Full characterization of ultrathin 5-nm low-k dielectric bilayers: Influence of dopants and surfaces 2 on the mechanical properties 3

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Ultrathin films and multilayers, with controlled thickness down to single atomic layers, are critical for advanced technologies ranging from nanoelectronics to spintronics to quantum devices. However, for thicknesses less than 10 nm, surfaces and dopants contribute significantly to the film properties, which can differ dramatically from that of bulk materials. For amorphous films being developed as low dielectric constant interfaces for nanoelectronics, the presence of surfaces or dopants can soften films and degrade their mechanical performance. Here we use coherent short-wavelength light to fully and nondestructively characterize the mechanical properties of individual films as thin as 5 nm within a bilayer. In general, we find that the mechanical properties depend both on the amount of doping and the presence of surfaces. In very thin (5-nm) silicon carbide bilayers with low hydrogen doping, surface effects induce a substantial softening-by almost an order of magnitude-compared with the same doping in thicker (46-nm) bilayers. These findings are important for informed design of ultrathin films for a host of nano- and quantum technologies, and for improving the switching speed and efficiency of next-generation electronics.

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I. INTRODUCTION

Advanced nanoelectronics, spintronics, and quantum de-28 vices are becoming increasingly three dimensional in de-29 sign, incorporating many layers of sub-10-nm ultrathin films. 30 Moreover, these heterostructures must maintain optimal me-31 chanical properties to avoid device failure. For example, 32 softening due to high hydrogenation (doping with hydrogen) 33 can lead to creep and delamination in semiconductor devices 34 [1,2]. Additionally, as devices push to ever-smaller character-35 istic dimensions, the larger influence of surfaces and inter-36 faces in nanoscale films can change the material properties 37 compared to bulk materials. Depending on the composition of 38 the film, nanoscale thickness effects have been shown to either 39 soften or stiffen ultrathin films [3,4]. 40

One mechanism for introducing a thickness dependence 41 of the elastic properties of ultrathin films arises from the 42 high proportion of atoms at the free surface of the material, 43 which have a reduced number of nearest neighbors compared 44 to atoms in the bulk volume. The low-coordinated surface 45 can either soften the film since surface atoms have fewer 46 constraints on their movement [5], or it can stiffen the film 47 as redistributed electrons induce charging or bond contraction 48 [6,7]. These mechanisms have been studied theoretically us-49 ing continuum and atomistic approaches [5,7-11], and were 50

measured experimentally in materials such as nitrides [12], 51 semiconductors [10,13], polymers [14], and metals [15–17]. 52 A second mechanism that can modify the elastic properties 53 of ultrathin films is the influence of interfaces in multilayers 54 [18]. For example, in few-nanometer-thick Ni/Ta bilayers, 55 while their density ratio is not meaningfully changed from 56 that expected in bulk, we have previously shown their elastic 57 properties are significantly modified-nickel softens while 58 tantalum stiffens, relative to their bulk counterparts [4]. How-59 ever, in this past work we could not extract both of the two 60 elastic constants that fully describe isotropic materials, which 61 are critical to understanding dielectrics for nanoelectronics. 62

Dielectric thin films such as SiC:H and SiOC:H promise 63 optimal electrical properties that are critical for continued 64 scaling of computing power, but struggle to maintain good 65 mechanical properties. To improve the efficiency and switch-66 ing speed of the final device, the dielectric constant, k, of the 67 material between the metallic circuit elements (the interlayer 68 dielectric) needs to be low, below that of the silica used 69 historically (k = 4.2) [1,19]. Methods to lower the dielectric 70 constant include introducing more nonpolar bonds via hydro-71 genation, or introducing pores into the interlayer dielectric. 72 However, the mechanical performance of the film degrades 73 when the network of bonds in the bulk of the film becomes too 74 disrupted, either by high levels of hydrogen bond termination [19–21] or porosity [22]. 76

To measure the elastic properties of such thin films, it is very challenging for most techniques to probe <50-nm

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thicknesses. Widely used techniques such as nanoindentation 79 can characterize films with thicknesses on the order of a 80 fraction of a micron, when combined with advanced modeling 81 [23,24]. Surface Brillouin light scattering, which uses the 82 interaction of light and acoustic phonons, has extracted the 83 full elastic tensor of films of thicknesses down to 25 nm [25]. 84 However, it has difficulty characterizing thinner films without 85 assuming one of the elastic constants. In past work, we used 86 coherent extreme ultraviolet (EUV) beams to characterize the 87 full elastic tensor of isotropic ultrathin films down to 11 nm in thickness [21]. This allowed us to simultaneously extract 89 the Young modulus and Poisson's ratio of low-k amorphous 90 SiC:H films with varying degrees of stiffness and hydrogena-91 tion, in a single measurement. 92

In this work, we show how dopants and surfaces inter-93 play to determine the elastic properties of low-k (k < 4.2) 94 dielectric films that are being developed for next-generation 95 nanoelectronics. We use coherent short-wavelength light to 96 97 fully and nondestructively characterize the mechanical properties of SiOC:H films and SiC:H bilayers with individual 98 layers as thin as 5 nm. This allows us to distinguish between 99 dopant-induced and surface-induced softening. For example, 100 in very thin (5-nm) silicon carbide films with low hydrogen 101 doping, surface effects induce a substantial softening-by al-102 most an order of magnitude-compared with the same doping 103 in thicker (46-nm) films. These findings are important for 104 informed design of ultrathin films for a host of nano- and 105 quantum technologies, and for improving the switching speed 106 and efficiency of next-generation electronics. 107

II. METHODS

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To distinguish between surface-induced softening and 109 dopant-induced softening, we compare two different sample 110 materials: high-hydrogenation amorphous SiC:H and low-111 hydrogenation amorphous SiOC:H. Each sample is fabricated 112 by plasma-enhanced chemical vapor deposition from diluted 113 organosilane precursors on 300-mm-diameter Si (001) by 114 Intel Corp., as described in Refs. [19,26,27]. To describe 115 the number of atomic bonds broken by hydrogenation in 116 each material, we use nominal values of network connectivity 117 (or average atomic coordination number), as determined by 118 Rutherford backscattering and nuclear reaction analysis [28] 119 at Intel Corp. See Supplemental Material for the nominal 120 film properties, including k [29]. As described by topological 121 constraint theory [30,31], an amorphous material transitions 122 from flexible to rigid when the number of constraints on 123 each atom, n, equals the number of degrees of freedom. 124 Accounting for the fixed bond lengths and the fixed bond 125 angles, the relation between network connectivity, $\langle r \rangle$, and 126 constraints, n, is $n = \langle r \rangle / 2 + (2 \langle r \rangle - 3)$. To constrain all 127 three degrees of freedom, $\langle r \rangle$ has a critical value at 2.4, 128 known as the rigidity percolation threshold, where the co-129 ordination is high enough for a rigid network of bonds to 130 percolate through the film volume. Our SiC:H samples have 131 $\langle r \rangle = 3.2$, and thus are rigid, while our SiOC:H samples have 132 been hydrogenated to the critical value of 2.4, which makes 133 films softer and less compressible, as we have previously 134 shown [21]. By characterizing the elastic properties of several 135 thicknesses of both materials, we investigate surface-induced 136

softening both above and below the critical level of hydrogen doping.

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We characterize the elastic properties of ultrathin films 139 using the EUV nanometrology technique described in 140 Refs. [4,21,32]. First, we deposit an array of Ni nanoline 141 grating transducers on each sample using e-beam lithography 142 and liftoff. Grating periods range from 1.5 μ m to 40 nm, 143 as characterized by atomic force microscopy [29,33]. We 144 laser excite these transducers using an ultrafast (30-fs), near-145 infrared (780-nm) pump pulse, as illustrated in Fig. 1(a). 146 The resulting impulsive thermal expansion of the nanolines 147 launches acoustic waves in the nanolines and the film. At early 148 times, a longitudinal acoustic wave propagates down into the 149 sample and reflects from any buried interfaces back to the 150 surface [Fig. 1(b)]. On longer timescales, a surface acoustic 151 wave dominates, with a wavelength defined by the grating 152 period [Fig. 1(c)]. The longitudinal breathing mode of the 153 nanolines is also excited, which we have previously used to 154 characterize the nanolines' elastic properties [4] [Fig. 1(c)]. 155

We measure these dynamic surface deformations by 156 diffracting a time-delayed, ultrafast (10-fs) coherent EUV 157 probe pulse from the surface. The probe has 29-nm wave-158 length (43-eV photon energy), obtained via high-harmonic 159 generation [34]. This photon energy is far from any absorption 160 edges in our sample materials, ensuring minimal sensitiv-161 ity to hot electrons, which can dominate visible-wavelength 162 measurements at the few-picosecond timescales of interest 163 to this work [e.g., Fig. 1(b)]. Our EUV probe provides a 164 direct and sensitive (~pm sensitivity) [35] measurement of 165 the surface acoustic waves and longitudinal acoustic waves 166 in the film. By fitting the acoustic wave velocities using 167 a finite-element analysis (FEA) procedure [21,36–38], we 168 extract the two independent components of the isotropic elas-169 tic tensor of the film: c_{11} and c_{44} , or equivalently Young's 170 modulus and Poisson's ratio (see Supplemental Material) 171 [29,39]. 172

III. RESULTS AND DISCUSSION

Our samples are deposited in two geometries: the SiC:H 174 samples are bilayers (a stack of two identical layers on a 175 Si substrate, see Fig. 2), and the SiOC:H samples are films 176 (a single film on a Si substrate). The two SiC:H bilayer 177 samples compare two different layer thicknesses, 46 and 5 178 nm, and we extract the elastic properties of the topmost laver 179 to check for softening due to the free surface. The SiOC:H 180 films compare three different thicknesses, 44, 19, and 11 181 nm, and have higher hydrogenation than the SiC:H bilayers, 182 as already described. Most importantly, all of the samples 183 we measure are nonporous. This isolates the influence of 184 the single free surface at the top of the film stack. Porous 185 SiOC: H has been shown to have constant elastic properties 186 down to 25-nm thickness [22], but the pores create multiple 187 surfaces throughout the volume of the film, complicating a 188 comparison between surface-induced softening and doping-189 induced softening in that case. 190

To extract the mechanical properties of the ultrathin films, our model requires *a priori* thicknesses and densities for each film. X-ray reflectivity provides these values for the SiOC:H films [29], but it is unable to distinguish the individ-194



FIG. 1. Dynamic EUV diffraction from transverse and longitudinal acoustic waves. (a) After ultrafast laser excitation, the hot Ni nanolines impulsively expand, launching acoustic waves in the sample. After a controlled time delay, an EUV probe pulse diffracts from the sample surface, and the scattered light is collected by a charge-coupled device camera. The acoustic waves dynamically change the EUV diffraction efficiency, as shown in (b) and (c). (b) A longitudinal acoustic wave is launched downward into the film (inset). Reflections from the film-substrate interface imprint a discrete series of echoes in the data (see arrows in inset). (c) At early times, we observe the longitudinal breathing mode (left) of the nanolines. At longer times, we observe a surface acoustic wave (right), whose penetration depth is confined to a fraction of the grating period. The surface acoustic wave and longitudinal acoustic wave velocities provide the two independent components of the isotropic film's elastic tensor.

ual layer thicknesses in the SiC:H bilayers. For the bilayers, 195 we instead utilize scanning transmission electron microscopy 196 (STEM) and energy-dispersive x-ray spectroscopy (EDS) to 197 validate the precise thicknesses and compositions of all the 198 layers we expect in the bilayer samples, as shown in Fig. 2. 199 During sample fabrication, a nitrogen plasma treatment was 200 performed before each SiC:H layer deposition, creating two 201 additional 2-nm N-rich layers [Fig. 2(b)]. Moreover, as STEM 202 was performed after the EUV measurements, we also observe 203

a layer of amorphous carbon on top of the sample, which both the electron and the EUV beams can deposit during the measurement [40]. We also observe strain-induced contrast in both the Si substrate and the lower SiC:H layer, which will be discussed below. 208

To enhance the EUV measurement sensitivity to the influence of a free surface on the elastic properties of the film, we confine the acoustic waves predominantly into the top SiC:H layer. We do this by launching surface acoustic waves 212



FIG. 2. Compositional characterization of the 5-nm SiC:H bilayer. (a) STEM image obtained using a HAADF detector. Strain from the deposited films blurs the atomic contrast peaks in the Si substrate, as expected [41]. HAADF intensity also drops at the interface of the two films in the bilayer. We attribute this to a reduction in the density of the bottom SiC:H layer, as it is strained by the layers above it. (b) EDS image of the sample showing Si (blue) and N (pink). Nitrogen exists at the bottom interfaces of the two SiC:H layers due to the nitrogen plasma clean applied to improve the film adhesion. (c) Horizontally binned lineout of the HAADF contrast for the full cross section taken for STEM characterization. This lineout extends upwards into the focused ion beam (FIB)-deposited Pt layer used for STEM, and downwards into the strained region of the Si substrate.



FIG. 3. Surface-induced softening compared to hydrogenation-induced softening. (a) Elastic constant ranges for the 5-nm SiC:H top layer (red circles), the 46-nm SiC:H top layer (green triangles), and the highly hydrogenated 44-nm SiOC:H film (blue squares). Each point represents a configuration simulated in the FEA model that agrees with the data, within uncertainty. The 46-nm film maintains a bulklike rigid bond network, due to its low hydrogenation and large thickness. The 5-nm film of the same material is significantly softened due to the terminated bonds at its surface (orange). This is distinct from the softening observed in highly hydrogenated SiOC:H, where hydrogenation breaks up the rigid bond network in the volume of the film (blue). Note the top schematics are to illustrate the differences between samples, and are not exact. The auxetic boundary, defined by $c_{11} = 2c_{44}$, is the limit where Poisson's ratio becomes negative. (b) The same data as in (a), expressed in terms of Poisson's ratio and Young's modulus.

with a 40-nm-period grating, which sets their wavelength and 213 confines their penetration depth to $\sim 1/\pi$ of this period 214 [36,37,42]. With most of the elastic energy confined to the 215 topmost layer, we are able to reliably fit the elastic properties 216 without any contribution from spurious effects from the lower 217 layers. We account for the effects of the EUV deposited car-218 bon, N-rich layers, and strain in our FEA model, as described 219 in the Supplemental Material [29,43–45]. 220

As shown in Fig. 3, we observe a strong softening in 221 the low-hydrogenation SiC:H top layer when the thickness is 222 reduced from 46 to 5 nm. This is mainly due to a reduced value 223 of c_{44} , while c_{11} stays approximately constant [Fig. 3(a)]. In 224 terms of Young's modulus and Poisson's ratio [Fig. 3(b)], the 225 5-nm SiC:H layer has a lower Young's modulus but higher 226 Poisson's ratio, i.e., it is softer and more incompressible, 227 like a polymer film. Note that the range of allowed elastic 228 properties defines a nonsquare region (shaded in Fig. 3), after 229 propagating the symmetric experimental uncertainty though 230 the analysis process (see Supplemental Material [29]). While 231 the 46-nm top layer maintains a bulklike rigid bond network, 232 due to its low hydrogenation and large thickness, the 5-nm 233 top layer of the same material is significantly softened due to 234 the terminated bonds at its surface (orange in Fig. 3). This 235

behavior is distinct from the softening observed in highly hydrogenated SiOC:H, where hydrogenation breaks up the rigid bond network in the volume of the film (blue in Fig. 3). This measurement of the 5-nm SiC:H layer represents the full characterization of a < 10 nm film without assuming any of the elastic constants. 230 240 241 241

The results on high-hydrogenation SiOC:H contrast with 242 the results on low-hydrogenation SiC:H. First, the SiOC:H 243 films have been hydrogenated past the critical number of bro-244 ken bonds, meaning that the thickest, 44-nm film is expected 245 to have lower c_{11} and c_{44} values than the SiC:H samples 246 [Fig. 3(a)]. This places the region of allowed elastic constants 247 close to the auxetic boundary, defined by $c_{11} = 2c_{44}$, below 248 which Poisson's ratio becomes negative. This appears as a 249 larger distortion after converting to Young's modulus and 250 Poisson's ratio [Fig. 3(b)]. Comparing different SiOC:H film 251 thicknesses, there is no discernible change in elastic properties 252 since the regions of allowed elastic properties overlap for 253 all three film thicknesses, with the only difference being an 254 increased experimental uncertainty for the thinner films. For 255 clarity, only the thickest SiOC:H film is shown in Fig. 3, but 256 all three films' results are shown in the Supplemental Material 257 [29]. 258



FIG. 4. STEM cross section of 46-nm SiC:H bilayer. (a) Full HAADF image. SiC:H films are known to strain Si substrates, and the full depth of the strained volume is visible here in the blurring of the atomic peaks imaged in the substrate region. We attribute the low-intensity region between the two SiC:H layers to a reduction in density as the top layer and N-rich layer similarly strain the bottom layer. Note the thickness of this strained region is similar to the layer thickness for the 5-nm bilayer in Fig. 2(a), indicating the same effect is lowering the HAADF intensity in the bottom layer of both SiC:H samples. (b) Lineout by horizontally binning (a). The faint intensity straitons within each SiC:H layer are a result of the four-step deposition process. Each step is identical, with no change in film composition.

From the results presented above, we observe no surface-259 induced softening in highly hydrogenated SiOC:H, but we ob-260 serve a significant surface-induced softening in lowly hydro-261 genated SiC:H. Our measurements indicate that this behavior 262 comes from a competition between bond termination at the 263 free surface, and bond termination in the bulk of the film due 264 to hydrogenation. Both of these mechanisms reduce the rigid 265 constraints on atoms, but only so many bonds can be termi-266 nated before a critical threshold is passed and the material 267 loses its rigidity. As described above, the SiOC:H films have 268 surpassed this critical threshold due to hydrogenation alone. 269 Thus, the free surface can only have a minimal effect as the 270 film thickness is reduced. The SiC:H bilayers, however, have 271 less hydrogenation, well below the critical threshold. This 272 allows the top layer to have a greater difference between the 273 rigid bond network in its volume, and the terminated bonds at 274 its free surface. When the layer thickness is reduced, the free 275 surface can then begin to dominate over the otherwise rigid 276 volume of SiC:H, softening the entire top layer. Importantly, 277 the softening we observe in the 5-nm bilayer is not due to 278 oxidation making the SiC:H equivalent to the SiOC:H films. 279 X-ray photoelectron spectroscopy has shown SiC:H to be 280 highly oxidation resistant [46,47], so any oxygen content in 281 the SiC:H bilayer is much lower than in the SiOC:H films and 282 likely is confined only to the outermost atomic layer, as indi-283 cated in Fig. 3 and corroborated by EDS (see Supplemental 284 Material) [29]. Moreover, oxygen is twofold coordinated, so 285 its presence still has the net effect of reducing the surface bond 286 coordination. 287

To confirm the interpretation of our results as a surface softening effect, we perform further STEM and EDS characterizations of the thicker SiC:H bilayer to verify that it is identical in composition to the thinner bilayer. In the STEM cross section of the 46-nm bilayer (see Fig. 4), we observe similar 292 changes in layer contrast as in the 5-nm bilayer. Specifically, 293 we observe a region of reduced high-angle annular dark-field 294 (HAADF) intensity at the top of the lower layer (just below 295 the N-rich layer). Because the SiC:H layers are thicker for 296 this sample, we can see here that the reduced intensity has a 297 finite penetration depth of about 4 nm into the lower layer. 298 This depth is comparable to the layer thickness of the thinner 299 bilayer sample, and so we attribute the intensity reduction in 300 both samples to the top layer straining the bottom layer. We 301 similarly observe a clear strain layer in the Si substrate due to 302 the lower SiC:H layer, as expected for such films on Si [41]. 303 For the EDS measurements, there is no direct sensitivity to 304 hydrogen, but they provide a self-consistent evaluation of the 305 relative amounts of Si, C, O, and N through the depth of each 306 bilayer sample. For both the top and bottom SiC:H layers of 307 both the 5- and 46-nm bilayers, we measure consistent values 308 of ~85% Si, ~12% C, and <2% each of O and N, which 309 is within the noise floor. See the Supplemental Material for 310 full EDS maps of each bilayer [29]. These characterizations 311 together confirm there is no large discrepancy in fabrication 312 between the two bilayer samples, and the primary difference 313 between them is layer thickness. 314

Finally, we rule out alternative explanations for the change 315 in elastic properties between the two SiC:H bilayers. First, 316 a large strain reducing the density of the lower layer cannot 317 explain our results. Reduced density would systematically 318 shift our results toward higher elastic constants (an opposite 319 trend to our observations)-moreover, our fitting procedure is 320 largely insensitive to the density of the lower layer. Second, 321 our observations cannot be explained by the nitrogen plasma 322 treatment. If the treatment was causing a change in the elastic 323 properties of lower layers, then the effect should be even more 324

pronounced when studying a multilayer SiC:H stack with a 325 plasma treatment on each layer. In Ref. [26], the authors 326 perform nanoindentation on exactly this case, with identical 327 SiC:H layers as in our samples. They observe no significant 328 difference in the average elastic properties of the multilayers, 329 both with and without the plasma treatment, down to a layer 330 thickness of 2.6 nm. This indicates buried N-rich layers do 331 not change the overall elastic properties of a stack of SiC:H 332 films. Our technique, however, enables us to isolate only 333 the top layer of our bilayer samples, where the free surface 334 dominates, independent of any plasma treatment on the lower 335 layer. Lastly, we cannot fit our data with nominal SiC:H 336 layer properties while only varying the elastic constants of the 337 N-rich interface layers. 338

IV. CONCLUSION

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We use coherent EUV beams to fully characterize the mechanical properties of films as thin as 5 nm. We find that in the top 5-nm layer of a SiC:H bilayer, surface effects induce a substantial softening—by almost an order of magnitude compared with thicker, 46-nm SiC:H bilayers. This contrasts with SiOC:H films at high hydrogenation levels, which have no significant surface-induced softening, down to 11 nm. We

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attribute this difference between the two sample sets to the 347 competing effects of terminated bonds in the volume of the 348 film due to hydrogenation, and the terminated bonds defining 349 the free surface of the film. For the free surface to change film 350 elastic properties, the surface atoms must be undercoordinated 351 compared to the atoms in the volume of the film. Once hy-352 drogenation terminates enough bonds in the bulk of the film, 353 atoms in the volume and surface of the film no longer have 354 significantly different coordination numbers, and no thickness 355 dependence is observed. These findings are important for 356 informed design of ultrathin, robust films for a host of nano-357 and quantum technologies, and particularly for improving the 358 switching speed and efficiency of next-generation electronics. 359

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