only resemblance between them; Au clearly 333 outperforms Ag in the etching of tortuous <10 nm pores for all Si grades. 334

f5

The specific characteristics of P++, UW, and N++ powders 335 336 after LL-MACE are summarized in Figure 5, and for MG and 337 MC10 powders in Figure S13.2 (Supporting Information). It can be seen that applying different metals, Si grades, and 338 temperatures, a wide range of specific surface areas, pore 339 volumes, and yields are accessible through LL-MACE. Depend-340 341 ing on the etching parameters, surface areas were found to vary 342 from 14 to 310 m²·g⁻¹, pore volumes from 0.07 to 1.03 cm³·g⁻¹, 343 and yields from 0.06 to 80%. The lowest values for surface areas 344 and pore volumes were measured for Cu and Pd, especially for 345 UW powder at HT. The highest values were unexpectedly 346 measured for N++ powder etched with Au. Yields were the

lowest when electropolishing appeared, particularly for Cu and 347 Pd at HT. 348

A general correlation between specific surface area and pore 349 volume can be observed; their increase (or decrease) correlates 350 with each other in most etchings. Yield, however, does not show 351 such a good correlation in many cases. This discrepancy is 352 because pore volumes were quantified with N₂ sorption and 353 calculated using the BJH method, which is applicable to 354 mesopore and small macropore range (2-75 nm). On the other 355 hand, yield accounts for all Si removed by the etch track pore 356 formation, etching of tortuous pores, and electropolishing. To 357 quantify the relative contribution of mesopores to the total 358 amount of Si etched, we introduce the ratio of $v_{\text{pore}}/v_{\text{etched}}$. It 359 connects two measured values: pore volume δ_{pore} from BJH 360 analysis and yield. The ratio can be calculated using the 361 following equation (see the derivation in Section S1.4, 362 Supporting Information) 363

$$\frac{\nu_{\text{pore}}}{\nu_{\text{etched}}} = \frac{\delta_{\text{pore}}\rho_{\text{Si}}Y}{(1-Y)} \times 100\% \tag{1}_{364}$$

where $\delta_{\text{pore}} [\text{cm}^3 \cdot \text{g}^{-1}]$ is the specific BJH pore volume, i.e., the 365 volume of pores significantly smaller than ~100 nm, $\rho_{\text{Si}} = 2.33 \text{ g} \cdot 366 \text{ cm}^{-3}$ is the bulk density of Si, and Y is the yield. 367

When v_{pore}/v_{etched} is close to 0%, but the yield is high, most of 368 the etching produced macropores. If v_{pore}/v_{etched} is close to 0% 369 and the yield is also small (Cu and Pd at HT), electropolishing 370 took place. Ag produced only etch track pores in UW powder 371 and $v_{pore}/v_{etched} = 10\%$. High values of v_{pore}/v_{etched} (as high as 372 50–90%), specific surface area, and pore volume were found for 373 Pt. Thus, Pt is the optimal catalyst to form mostly tortuous <10 374 nm pores; the large etch track pores visible on SEM images 375



Figure 5. BET specific surface area, yield, BJH pore volume, and ratio of $v_{\rm pore}/v_{\rm etched}$ for P++, UW, and N++ powders after Cu, Ag, Pd, Pt, and Au LL-MACE at three different temperatures. The value of $v_{\rm pore}/v_{\rm etched}$ describes the ratio of the BJH pore volume (pores < 150 nm) to the total volume of etched-out Si.

376 account for a minority of the etched Si. Pt is, however, limited to 377 producing a relatively small surface area for UW powder. 412

Au is the optimal catalyst for etching of tortuous pores in all Si 378 grades. It is the metal that created the highest specific surface 379 areas of 310 m²·g⁻¹ for N++ powder and 190 m²·g⁻¹ for P++ 380 powder, and it is the only catalyst to produce up to 100 m²·g⁻¹ 381 surface area and 0.5 cm³·g⁻¹ pore volume in UW Si at HT. The 382 ratio of ν_{pore}/ν_{etched} adopts moderate values of 25–50% 383 depending on temperature, possibly due to etch track pores 384 that are larger than 100 nm and/or partial electropolishing. 385

Finally, we compared all of the Si grades at specific 386 temperatures for all metals to (1) understand the role of 387 impurities in MG and MC10 powders and (2) evaluate the effect 388 of intermediate dopant levels in P+, P, and N+ powders on pore 389 size distributions (Figure 6). The temperatures were selected to 390 f6 avoid electropolishing and to produce a high number of tortuous 391 pores. Specific surface areas, pore volumes, yields, and $v_{pore}/392$ v_{etched} values can be found in Section S14, Supporting 393 Information. Figure 6 demonstrates unambiguously that there 394 is a threshold for the appearance of a high volume of tortuous 395 pores. They are etched efficiently in highly doped nearly metallic 396 P++ and N++ powders, as well as 99.6% pure MG Si, 397 independent of the metal catalyst used. Interestingly, N++ 398 outperforms P++ in all cases, although it was generally 399 considered that in MACE remote etching appears mainly in 400 the highly doped p-type Si.²⁸ All moderately doped Si powders, 401 UW Si, and 99.997% pure MC10 powder show nearly the same 402 outcome. MC10 has somewhat larger pore volumes and surface 403 areas than P+, P, UW, and N+ powders because of the smaller 404 particle size, 1.5–10 μ m compared to 11–25 μ m for others. 405 Thus, a higher initial exterior surface area is accessible for 406 etching. The only difference between P+, P, UW, and N+ 407 powders can be observed for Pt-etched P+ Si, which has a larger 408 volume of tortuous pores than the others. MG powder is 409 somewhere in between N++ and P++. With Cu, MG resembles 410 N++, while with Au it resembles P++. 411

DISCUSSION

Mechanism of LL-MACE. The presence of a hole near the Si 413 surface is the essential requirement to initiate Si etching in HF- 414 based solutions. Holes polarize surface Si-H bonds making 415 them susceptible to fluoride ion attack and eventually resulting 416



Figure 6. BJH pore size distribution after (a) Cu, (b) Ag, (c) Pd, (d) Pt, and (e) Au LL-MACE of different Si grades at the selected temperatures. The temperatures producing a high volume of tortuous pores are shown.

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Si type $E_{\rm F} ({\rm eV})^a$	Cu 4.94 eV ^b	Ag 4.74 eV valence 2^{c}	Pd 5.6 eV valence 2/4	Pt 5.93 eV valence 4	Au 5.31 eV valence 2
P++	-0.37 eV ^d /8.6 nm ^e	−0.57 eV/10 nm	0.29 eV/1.4 nm	0.62 eV/1.4 nm	0.004 eV/1.4 nm
0.04	P depletion ^f	P depletion	P accumulation/degeneracy	P accumulation/degeneracy	P weak accumulation
P+	−0.3 eV/23 nm	−0.5 eV/28 nm	0.36 eV/5.3 nm	0.69 eV/5.3 nm	0.07 eV/5.3 nm
0.11	P depletion	P depletion	P accumulation/degeneracy	P accumulation/degeneracy	P weak accumulation
Р	-0.14/169 nm	−0.34 eV/230 nm	0.52 eV/130 nm	0.85 eV/130 nm	0.23 eV/130 nm
0.27	P depletion	P depletion	P accumulation/degeneracy	P accumulation/degeneracy	P accumulation
UW	$0.02/4~\mu{ m m}$	$-0.18 \text{ eV}/1.5 \ \mu\text{m}$	0.68 eV/2.9 μm	1.01 eV/2.9 μm	0.39 eV/2.9 μm
0.43	P weak accumulation	P depletion	P accumulation/degeneracy	P accumulation/degeneracy	P accumulation
N+	0.48/320 nm	0.28 eV/250 nm	1.14 eV/200 + 60 nm	1.47 eV/200 + 60 nm	0.85 eV/200 + 60 nm
0.88	N depletion	N depletion	N depletion + inversion	N depletion + inversion	N depletion + inversion
N++	0.66/21 + 6 nm	0.46/9.9 nm	1.32 eV/21 + 6 nm	1.65 eV/21 + 6 nm	1.03 eV/21 + 6 nm
1.06	N depletion + inversion	N depletion	N depletion + inversion	N depletion + inversion	N depletion + inversion

^{*a*}Fermi energy $E_{\rm F}$ is calculated from the top of the valence band $E_{\rm V}$, which is set to 0. Si work function in bulk is $\Phi_{\rm Si} = \chi_{\rm Si} + E_{\rm g} - E_{\rm F}$, where $\chi_{\rm Si}$ is the Si electron affinity and $E_{\rm g}$ is the band gap. ^{*b*}Metal work functions $\Phi_{\rm M}$ for (111) planes are taken from ref 56. ^{*c*}Molar stoichiometry $n({\rm VO}_2^+)/n({\rm Si})$ for MACE with ${\rm VO}_2^+$.³² ^{*c*}dIdeal magnitudes of band bending $E_{\rm BB}$ calculated by subtracting the Si work function $\Phi_{\rm Si}$ from metal work function $\Phi_{\rm M}$. For $E_{\rm BB} < 0$, valence and conduction band edges bend downwards (hole depletion for the p-type Si), for $E_{\rm BB} > 0$ —upwards (electron depletion for the n-type Si). ^{*e*}Schottky depletion space-charge layer $D_{\rm dep}$ width was calculated using eq S15.1 (Supporting Information). Equation S15.4 was used for inversion layer width. The width of the accumulation layer is the Debye length $L_{\rm D}$ calculated using eq S15.3. ^{*f*}Classification of the space-charge layer based on the doping type, density, and magnitude of band bending $E_{\rm BB}$.

417 in the dissolution of Si. Various Si etching methods, including 418 electrochemical anodization, stain etching, and MACE, require 419 holes and they satisfy it using different approaches. In 420 electrochemical anodization of Si, holes present in Si due to 421 doping, light illumination, or spontaneous bulk generation are directed to the surface by an applied bias. To form porous Si, the 422 423 bias should induce a small enough current density to make the 424 current-doubling pathway of Si dissolution dominant. In this 425 pathway, an injection of 1 hole into the valence band and 1 426 electron into the conduction band (that is, a valence 2 process) 427 is involved in the dissolution of 1 Si atom.⁴⁴ With the increase of 428 bias strength, the current-quadrupling (valence 4) pathway 429 involving 4 holes per Si atom becomes dominant and results in 430 complete dissolution of Si, i.e., above-critical-current density $_{431}$ etching. In stain etching, a strong oxidant (such as HNO₃) 432 present in the HF solution is able to oxidize Si directly, thus 433 injecting holes into the valence band.⁴⁵ Depending on the concentrations, stain etching can form porous Si or extremely 434 435 flat surfaces by electropolishing.

In MACE, H_2O_2 is the most commonly used oxidant.^{16,28} It 436 437 has very poor hole injection kinetics into the Si valence band, 438 and the etch rate in HF/H₂O₂ is far below 0.005 nm \cdot s⁻¹⁴⁶ for the $_{439}$ H₂O₂ concentrations used in this study. The inability of H₂O₂ to 440 induce Si etching directly is possibly due to the need to break 441 H₂O₂ into two -OH radicals for charge transfer to occur. In HF 442 solutions, Si surfaces are H-terminated and cannot dissociate $_{443}$ H₂O₂ at an appreciable rate. Thus, a catalyst is necessary to 444 reduce H₂O₂ and to inject holes into Si; many different metals 445 were studied for MACE including most commonly Ag, Au, Pd, 446 Pt,^{16,28,32} and Cu³⁶ and more exotically Ru,⁴⁷ Rh,⁴⁸ and Ni.⁴⁹ Ag 447 and Au are the most frequently applied catalysts, as their high 448 loading forms ordered arrays of Si NWs after HL-MACE. Cu 449 does not deposit well on Si surfaces and dissolves after oxidation 450 with high-concentration H2O2, thus making it impossible to etch 451 Si NWs.

The electrochemical reactions involved in MACE consist of a $_{453}$ cathodic reaction on the metal catalyst and an anodic reaction $_{454}$ on Si. The overall reaction can be written as 16,50

Si + 6HF +
$$\frac{n}{2}$$
H₂O₂ \rightarrow H₂SiF₆ + n H₂O + $\frac{4 - n}{2}$ H₂↑
(2) 455

where *n* can be 2, 3, or 4. n = 2 is the valence 2 or current- 456 doubling pathway, which is attributed to the formation of porous 457 Si and porous Si NWs during MACE (or more precisely HL- 458 MACE).⁵¹ When n = 4, the current-quadrupling or valence 4 459 pathway is in action, which can trigger electropolishing⁵² or 460 porosification, based on the model first proposed by Kooij and 461 Vanmaekelbergh⁵³ and explained in detail by Kolasinski et al.³² n 462 = 3 results from a mixture of divalent and tetravalent processes. 463 In HL-MACE, it is believed that etching above the critical 464 current density for porosification occurs in the vicinity of metal 465 particles because the concentration of holes injected into Si is 466 the highest there. However, some of the holes can diffuse further 467 from the metal particles. These holes induce nonlocal or remote 468 etching typically in the current-doubling regime producing 469 porous Si and porous Si NWs. 470

To determine the prevailing pathway, Chartier et al.⁵¹ $_{471}$ suggested using the ratio of HF and H₂O₂ concentrations $_{472}$

$$\rho = \frac{[HF]}{[HF] + [H_2O_2]}$$
(3) 473

They found that for Ag HL-MACE and $\rho \ge 0.8$ the reaction 474 proceeded predominantly according to the valence 2 pathway, 475 while for $\rho \le 0.2$ a valence 4 pathway and electropolishing were 476 observed with no Si NWs formed. In the middle range of $0.2 \le \rho$ 477 ≤ 0.8 , the mixture of pathways was present. This is currently the 478 widely accepted view on the HL-MACE mechanism supported 479 by many studies^{28,33} and comprehensively investigated for p- 480 type Si wafers.³¹

Applying this criterion to LL-MACE, ρ would be equal to 0.92 482 if all 0.037 mol of H₂O₂ were added initially to the etching 483 solution. However, ρ is never this large because during injection 484 H₂O₂ reacts at a rate that is comparable to the rate of injection. 485 This maintains a very low steady-state value of [H₂O₂] 486 throughout etching. Thus, if we were to extrapolate the ρ 487 dependence of etching in the presence of Ag to other metals, the 488 current-doubling pathway should be the dominant one for LL- 489

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Figure 7. (a) Schematic energy band diagrams for Ag and Au LL-MACE of P++, UW, and N++ powders at low (LT, ice bath) and high (HT, water bath at 43 °C) temperatures. Number of injected holes at HT is larger than at LT and is different for Ag and Au. Due to hole accumulation in the metal, the quasi-Fermi level shifts downwards changing the band bending. The holes (h+) are depicted as filled red dots, and the ionized donors ND+ or acceptors NA- in SCLs of N++ and P++ Si are depicted as open circles with a positive or negative sign, respectively. (b) BJH pore size distributions for all metals after LL-MACE of P++, UW, and N++ powders at HT.

490 MACE regardless of the type of metal because we always 491 perform LL-MACE with a value of ρ very close to 1.

As one can see, eq 2 does not include any information specific 492 to the metal catalyst, which our results demonstrate to play an 493 essential role in MACE. When VO_2^+ is used as an oxidant, the 494 chemical identity of the metal and its catalytic activity change the 495 mechanism of etching from the current-doubling pathway for Ag 496 and Au to a current-quadrupling pathway for Pt and a 497 combination of them for Pd.³² Unfortunately, the results for 498 499 H_2O_2 were less distinct, and in ref 32, it was concluded that the 500 effective valence can be affected by many factors including side 501 reactions, temperature, and metal coverage.

t2

Table 2 summarizes the Fermi levels of Si powders used here 502 503 as well as metal work functions, magnitudes, widths, and 504 classification of band bending in the ideal scenario and 505 equilibrium conditions after metal deposition (no H₂O₂ 506 injection). Details of the calculation of these parameters can 507 be found in Section S15, Supporting Information. The diameter of the metal particles was set to 20 nm. A clear difference can be 508 observed for the p-type Si between Cu, Ag and Pd, Pt, Au. Due to 509 510 their low work function, Cu and Ag create a hole depletion space-charge layer (SCL) in the p-type Si, while Pd, Pt, and Au 511 with sufficiently high work function such that $\Phi_M > \Phi_{Si}$ result in 512 hole accumulation and partly degenerate SCLs. Au on P++ 513 powder is a particularly interesting case as there is almost no 514 band bending. This represents an Ohmic contact; it follows a 515 516 linear current-voltage curve and poses no barrier for charge 517 flow. For the n-type Si, all of the metals build up electron 518 depletion SCLs, and metals with high work function may result 519 in inversion layer formation. Note, however, that in real metal/Si 520 contacts the magnitude of band bending is significantly lower ⁵²¹ than in the ideal case used for calculation here.³⁸ Therefore, it is

unlikely that in real samples inversion and degenerate parts of 522 SCLs are formed. 523

Overall, the magnitude of band bending depends on the 524 difference between the metal and Si work functions; the latter in 525 turn depends on the Si doping type and density. Additionally, 526 surface states can appear in the Si/metal interface, and the 527 density of these states may largely affect the Si Fermi level up to 528 the point where it gets pinned.⁵⁴ This, however, should not be 529 the case, as relatively defect-free hydrogen-terminated surfaces 530 form during MACE in HF-based solutions. Finally, H₂O₂ 531 reduction injects holes and adsorbates can inject either electrons 532 or holes into metal particles; holes then accumulate in particles 533 due to an imbalance between hole generation, consumption, 534 and/or diffusion and make the particles charged. Charge 535 accumulation acts as an applied voltage bias and creates a 536 local electric field, which penetrates into Si as it cannot be 537 efficiently screened by the low concentration of charge carriers. 538 A potential drop at the Si/metal interface due to the bias can also 539 be viewed as a decrease of the quasi-Fermi level⁵⁵ in the metal, 540 thus favoring the holes to flow from metal to Si. The applied bias 541 can sufficiently affect band bending and even change the SCL 542 from, e.g., depletion to accumulation or the reverse. 543

Chiappini et al.³¹ showed the difference in porosity for Si 544 NWs after Ag and Au HL-MACE of the p-type Si at various 545 H_2O_2 concentrations. It is widely accepted that porosity is 546 introduced by remote etching, i.e., the nonlocal etching induced 547 by holes further away from their injection site at the metal/Si 548 interface. The tortuous pores in LL-MACE are due to remote 549 etching, and we demonstrate here that remote etching depends 550 heavily on band bending. 551

Herein, we discuss a model that considers the rate of hole 552 generation at the solution/metal interface and the rate of hole 553

554 escape from the metal/Si interface. The balance of these rates 555 determines the balance between local and remote etching. We 556 assume that the rate of escape does not influence the rate of 557 generation. As we shall see, to a first approximation, this model 558 explains the trends found in our data. A further refinement would include the potential for the coupling of these two rates. 559 560 In LL-MACE, we used the same amount of H_2O_2 in all s61 etchings. Thus, we assume that the role of H_2O_2 concentration 562 in increasing carrier density is similar to that of temperature, in 563 that either the increase of temperature or increase of H_2O_2 564 concentration increases the availability of holes.^{33,57} Even 565 though the total amount of injected H₂O₂ is independent of 566 temperature, the rate of injection is higher at higher temper-567 atures because of faster diffusion and possible changes in the ratio of H₂O₂ used in hole injection versus side reactions. We 568 569 further assume that each metal particle accumulates the same 570 number of holes at each temperature independent of the Si doping type. For example, Ag accumulates holes at a certain rate 571 572 at LT for all Si types and at a higher rate at HT, but again the rate 573 is the same for all Si types. Also, Au accumulates holes at a 574 different rate than Ag at LT and at HT, but still, the rate is the same for all Si types. 575

During H_2O_2 reduction, metal particles accumulate a positive charge, which acts as the applied potential and lowers the quasi-Fermi level of the metal. This in turn leads to a decrease of band bending in the presence of H_2O_2 , $E_{BB}^{H_2O_2}$, by the value of eV_a from the ideal equilibrium values, E_{BB} , listed in Table 2

$$_{581} \qquad E_{\rm BB}^{\rm H_2O_2} = E_{\rm BB} + eV_{\rm a} \tag{4}$$

 $_{\rm 582}$ where a potential $V_{\rm a}$ is gained due to accumulation of $N_{\rm h}$ holes $_{\rm 583}$ and can be estimated as $^{\rm 50}$

$$V_{\rm a} = \frac{1}{4\pi\varepsilon_0} \frac{eN_{\rm h}}{r_{\rm M}} \tag{5}$$

⁵⁸⁵ where ε_0 is the dielectric permittivity of vacuum, *e* is the ⁵⁸⁶ elementary charge (positive), and r_M is the radius of the metal ⁵⁸⁷ particle. With temperature increase, more holes accumulate in ⁵⁸⁸ the metal lowering its quasi-Fermi level, increasing applied ⁵⁸⁹ potential V_{a} and decreasing band bending.

We first focus on explaining the differences between Ag and 590 591 Au because these are the two most often used metals in HL-592 MACE, and in LL-MACE, they exhibit clearly distinctive pore 593 size distributions. For P++ Si, Ag and Au at equilibrium create 594 hole depletion and weak accumulation SCLs, respectively (Table 2). Accumulation of holes in the metal always results 595 596 in the decrease of its quasi-Fermi level, thus either making depletion stronger, as for Ag, or creating a weak depletion SCL, 597 as in the case of Au (Figure 7a). The striking difference between 598 <10 nm pore volumes can be observed in Figures 7b and \$9.1-599 S9.3, Supporting Information. In contrast, the volumes of etch 600 601 track pores >10 nm produced by the metal particle movement 602 are nearly identical. Thus, the availability of holes used for the above-critical-current etching near the particles is similar for Ag 603 and Au, whereas the availability of holes that induce remote 604 etching is minimal for Ag. 605

The Ag deposition-induced depletion SCL in P++ Si 607 efficiently pushes the injected holes and holes present due to 608 doping ($N_A = 4 \times 10^{18}$ cm⁻³) away from the metal/Si interface. 609 Part of the injected holes may additionally recombine with the 610 fixed negatively charged acceptors N_A^- in the SCL. The layer is, 611 however, rather thin (~10 nm, Table 2) due to high doping 612 density. Thus, assuming the distance between 20 nm metal particles is on the order of their size or larger,²⁰ there are regions 613 of Si surface that are not affected by hole depletion. Combined 614 with a high abundance of holes in the bulk and electric fields 615 created by charged metal particles, holes can end up near the Si 616 surface away from the SCL and, although not very efficient, 617 nonetheless induce remote etching of tortuous pores. At HT, 618 more holes get injected into the metal than at LT, decreasing the 619 quasi-Fermi energy. This leads to the increase of potential eV_{ay} 620 making depletion even stronger while slightly increasing the 621 width of SCL. On the other hand, correspondingly more holes 622 are injected into Si resulting in a higher volume of tortuous pores 623 at HT than at LT (graphs for Ag in Figure 4). 624

Band bending in P++ Si after Au deposition differs from that 625 induced by Ag (Figure 6a). Since $\Phi_{Au} \sim \Phi_{Si,P++}$, a nearly Ohmic 626 contact is formed with almost flat bands (weak accumulation 627 SCL; Table 2). Thus, there should be no regions of Si where 628 holes are unable to reach the Si surface and participate in remote 629 etching. However, during H₂O₂ injection, a potential eV_a builds 630 up that affects depletion layer formation, and at LT, the volume 631 of pores <10 nm is small (Figures 4 and S9.1, Supporting 632 Information). With the increase of temperature, more holes get 633 injected, and due to much weaker and shorter depletion than in 634 the case of Ag, a lot of tortuous pores are etched (Figures 4 and 635 7).

Properties of SCLs in UW differ considerably from P++ Si. In 637 addition to metal work functions, Si doping density affects two 638 main parameters of the band bending. First, $E_{F,UW}$ is now close to 639 the middle of the band gap, changing the value of $\Phi_{
m M}$ – $\Phi_{
m Si,UW}$ 640 and, therefore, the magnitude of band bending. Second, low 641 dopant density makes the SCL span over several micrometers 642 (Table 2) covering the whole Si surface. Thus, there is no gap 643 between metal particles where an SCL does not exist. Due to $\Phi_{
m Ag}$ 644 $\sim \Phi_{
m Si,UW}$, a weak depletion layer is formed at equilibrium, which 645 becomes stronger during the injection of H_2O_2 as the Ag quasi- 646 Fermi level decreases. However, there are almost no free holes 647 present in bulk UW Si compared to those in P++ Si, and all of the 648 holes injected to Si either diffuse deep into the bulk or 649 participate in local etching in the vicinity of Ag particles, 650 producing only etch track pores (Figures 3 and S11.1, 651 Supporting Information). High temperature makes depletion 652 even stronger by an increase of eV_a . Thus, all holes, which did not 653 participate in local etching, travel away from the Si surface, and 654 no tortuous pores are etched (Figures 4 and 7). 655

The large work function of Au means that $\Phi_{Au} > \Phi_{Si,UW} > \Phi_{Ag'}$ 656 and at equilibrium, Au creates a hole accumulation SCL in UW 657 Si (Figure 7a). It is again spatially large as there are almost no 658 holes from dopants to compensate the negative charge in Au. 659 During H_2O_2 injection at LT, the magnitude of SCL somewhat 660 decreases, which, however, is not enough to fully compensate 661 the band bending. Thus, most of the holes accumulate at the Au/ 662 Si interface and are consumed for local etching of etch track 663 pores (Figures 4 and S11.1, Supporting Information). The 664 situation, however, changes significantly at HT when more holes 665 are injected into Au, further lowering the magnitude of band 666 bending. We assume that eV_a at HT becomes large enough to 667 almost completely eliminate band bending, creating an Ohmic 668 contact or a slight hole depletion layer. This would efficiently 669 inject holes from Au into Si and spread them across the whole Si 670 particle with virtually no barrier to approach the surface to 671 participate in remote etching. We indeed clearly observed a 672 significant increase of tortuous pore volume and specific surface 673 area with the increase of temperature for UW Au LL-MACE 674 (Figures 4 and S11.1–S11.3, Supporting Information). 675

Considering N++ powder, we note that there are extremely 676 $_{\rm 677}\,$ few holes present at equilibrium in Si, and its $E_{\rm F,N++}$ is quite close 678 to the conduction band, making $\Phi_{Si,N++} < \Phi_{Me}$ for all metals. 679 Thus, both Ag and Au create electron depletion layers consisting 680 of fixed ionized donors $N_{\rm D}^{+}$, which are positively charged phosphorus atoms as their electrons diffused into the metal due 681 682 to differences in work functions (Figure 7a). When band 683 bending is strong enough, the intrinsic level E_i can cross $E_{\rm FN++}$. 684 Then, according to the definition of the p-type Si, N++ Si 685 changes from n-type to p-type in the area of SCL, where $E_i >$ 686 $E_{\rm F,N++}$. An inversion layer is formed for all of the metals except 687 Ag in the equilibrium conditions and ideal band bending. We, 688 however, point out again that the formation of an inversion layer 689 on real Si surfaces without the additional applied bias is highly 690 unlikely.³⁸ In most cases, only an electron depletion layer is built up with a width of 8-10 nm instead of ~ 27 nm as listed in Table 691 692 2 for ideal band bending case (but see the discussion of Pd and Pt after Ag and Au). 693

Similar to UW Si and if no inversion layer is formed, only the 694 695 holes available during LL-MACE of N++ Si with Ag and Au are 696 the holes injected first into the metal particles by H_2O_2 697 reduction, which subsequently diffused into Si. Herein, we 698 note the approximately twice higher volume of tortuous pores etched by both Ag and Au in N++ Si compared to that of P++ Si 699 (Figures 4 and 6). We propose that in addition to SCL 700 differences, two other factors play significant roles here. The first 701 one is the charged donors in the N++ electron depletion SCL, 702 which may take part in LL-MACE and at least partially take a 703 704 role of holes to induce etching. Since donors are fixed in the Si 705 lattice, they are not susceptible to diffusion, and it can be shown 706 that all donors are ionized in the SCL.⁵⁴ The second and more 707 important factor is the many orders of magnitude larger hole diffusion current in N++ compared to that in P++. To produce 708 the observed yields, the number of holes injected by H₂O₂ 709 710 dissociation and consumed per Si particle during etching is on the order of 4×10^{10} s⁻¹. This value is slightly higher than $N_{\rm h}$ = 711 712 2.3 \times 10¹⁰ per Si particle in P++ powder and 10¹⁷ times larger 713 than $N_{\rm h} = 1.5 \cdot 10^{-7}$ per Si particle in N++ powder. Thus, in N++ 714 Si due to the large difference between the density of injected 715 holes compared to the equilibrium density of holes, the SCL 716 cannot act as a reservoir to counteract the redistribution of these 717 holes from their site of injection. This leads to efficient hole diffusion away from metal particles and etching of numerous 718 719 tortuous pores despite the recombination of injected holes with 720 bulk electrons.

Pd and Pt are the two metals with the largest work functions 721 722 among the set of metals studied. Both create upward band bending independent of Si type, producing hole accumulation 723 724 layers in the p-type Si and electron depletion layers in the n-type Si (Table 2). $\Phi_{Pt} > \Phi_{Pd}$ and consequently ideal magnitudes of 725 band bending are the highest for Pt. Thus, we cannot exclude the 726 727 formation of an inversion layer with mobile holes in N++ Si. Indeed, we observed slow gas bubble generation for both Pt- and 728 Pd-deposited N++ Si even without H₂O₂ injection, which 729 indicated the presence of slow etching. No gas generation was 730 found for other Si grades and metals after deposition. 731

Pt was highly efficient in the etching of tortuous pores in all Si rag grades with the highest values of v_{pore}/v_{etched} (Figures 4, 5, and rad S12.2, Supporting Information). We attribute this to several rad factors. First, Pt particles tended to aggregate to a smaller rad number of larger chunks (Figures 2 and 3); etching of etch track raz pores by the aggregates was rather local, possibly due to the ras prevalence of the current-quadrupling pathway for Pt MACE.³² www.acsami.org

Second, the high magnitude of band bending resulted in hole 739 accumulation SCL for the p-type Si and buildup of an inversion 740 layer in the n-type Si. Finally, the SCLs are wide enough to span 741 between metal particles attracting holes to the Si surface and 742 inducing remote etching. The only metal that was able to 743 outperform Pt to create higher volumes of tortuous pores and 744 specific surface area was Au and only for UW at HT: 0.038 versus 745 0.106 cm³·g⁻¹ (Figures 4, 5, and 7b and Tables S6.4 and S7.4, 746 Supporting Information). We propose that the Pt-induced band 747 bending favors the injected hole accumulation near the Si 748 surface and makes more holes available for participation in 749 etching than for Au. With temperature increase, this leads to 750 increased local etching, which can be clearly observed by the 751 growth of the average pore size and decrease of v_{pore}/v_{etched} ratio. 752 On the other hand, due to the large distance between aggregated 753 Pt particles, there is more Si without SCL where hole diffusion 754 current brings holes to produce tortuous pores. 755

The behavior of Pd is the most unique. Based on the 756 discussion above, it should produce porosity somewhere 757 between Au and Pt. However, this is not the case. Moreover, 758 25 μ mol of Pd as used for other metals resulted in complete 759 dissolution of Si. Consequently, we used 2.5 μ mol for all of the 760 etchings with Pd presented here, yet still, the outcome of LL- 761 MACE was very different. Electropolishing was observed for 762 UW powder at all temperatures, while for P++ wafer the volume 763 of tortuous pores decreased with increasing temperature (Figure 764 3 and Table S5.2, Supporting Information). For N++ powder, 765 etching was noticed after Pd deposition similar to Pt. Thus, Φ_{Pd} 766 may be high enough to create an inversion SCL in Si. 767 Additionally, Pd itself was found to be highly catalytically active 768 toward etching of Si even without oxidant injection, which was 769 attributed to the effects of dissolved O2.58 Consistent with the 770 concurrent etching by both valence 2 and valence 4 paths of 771 etching reported by Kolasinski et al.,³² Pd seems to catalyze 772 electropolishing instead of tortuous pore formation, especially at 773 HT. 774

Among all of the tested metals, Cu dissolves most readily 775 upon H2O2 injection because it has the lowest reduction 776 potential. For example, Cao et al.⁵⁹ measured nearly 90% Cu 777 mass reduction during the first 60 s of MACE. Therefore, no 778 HL-MACE was reported with Cu to produce ordered arrays of 779 etch track pores and Si NWs, although surface roughening was a 780 common outcome.^{60,61} Furthermore, the band bending induced 781 by Cu is similar to Ag; it is unfavorable for remote etching as a 782 depletion SCL is formed in the p-type Si (Table 2). For UW, 783 only etch track pores were observed analogous to Ag (Figure 4). 784 With temperature increase, the reduction of H2O2 becomes 785 faster and more Cu is dissolved than at LT. We associate the Cu 786 dissolution with a significant change of pore size distributions 787 with temperature for P++ and N++ powders and electro- 788 polishing for UW powder (Figures 4 and 7b) compared to other 789 metals. Therefore, only at LT, a reasonable volume of tortuous 790 pores was etched for P++ Si and etch track pores for UW Si. On 791 the other hand, the yield was 72 and 78% for P++ and UW, 792 respectively, clearly indicating only partial etching of Si particles 793 (Figures 2 and 5). When most of the Cu is dissolved, etching 794 may switch to a more regenerative electroless etching-like 795 mechanism.²⁷ In this case, Cu ions inject holes into Si directly 796 and then get regenerated by H_2O_2 to repeat the process. At HT, 797 the rate of injection increases, resulting in electropolishing, while 798 some etch track pores produced by large Cu particles are still 799 generated. 800

Intermediately doped P+, P, and N+ Si powders show nearly 801 802 the same outcomes as UW Si for each metal (Figure 6). These 803 results support the previous assumption that the injection of 804 holes into Si is independent of Si grade and is determined mainly 805 by the metal. Since the rate of holes that must be injected per Si so particle is on the order of 4×10^{10} s⁻¹, which is much larger than 807 the number of dopant atoms in the particle, there is a large gradient in hole concentration between the metal nanoparticle 808 809 and bulk Si. The large gradient then drives the diffusion of holes ⁸¹⁰ from the metal into the Si bulk, and some of the holes re-emerge 811 near the Si surface to induce remote etching. The appearance of 812 holes far from metal nanoparticles near the Si surface is in turn 813 governed by the type of SCL, magnitude of the electric field in it, 814 and SCL width. For P+, P, UW, and N+ powders, SCL width 815 becomes large enough to leave no gaps between metal 816 nanoparticles (Table 2). Thus, depending on the type of SCL, 817 holes get effectively pushed into the bulk by the electric field (as 818 for Cu and Ag) or stay close to metal nanoparticles (as for Pd, Pt, 819 and Au). Only when there are enough holes accumulated in Au 820 at HT and the magnitude of band bending on the Si/Au 821 interface along with the electric field is decreased, holes are able 822 to diffuse further from metal nanoparticles and tortuous pores 823 start to appear. Pt is the exception here; it produces tortuous 824 pores for all Si grades and at all temperatures (Figures 4 and 6 825 and Section S6, Supporting Information). In our view, this is due 826 to upward band bending in all Si grades and large size of Pt 827 nanoparticle aggregates. The latter results in the parts of Si 828 surface not affected by SCL.

The rate of hole generation in metal nanoparticles, their 829 830 charging, and further diffusion of holes into Si are affected by H_2O_2 concentration near the nanoparticles. During etching, the $_{832}$ H₂O₂ needs to diffuse from the injection site in the solution, thus 833 making the H₂O₂ concentration in the pores unknown. The size of metal nanoparticles and correspondingly the size of etch track 834 pores may therefore play an important role in the transport of 835 $_{836}$ H₂O₂ and other species involved in etching. For example, a few 837 large metal nanoparticles (as in the case of Pt) generate wider 838 etch track pores that facilitate a faster influx of etchant and efflux 839 of etch products. A steady-state concentration gradient may be 840 reached around metal particles in the tortuous pores due to the balance between the reaction and diffusion rates. Concentration 841 gradient profiles can be further affected by, e.g., temperature, 842 metal catalyst, pore size, and depth and contribute to the pore 843 844 morphology of the product.

845 To summarize, LL-MACE resulted in distinctive specific 846 surface areas, yields, and pore volumes depending on the Si 847 grade, metal catalyst, and temperature. Three major factors have 848 been identified that contribute to the distinctive features of LL-849 MACE:

(1) Band bending and space-charge layer (SCL) formation 850 clearly play a significant role in defining the observed 851 outcomes. A distinct difference exists between the highly 852 doped nearly metallic Si and moderately doped Si. The 853 SCL width in relation to the distance between metal 854 nanoparticles is crucial. The SCL is narrow for N++ and P 855 ++ Si, leaving gaps on the Si surface between metal 856 nanoparticles without the electric field. For HL-MACE, it 857 is believed that remote etching is only important in P++ 858 Si. In LL-MACE, we find that the orders of magnitudes 859 larger gradient of hole concentration in N++ facilitate 860 high hole diffusion current and efficient remote etching. 861

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- (2) Chemical, not just electronic, differences between metals 862 lead to distinctive behaviors. Dissolution and redeposition 863 of metal nanoparticles affect the nanoparticle structure 864 during etching, which in turn affects the resulting pore size 865 distributions. We observed peculiar comet-like structures 866 of Cu, Pd, and Pt particles (Figure 2), which 867 unambiguously demonstrates the dynamic nature of 868 metal nanoparticles. The comet-like structure is not 869 required for remote etching because Au never shows a 870 comet-like structure and Pt does not always show it even 871 though they are efficient catalysts of remote etching. The 872 reactivity of Pd is unique, and it must be used at a lower 873 coverage than the other metals to enable porosification. 874
- (3) Mass transport and concentration of H_2O_2 affect the rates 875 of both etching and metal nanoparticle dissolution and are 876 important for morphological evolution. Cu is the most 877 easily dissolving metal, and at high temperature, it 878 catalyzes electropolishing rather than porosification. 879 Nonetheless, a low concentration of H_2O_2 achieved 880 with controlled injection and low temperature enables 881 highly efficient and controllable LL-MACE also with Cu 882 (Figures 3 and 4). 883

CONCLUSIONS

Our comprehensive study has illuminated the mechanisms of 885 MACE and enhanced our ability to exploit it for technical 886 applications. We built upon the recognition of the role of band 887 bending in Si³⁹ in the vicinity of deposited metal nano- 888 particles^{38,62} to gain control over the structures formed by 889</sup> etching. We show that band bending alone is unable to fully 890 describe the observed LL-MACE outcomes. Instead, the 891 chemical identity of metals, their catalytic activity toward the 892 reduction of H₂O₂, the ability to induce different etching 893 pathways, and the tendency to dissolve all influence the results of 894 etching. We demonstrated the generality of LL-MACE and that 895 it can be applied to any kind of Si. Moreover, the judicious 896 choice of the etching parameters (including H₂O₂ concen- 897 tration, metal catalyst, temperature, and the type and density of 898 Si doping) facilitates considerable control of the mean pore size, 899 surface areas, and pore volumes of etched Si powders. The large 900 variety of accessible surface areas, pore sizes, and pore volumes 901 combined with high scalability and economical use of reagents 902 will make LL-MACE highly appealing for use in both research 903 and industry. 904

ASSOCIATED CONTENT	905
Supporting Information	906
The Supporting Information is available free of charge at	907
https://pubs.acs.org/doi/10.1021/acsami.0c13980.	908

Experimental details, description of band bending 909 calculation, and detailed etching data for all Si types 910 including metallurgical-grade Si (PDF) 911

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