Sculpturing desired shapes in single crystal diamond is ever more crucial in the realization of complex devices for nanophotonics, quantum computing, and quantum optics. The crystallographic orientation dependent wet etch of single crystalline silicon in potassium hydroxide (KOH) allows a range of shapes to be formed and has significant impacts on microelectromechanical systems (MEMS), atomic force microscopy (AFM), and microfluidics. Here, a crystal direction dependent dry etching principle in an inductively coupled plasma reactive ion etcher is presented, which selectively reveals desired crystal planes in monocrystalline diamond by controlling the etching conditions. Using this principle, monolithic diamond nanopillars for magnetometry using nitrogen vacancy centers are fabricated. In these nanopillars, a half-tapering angle up to 21° is achieved, the highest angle reported in the literature, which leads to a high photon efficiency and high mechanical strength of the nanopillar. These results represent the first demonstration of a crystallographic orientation dependent reactive ion etching principle, which opens a new window for shaping specific nanostructures which is at the heart of nanotechnology. It is believed that this principle will prove to be valuable for the structuring and patterning of other single crystal materials as well.

The ability to transform single crystalline materials into desired shapes is vital in nanotechnology. In micro- and nanoscale fabrication, controlling etch direction is essential to achieve the specific shapes in single crystal materials required by device applications. Developing new etching techniques and processes is therefore critical for successful realization of complex devices. Focused ion beam for sculpting specially shaped individual elements[1] and ion beam milling for controlled angle etch[2] are two examples of sculpting techniques that are largely insensitive to crystalline directions. Taking advantage of the anisotropic nature of monolithic materials, crystal direction dependent wet etching techniques have been developed and are most well-known for etching Si in KOH. In a KOH solution, the kinetics of chemical reactions vary on Si {100}, {110}, and {111} planes, leading to a crystallographic dependent etch.[3–9] This wet etch recipe can process large amounts of samples in parallel and is one of the most important assets in modern microelectromechanical systems (MEMS) technology.[10] Having such similar techniques in a dry etch process would be desirable as a tool for more advanced MEMS fabrication. In addition, it would be particularly desirable in nano-electromechanical (NEMS) systems because wet chemical processes can be difficult to control precisely, especially in case of delicate nanoscale devices. Here, we demonstrate that crystal direction dependent etch can be achieved as a dry process on a single crystal diamond.

Diamond is a metastable allotrope of carbon, where the carbon atoms are arranged in a variation of a face-centered cubic crystal structure called a diamond lattice. It has broad applications in science and technology due to its mechanical strength,[11] chemical inertness, thermal properties,[12] and wide-band optical transparency.[13] Diamond further serves as a host material for a variety of atomic defects, some of which show interesting quantum-mechanical spin and optical properties.[14–17] The presence of such atomic color centers has given diamond an important role in quantum computing,[18–21] magnetometry,[17,22–28] and photonics.[13,20,29] For many such applications, optimizing the diamond structure in relation to the color center on the micro- and nanoscales and along particular crystal directions is important.[28,29,31–33]

Developing new etching techniques and processes is a critical step for successfully fabricating devices in diamond. Dry etching processes were developed to make antireflection coating,[34,35] solid immersion lens,[11] nanocavities,[30,36,37] nanopillars,[29,31,33] nanobeams,[28,30,38–42] and microdisk.[35,37] Faceting in diamond was observed in cleaving,[11,43] pure chemical etching in high temperature furnaces filled with O2 gas,[44] and reactive ion etching at high inductively coupled plasma (ICP) power and high substrate temperature.[17] Though {111} crystal planes were observed at zero substrate power, this mechanism remained unexplored. Thus, crystallographic
orientation dependent dry etching has not yet been controlled. In this work, we demonstrate that anisotropic etching along multiple crystal directions in diamond is achieved by controlling the oxygen plasma conditions in an inductively coupled reactive ion etcher (ICP-RIE). Further, we present the underlying etching principle to shed light onto diamond crystal direction dependent etching mechanism. Using this principle, Si-KOH etch is resembled on diamond as an encouraging sign that the principle can be applied to other single crystal materials.

In RIE, etching mechanisms include chemical reactions on exposed surfaces that form volatile byproducts and physical ion bombardments to enhance etch rate and directionality. Major factors controlling etch dynamics include (i) reactive ion flux impinging exposed surfaces, which mainly depends on the concentration of reactive ions in plasma, (ii) the kinetic energy of ions that arrive onto exposed surfaces, which is determined by the negative DC bias between plasma and substrates without considering collisions in the cathode charge region,[45] and (iii) the energy barrier for chemical reactions taking place, which is determined by substrate materials and can be anisotropic in certain single crystals. Under a constant reactive ion flux, the etching process is dominated by either the ion’s kinetic energy or the energy barrier for chemical reactions. Only when the ion energy is closely tuned to this energy barrier and the chemical reaction limits the etching process, does a high etch selectivity along crystallographic directions emerge. This principle is demonstrated in this work.

Figure 1 shows a schematic illustration of the formation of a V-shaped groove in a condition of crystal direction dependent etching. Starting with a rectangular etch window defined on a (100) oriented crystal substrate, if the etch rate in <hkl> direction is slower than that in the vertical direction <100>, tapered {hkl} sidewalls will develop and grow until a V-shape is fully formed.

Throughout this work, ultrapure, monocrystalline, electronic grade (100)-oriented chemical vapor deposition (CVD) synthetic diamond substrates (13C natural abundance, Element Six) are used. The diamond surface is polished by a commonly used method in polishing industry. Atikian et al. showed that 1–2 nm of rms roughness in diamond can be achieved after polishing.[46] The diamond surface is further strain relieved to achieve an rms roughness of less than 1 nm.[46,47] Prior to processing, diamond substrates are cleaned in a boiling mixture consisting of equal parts sulfuric acid, nitric acid, and perchloric acid to remove organic contaminations and to oxygen terminate the surface. To define an etch mask using electron beam lithography, a layer of flowable oxide (FOX) is spin coated on the sample using a 10 nm thick titanium layer as adhesion promoter. On each sample, one group of rectangular etch windows is aligned with its edges parallel to the <110> direction, while another group of windows is aligned to the <100> direction. The crystal orientation of the samples is independently verified in nitrogen vacancy (NV) center magnetometry experiments revealing <111> crystal axis.[48] Etching experiments are conducted in the Plasma-Therm Versaline ICP–RIE system using 900 W ICP power, 40 sccm O₂ flow rate, 10 mTorr pressure, and 10 °C substrate temperature by varying the substrate power from 0 to 120 W. An etch depth of 2–3 µm is achieved for each sample by adjusting the etch duration for each given substrate power. For etch rates at various substrate powers and more details on fabrication, please see the Supporting Information.

As an example of the crystal orientation dependent etch, Figure 2 shows several forms, such as a truncated square pyramid in Figure 2a–c, a V-shaped groove and a truncated rectangular pyramid in Figure 2d. These forms are from the etching masks aligned to the <110> direction and etching at 5 W substrate power for 70 min. The faceted sidewalls belong to the {332} family and have an angle of 25° with respect to [110] vertical planes. Flat etched surfaces and fine straight intersection lines are observed at the corners and between sidewalls and the bottom surface, as shown by the high magnification image in Figure 2e. The visible polishing marks of ≈1 nm rms roughness on the top surface are due to the initial polishing of the diamond and are not a result of the etching process. In contrast, the roughness of the etched sidewalls is not resolved indicating an rms roughness of far less than 1 nm. For the square ring shown in Figure 2a, additional facets appeared around the outside corners, which are symmetric with respect to the <100> diagonals. Their intersections with the bottom (100) surface are close to <740> direction with an average angle of 60.4° to <100>. These corner facets are the result of different ion fluxes and diverted ion trajectories at the corners of the etch window. As shown in Figure 2d, for steady ion bombardment along the long sides of the rectangular features, except at the end corners, only one facet developed.
On the same sample, when the etch masks are aligned parallel to the \( <100> \) direction, the resulting etch profiles are dramatically different. Additional surfaces at inner corners emerged as shown in Figure 3a–d. These surfaces have an orientation very close to \( \{111\} \) but are not flat. Similarly, the etched sidewalls parallel to \( <100> \) are curved and their intersections with the corner surfaces form arc lines. These results imply that the faceting did not fully develop in \( <100> \)-oriented windows at 5 W substrate power. However, when the substrate power was decreased further to 0 W, the faceting along \( \{111\} \) planes at corners and \( \{100\} \) vertical sidewalls did appear as shown in Figure 3e, as indicated by the straight intersection lines and smooth flat etched surfaces. This crystal faceting at zero substrate power was also observed under etching conditions of 3000 W ICP power and 250 °C substrate temperature as has been previously reported.[37]

For etch masks with edges parallel to \( <110> \), the angles between the faceted surfaces etched at different substrate powers and \( \{110\} \) vertical planes were measured with SEM and the respected Miller indices are assigned accordingly (shown in Table 1). These faceted planes have relatively low indices, intersect with the \( <110> \) direction, and rotate around \( <110> \) axis from \( \{111\} \) family to \{331\} family as the substrate power increases from zero to 40 W, as illustrated in the inset of Table 1. These results imply that desired crystal planes can be revealed by varying the substrate power.

In contrast to the low power regime where selectivity is observed, at high substrate powers the etching anisotropy diminished. Figure 3f,g shows the images of a sample etched at 80 W substrate power with windows aligned to \( <110> \) and \( <100> \). The faceting disappears and cavities form at the corners due to heavier ion bombardment. The etching morphologies become identical despite these two differently oriented etching masks. Assuming that ions have no collisions after they move into the cathode charge region above the substrates,[45] the potential energy of ions in the plasma will be entirely transferred to kinetic energy when ions arrive at the substrate surface. Therefore, we find kinetic energy \( \leq 60 \text{ eV} \) (40 W substrate power) to be the threshold to observe crystallographic etching in diamond. This demonstrates that diamond crystallographic etching follows the dry etching principle presented above.

A direct application of this crystallographic orientation dependent etching technique is the fabrication of monolithic nanopillars with large tapering angles. Such waveguiding structures have been shown to increase the photon collection efficiency of color centers in diamond due to a combination of optical wave guiding and adiabatic changes of the effective local refractive index.[33] As shown in Figure 4a–d, tapering angles (half apex angle) of up to 21° are achieved using the presented technique. The taper angle as a function of the substrate power is measured using SEM and shown in Figure 4e. The angle varies linearly with the substrate power when it is \( \geq 40 \text{ W} \) and is discrete at the lower power region. The switching effect indicates that the etching mechanism enters the surface chemical reaction limited regime where the kinetic energy of ions and crystalline bonding energy play major roles. This is consistent...
To verify the waveguide properties of the different nanopillar geometries, the saturated fluorescence intensity of a single NV center implanted 10 nm below the pillars top surface is measured. A group of 10–20 nanopillars containing single NV centers is studied at each taper (half apex angle) angle. All nanopillars shown here have a top diameter of 350 nm and a length of 1.5 μm. Their average saturation count rate and standard deviation are shown in Figure 4f. The results are found to be consistent with previous reports that a larger taper angle yields a higher photon collection efficiency.[33] With the technique presented here, larger taper angles compared to previous studies can be achieved. Beyond higher fluorescence collection efficiency, the larger taper angle of nanopillars strongly enhances the mechanical strength of the structure. This is particularly important in NV center-based scanning probe applications where the diamond nanopillar is scanned in contact over the sample surface.

In summary, the crystal direction dependent reactive ion etching principle is presented for selectively revealing crystal planes in monocrystalline diamond by varying etching conditions in an ICP-RIE system. The physical intuition is to adjust the reactive ion energy to become comparable with the energy barrier for chemical reactions to take place on crystal planes, which leads to crystal direction dependent etch rates. As a direct application of this technique, we demonstrate the fabrication of monolithic diamond nanopillars with tapering angles of up to 21°, which yield high photon collection efficiencies from single NV centers and high mechanical strength. We believe that the etching principle presented here is applicable to other single crystal materials that hold crystallographic anisotropy and in other types of dry etching systems, such as reactive ion beam etch. This method will enable forming a wide range of shapes in different single crystal materials for a broad variety of applications.

### Experimental Section

**Diamond Substrate Preparation:** The electronic grade diamonds (4 × 4 × 0.5 mm³) provided by Element Six were cut and polished by Delaware Diamond Knives. Cleaning the diamonds in a boiling mixture of equal parts of sulfuric, nitric, and perchloric acid was required to remove contamination and impurities. Subsequently, one surface of each diamond substrate was strain relieved using an Ar/Cl and O₂ RIE process. By doing so, a few micrometer of the top diamond surface was removed. This top layer was known to have a large concentration of defects and dislocations accumulated during the polishing process. The smoothness of the diamond surface also benefitted from this strain relief process as shown in a previous report.[43] The strain relief etch parameters are presented in Table S1 of the Supporting Information.
Diamond Fabrication: The diamond was mounted onto a Si carrier chip (1 \times 1 \text{cm}^2) with strain relieved surface facing up using crystal bond for easy handling. This could be done using a hot plate at 150–180 °C to melt crystal bond. To promote the adhesion between e-beam resist and substrate, 10 nm of Ti was evaporated onto the substrate. Three layers of flowable oxide (FOX-16, Dow Corning) were spin coated with each spun at 3000 RPM for 45 s and baked at 100 °C for 10 min. After baking, the FOX layer was about 1 \mu m thick. The FOX layer was then directly exposed with e-beam lithography at 100 keV energy and 5400 \mu C per cm² dosage. The exposed FOX layer was developed in 25 wt% tetramethylammonium hydroxide (TMAH) for 30 s followed by a deionized (DI) water rinse and isopropyl alcohol (IPA) cleaning. This formed the etch mask for the RIE process. First, an Ar/Cl recipe was used to remove the 10 nm Ti layer in the regions not covered by FOX. This exposed the bare diamond surface for the O₂ etch process described in Table S1 in the Supporting Information. After the RIE process, the substrate was dipped in HF to remove residual Ti and FOX.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author.

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Conflict of Interest
The authors declare no conflict of interest.

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crystal orientation, diamonds, nanophotonics, reactive ion etching, single crystals

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