Nano- and Microstructured Copper/Copper Oxide Composites on Laser-Induced Carbon for Enzyme-Free Glucose Sensors

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1. INTRODUCTION

Glucose detection is essential in the healthcare monitoring of people suffering from diabetes mellitus. Increased levels of blood glucose directly relate to the risks in the development of cardiac arrests, renal failure, blindness, and cerebral, and neuronal damage. Globally, the number of people diagnosed with diabetes mellitus is currently estimated at about 500 million by the International Diabetes Federation, and the number is steadily rising, which lays a huge burden on the healthcare system. Thus, direct monitoring of glucose concentration in blood and sweat is essential for timely assistance and prevention.

A new generation of low-cost point-of-care sensors addresses this problem and provides direct analysis of the glucose concentration. Traditional glucose sensors are mostly of enzymatic nature, where a glucose oxidase enzyme is immobilized on a sensing material. The sensing mechanism of such sensors relies on the oxidation of glucose to D-glucuronolactone, the subsequent generation of hydrogen peroxide, and further on to gluconic acid. However, the extraction of the enzymes from fungus, such as industrially accepted Aspergillus niger, is expensive. Furthermore, long-term stability and storage of the enzymatic sensors are of major concern. Therefore, in recent years, significant attention has been paid to enzyme-free biosensors, where several noble metals, metal nanoparticles, and transition-metal oxides have been exploited as the electroactive material for glucose sensing. The use of these materials for biosensing majorly relies on the fabrication of composite electrodes, typically with a carbon-based material such as graphene, carbon nanotubes, and glassy carbon, which further reinforces the essential biocompatibility of the sensing material. Among various electroactive materials, copper and copper oxide exhibit a significant electrocatalytic activity for glucose oxidation, due to its lower overpotential induced by an electron transfer. Moreover, copper oxide nanowires allow an improvement in the sensitivity of detection due to their higher surface-to-volume ratio. Notably, the surface area can be further increased by the growth of nanowires on the spheroids, leading to an urchin-like morphology. Because of their superior morphologies, urchin materials have demonstrated significantly higher sensitivity for glucose detection compared to other copper-based materials. However, the CuO urchin-based glucose...
sensors reported to date were on rigid substrates, used for the
detection of $\text{H}_2\text{O}_2^{35}$ and humidity.\textsuperscript{36} Even though the
fabrication of flexible biosensors has been reported using
bulk copper oxide\textsuperscript{37} and its nanowire variant,\textsuperscript{11,25,26,32,38}
urchin-like morphologies are still unexplored for flexible sensor
devices. Furthermore, current state of the art for the synthesis of
urchin morphologies only includes the hydrothermal
method and some chemical methods.\textsuperscript{35,36} Here, we focus on
the fabrication of a CuO urchin-based flexible glucose sensor,
where the urchin structures are achieved solely by thermal
annealing, together with a straightforward variation of the
working environment.

As a promising method for the development of a new
 generation of economic point-of-care devices, laser-induced
carbonization (LIC) has manifested itself as a rapid and low-
cost fabrication method.\textsuperscript{35,40} The fabrication process of LIC
utilizes localized laser irradiation for immediate conversion of
the polymer substrate to a conductive film. This process works
for a wide range of possible wavelengths.\textsuperscript{39–41} The laser carbon
(LC) is induced by a combination of photochemical and
photothermal mechanisms,\textsuperscript{39,42} forming conductive, highly porous,
highly graphitized, carbon-rich films, which are embeded directly in
the polymer precursor.\textsuperscript{40} A number of high-temperature thermosets can be used as precursors for
LIC, including polyaramid Nomex.\textsuperscript{43,44} Laser carbon derived from Nomex exhibits a superior conductivity ($\approx 45$ S cm$^{-1}$) compared to other precursors converted with 10.6 $\mu$m wavelength laser radiation.\textsuperscript{44} LIC has revealed itself as a suitable process for the fabrication of chemical sensors,\textsuperscript{45–47} and biological sensors,\textsuperscript{48,49} due to its high surface activity. These advantages of LIC allow accurate detection with a low concentration of such biological metabolites as urea,\textsuperscript{48} dopamine,\textsuperscript{50} $\text{H}_2\text{O}_2$,\textsuperscript{51} thrombin,\textsuperscript{52} and glucose.\textsuperscript{45,50,53}

Here, we report on the fabrication of copper microspheres
(Cu-MS) and copper oxide urchin (Cu-O-U)-based flexible
bioelectrosensors. The sensor electrodes are fabricated by
laser-induced carbonization of flexible Nomex sheets, followed
by electroplating of copper, high-temperature annealing, and
preferential oxidation toward obtaining either Cu-MS or CuO-
U. We study the effect of different processing parameters to
understand the formation of both materials. We characterize
the material properties using scanning electron microscopy
(SEM), X-ray diffractometry (XRD), transmission electron
microscopy (TEM), and X-ray photoelectron spectroscopy
(XPS). We then present the electrochemical performances of these
materials with potentiostatic cyclic voltammetry (CV)
and use them for the amperometric glucose detection with
real-time amperometry. We end with a discussion on the
perspective of our electrode materials in point-of-care
applications.

2. MATERIALS AND METHODS

2.1. Fabrication. The fabrication of the electroplated film on laser
carbon was derived from Mamleyev et al.\textsuperscript{45} Poly(1,3-phenylene
isophthalamide), commercially known as Nomex Type 410 insulation
paper (DuPont; supplied by RS Components), of thickness 130 ± 13
$\mu$m, was rinsed with isopropanol and dried for 10 min naturally. The
films were fixed on glass slides with acrylic adhesive films (3M-
467MP) to prevent the films from warping during fabrication. LIC on
Nomex was carried out with a CO$_2$ laser (ULS Versa Laser 3.50,
wavelength: 10.6 $\mu$m, lens 2.0 in., beam diameter 120 $\mu$m at focus).
The fabrication was performed at a 3 W power, a 13 cm s$^{-1}$
scanning speed, and a focal distance of 2.5 mm. The pulse throughput was
controlled by pulses per inch (PPI) and was set to 1000 along and
perpendicular to the rasterization direction. During consecutive
passes, the perpendicular PPI was reduced to 750, and during the
third pass, the speed was increased to 32 cm s$^{-1}$. The supporting
electrode for bent amperometric tests was fabricated using only two
passes with 750 PPI and scan speeds of 13 and 32 cm s$^{-1}$, which
imparted flexibility to the carbonized films.

The copper electroplating was conducted in a custom-built
electrolyte bath system (Cuprostar LP-1, Enthone-OMIGmbH) with
$\text{H}_2\text{SO}_4$ (2 M), $\text{H}_2\text{O}_2$ ($7 \times 10^{-4}$ M), and $\text{HCl}$ ($7 \times 10^{-4}$ M), following
the manufacturer’s specification for promoter additives. The bath was
constantly stirred by an electric pump. A direct current of 30 mA was applied during the plating, and the deposition was
performed for 600 s. The electrode exposed to the electrolyte had a
square shape with a 15 mm side length. The supporting electrode for
bend tests was electroplated under the same current density.
However, the plating time was increased to 3000 s due to the
enlargement of the total area to 3.5 cm$^2$ and an increase in resistance
for carbon films prepared during the double laser passes.

The annealing was performed in a horizontal tube furnace
(Carbolute Gero FHA 13) under a constant nitrogen flow ($\approx 0.8$ L
min$^{-1}$) for 1 h, with a heating rate of 5 °C min$^{-1}$. To investigate the
phenomenon of folding into spheroids, annealing was conducted at
800, 900, 1000, and 1100 °C, followed by natural cooling under
$\text{N}_2$ flow. For the growth of the CuO-U, we introduced air after 450 °C
during the cooling step, following a specific regime for nanowire
growth.\textsuperscript{54,55} Both types of the annealed samples fabricated at 1000 °C
were selected for the glucose sensing and used for the amperometric
studies.

2.2. Characterization. The surface morphology of each film was studied with a Carl Zeiss AG SUPRA 60VP SEM equipped with
a Bruker X-FLASH 5010 energy-dispersive X-ray (EDX) spectroscopy
detector. XRD was conducted on a Bruker D8 Advance diffractometer in a $\theta-\theta$ geometry using Cu K$_\alpha$$_2$ radiation. Bragg’s formula was used
for the determination of the interplane distances. Rietveld refinement
was employed for the determination of the crystallite sizes and the
internal stresses.\textsuperscript{56} TEM was carried out on a Thermo Fisher
Scientific Themis300 operated at 300 kV, with selected area electron
diffraction (SAED) patterns collected at a camera length of 600 mm
on a bottom-mounted Ceta 16M camera from studied specimens.
Diffraction patterns were evaluated using the “PASAD-plugin” of
the Gatan Digital Micrograph software. Laser carbon and Cu-O-U
dispersions were dispersed in ethanol and ultrasonicated for 1 h, and then
deposited on Quantifoil Holey Carbon Film supports with circular
holes, type R1.2/1.3 $+ 2$ nm additional continuous C layer on Au grid
with 200 mesh. XPS was performed using an Al K$_\alpha$ radiation source ($E = 1486.6$
$eV$), and the spectra were collected with the hemispherical energy
analyzer RG Scienta 4000. The survey and individual spectra were
collected with 1 and 0.05 eV increments, respectively. The spectra
were analyzed using the CasaXPS software; all spectra were fit with a
Gaussian–Lorentzian function and a Shirley background, excluding
Cu 2p spectra, which were fit with a spline Tougaard background.
The built-up charge accumulation resulted in the shift of the binding
energy in all spectra, which was corrected by adjusting to the center
adventitious carbon centered at 284.6 eV.

CV was performed on a modular AutoLab PGSTAT128N
potentiostat/galvanostat equipped with NOVA software.
The measurements were carried out in a three-electrode system: a working
electrode, a platinum counter electrode, and a Ag/AgCl reference
electrode immersed in 1 M KCl solution. All experiments were
performed at 1000 mV s$^{-1}$ with the presence and absence of a 1 mM glucose solution, within a potential range between
$-0.8$ and 0.8 V for the determination of the characteristic redox
potentials, and between 0 and 0.8 V to study only the oxidation
behavior. The selected electrolyte provided a charge transfer during
the glucose sensing and showed some influence of the concentration

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of NaOH in the amperometric signal, but without any significant contribution above 100 mM, which was chosen as a standard concentration for the solution for the electrochemical tests.\textsuperscript{11}

Amperometric glucose detection was carried out using the same setup, with the potential fixed to 0.5 V. The detection was performed in a 100 mM NaOH electrolyte with separate glucose administration with increasing concentration 1-fold in the whole volume after three consequent injections of same molarity starting from 10 nM and ending at 10 mM injections. Each injection had a duration of \( \sim 10 \text{ s} \) with a 100 s interval between injections; after injection of 10 mM, the interval was increased to 200 s due to the saturation current rise at a stable and reproducible level. Determination of the specific selectivity to glucose was conducted by adding 1 mM of the following chemicals: \( \beta \)-glucose, sucrose, urea (all supplied by Merck), acetic acid (50%, supplied by VWR), and uric acid (supplied by Sigma-Aldrich). The sensitivity was derived from the slopes of the fit curves, and the limit of detection (LOD) was calculated using the formula

\[
\text{LOD} = \frac{3.3 \times \text{sd}}{s},
\]

where \( \text{sd} \) is the standard deviation of the signal in a single step and \( s \) is the sensitivity.\textsuperscript{58}

3. RESULTS

The fabrication of the glucose sensor is illustrated in Figure 1. Flexible Nomex insulation sheets were irradiated with an infrared laser, leading to the formation of a carbon-rich conductive seed layer. The carbon seed layers on the Nomex sheets were used for subsequent electroplating of copper. The copper/carbon composite films were then annealed in a tube furnace under a nitrogen atmosphere. During the cooling step of the heat treatment process, two different approaches were implemented to form either copper microspheroids (Cu-MS) or copper oxide urchins (CuO-U). These materials further adhered to a copper electrode that was electroplated onto a laser carbon/Nomex substrate to fabricate the new electrodes.

Figure 1. Schematic representation of the process flow. (I) Nomex paper attached to a glass slide; (II) fabricated electrode patterned with IR laser; (III) electrode electroplated with Cu; (IV) detached substrate annealed in nitrogen (for Cu-MS) and nitrogen/air (for CuO-U) working environments; and (V) CuO-U film attached to the plated contact pad and tested bent multiple times.

Figure 2. SEM images of (A) an initial laser carbon film, (B) a copper plated on laser carbon, and composites (C) Cu-MS and (D) CuO-U, (E, F) with a magnification on the copper nanowires.
Figure 2A shows a scanning electron microscopy (SEM) image of the laser carbon film, which featured a porous morphology with a pore size ranging from ~50 nm to ~10 μm. The electroplated copper film exhibited a morphology featuring elongated crystals ~3.5 μm in length and ~1 μm in width, centered at the adhesion site and uniformly distributed across the precursor surface (Figure 2B). The annealing and cooling in the nitrogen environment resulted in aggregation into bigger copper clusters, partially melted and aggregated into microspheroids and nanoparticles (Figure 2C). The shape of such agglomerates, the microspheroids’ diameters, their quantity, and uniformity of distribution on laser carbon were directly proportional to the plating duration (Figure S1 and Table S1). However, a plating time higher than 10 min resulted in the formation of asymmetrical agglomerates, yielding random shapes as shown in Figure S1D. At an annealing temperature below 1000 °C, the plated film did not form any microspheroids, rather, it partially melted and agglomerated into bigger crystals, decreasing the roughness and areal coverage on the underlying laser carbon (Figure S2). Introduction of air at 450 °C during the cooling step yielded shell oxidation and growth of copper oxide nanowires on Cu-MS, resulting in urchin-like microstructure (Figures 2D and S3) with a small increase in the diameter of the microspheroids (Tables S2 and S3). The morphology was uniform throughout the surface. The CuO-U obtained from a precursor sample with a plating time above 10 min resulted in a fracturing of the oxidized shell of the microspheroids (Figure S3D). This was caused by excessive stress release, resulting in deterioration of their adhesion to the laser carbon substrate and negatively affecting the sensing capability of the composite electrode. The densely grown nanowires featured diameters ranging from 15 to 150 nm, with lengths up to 5 μm as shown in Figure 2E,F. For both types of films, the samples obtained for annealing at 1000 °C exhibited the highest amperometric response without any significant deterioration of the signal due to saturation (Figure S4). Therefore, these samples were selected for further detailed studies.

Figure 3A depicts X-ray diffractograms of the studied films. Laser carbon features peaks at 25.8, 42.9, and 53.4°, which are characteristic of a (002) reflection, originating from the perpendicular arrangement of the basal planes in graphite, (100) reflection corresponding to the hexagonal planes, and (004) of second order for the basal plane reflections, respectively.59,60 The interplane distances determined using Bragg’s law for the (002) and (100) peaks are 3.45 and 2.10 Å, respectively, and listed in Table 1. The XRD pattern of the electroplated copper featured characteristic peaks of Cu at 43.3, 50.4, and 74.1°, which correspond to (111), (200), and (220) reflections, and attributed to JCPDS #04-0836. Cu-MS also exhibited similar peaks. However, the intensity of all peaks dropped slightly, indicating a reduction in the total acquired signal. CuO-U films showed a significant reduction in Cu reflections. In addition, peaks corresponding to Cu2O and CuO appeared in the diffractogram, which matched to JCPDS
The presence of both chemical modifications confirmed a phase transition of copper to an oxidized phase. The interplane distances were calculated using Bragg’s law, and the crystallite sizes and the internal stresses for each component were derived after using Rietveld refinement, and the results are listed in Table 1. Figures 3B,C and S5 show transmission electron microscopy (TEM) micrographs of laser carbon, featuring thick stacks of graphene sheets and the absence of a long-range order characteristic of polymer-derived carbons. Additionally, the selected area electron diffraction (SAED) patterns were collected from the same sample and are depicted in Figure 3D, which exhibitsdiffusion rings with high-order diffraction patterns previously not detected with XRD. Figures 3E–G and S6 show TEM images from CuO-U nanowires and neighboring areas, featuring a long-range order and presence of two phases: cubic Cu2O at the Urchin with a unit cell determined at 4.60 ± 0.34 Å, and monoclinic CuO with a lattice distance of 2.53 Å calculated for the (111) plane. Figure 3H shows SAED collected from a single nanowire with the electron beam focused along the [1̅10] orientation with the reflected planes marked accordingly. Raman spectroscopy

Figure 4. Detailed XPS spectra of C 1s, O 1s, N 1s, and Cu 2p for (A–C) laser carbon, (D–F) after electroplating of Cu, and the composites (G–I) Cu-MS and (J–L) CuO-U on LC.
Figure 5. Cyclic voltammograms performed in 100 mM NaOH and 1 mM glucose with variation of the scanning rate on (A) Cu-MS and (B) CuO-U. (C) Oxidation current density at 0.5 V on the speed of scanning rate. (D) Comparison of the cyclic voltammograms between different substrates performed at a 20 m V s⁻¹ scan rate in 100 mM NaOH electrolyte, with and without 1 mM glucose.

Further conﬁrmed the formation of laser carbon and CuO (Figure S7). The Raman spectrum of laser carbon featured the characteristic first-order peaks for the D-band at 1346 cm⁻¹, arising from an A₁g breathing mode of the hexagonal rings at the K-point initiated by disorder, and for the G-band at 1593 cm⁻¹ from an E₂g mode at the Γ-point, responsible for the stretching vibrations in carbon.⁶²,⁶³ In addition, the second-order 2D overtone at 2683 cm⁻¹ from sp² carbons and a D + G combinational peak at 2931 cm⁻¹ can also be observed. The calculated intensity ratios I_G/I_D and I_2D/I_G were 0.68 and 0.21, respectively. Figure S7B shows the Raman spectrum of the CuO-U ﬁlm, which featured three characteristic peaks of CuO (inset of Figure S7B) along with the peaks of the laser carbon. The bands in LC shifted insigniﬁcantly from the reference values, and the I_G/I_D and I_2D/I_G ratios were 0.69 and 0.25, respectively, suggesting improvement in the graphitization of the carbon materials. The CuO peaks originate from 3 Raman active modes, out of 11 possible optical modes, which correspond to A₁g at 294 cm⁻¹ and two B₁g modes at 341 and 627 cm⁻¹.⁶²,⁶⁴ The positions of these bands matched with Raman shift toward the lower wavenumbers for nanocrystalline powders,⁶⁴ which further conﬁrmed the crystallite sizes determined with the XRD.

Surface chemical compositions of the fabricated ﬁlms were studied using X-ray photoelectron spectroscopy (XPS). The survey spectrum is shown in Figure S8, which features the presence of C 1s, O 1s, and N 1s peaks in laser carbon, and a peak of Cu 2p was observed after plating and annealing. A few contaminants were also detected on the surface. For example, peaks of Na 1s and Auger Na_KLL were observed in the XPS spectra of the laser carbon sample, which might have originated from the manufacturing process of the Nomex sheets. The spectra for copper-coated ﬁlms featured visible peaks of F 1s, S 2p, and Si 2p, which might have originated from the electroplating bath and precipitation of silicon nanoparticles during dicing of the glass slide. After the annealing procedures, the presence of these contaminants was signiﬁcantly reduced. However, the spectra of the annealed samples featured insigniﬁcant contributions from Al 2p, which probably originated from sample handling. Table S4 indicates total atomic composition corrected to the elements of interest for each studied ﬁlm. Figure 4 presents deconvoluted C 1s and O 1s XPS spectra of all of the four types of samples. Deconvoluted N 1s spectrum was collected only for laser carbon, due to its signiﬁcant contribution, whereas Cu 2p spectra were collected from the samples obtained after plating and annealing.

The C 1s peaks of the XPS spectra of each sample (Figure 4A,D,G,J) are deconvoluted into six species: C=C, C−C/C=H at 284.6 eV, C−N at 285.6 eV, a hydroxyl (C−OH) at 286.1 eV, a carbonyl (C=O) at 287.4 eV, a carboxyl (HO−C=O) at 288.8 eV, and π−π* shake-up at 291.0 eV.⁴⁰−⁴⁸,⁶⁶−⁶⁷

The ﬁtting was reduced to ﬁve species of both samples obtained after annealing (Figure 4GJ), and the C−N species was removed due to insigniﬁcant contribution from the N 1s peak. Figure 4B,E,H,K shows the O 1s peak, which featured
three species in all samples, including an aromatic carbonyl at 530.9 eV, an aliphatic carbonyl at 532.3 eV, and a hydroxyl at 534.1 eV. The fitting for laser carbon was performed with an additional species from amide at 531.9 eV and overlapping with some minor contamination originated from a hydroxyl group, chemisorbed water, and possible Auger NaKLL, which were fitted in one peak at 536.0 eV. The O 1s peak (Figure 4E,H,K) after deposition of copper featured two supplementary species, which were a cupric oxide (CuO) at 529.7 eV and a cuprous oxide (Cu2O) at 530.3 eV. Figure 4C shows the deconvolution of the N 1s peak into four nitrogen-containing species: a pyridinic (N-6) at 398.80 eV, a pyrrolic (N-5) at 399.8 eV, a quaternary/graphitic (N-Q) at 401.8 eV, and an oxidic (N-Ox) at 404.2 eV. Figure 4F,I,L shows Cu 2p peaks, which were fitted into the five species: a copper, a cuprous oxide, a cupric oxide, a copper hydroxide (CuOOH), and a copper sulfate (CuSO4). The contribution from copper sulfate after annealing was negligible and was therefore omitted in the further analysis. The detailed results after the deconvolution of all spectra are presented in Table S5.

The electrochemical properties of the annealed films (Cu-MS and CuO-U) were investigated using cyclic voltammetry (CV). The redox potentials were observed within the potential range −0.8 to 0.8 V for both samples, as shown in Figure S9. The electrochemical measurements were independently performed in the initial alkaline electrolyte, in the presence and absence of 1 mM glucose and independently with sucrose. Figure S9A shows cyclic voltammograms of the Cu-MS films. The oxidation peaks were observed at −0.32 and −0.10 V, which can be attributed to the transition from Cu0 to Cu1+ (CuO) and from Cu1+ to Cu2+ (CuO), respectively. The reduction was observed at −0.54 and −0.74 V, which correspond to the transition from Cu2+ to Cu1+ and then to Cu0. The addition of glucose to the electrolyte resulted in a decrease in the intensities of all of the peaks, and a shift of the oxidation peak to −0.23 V, which was due to the transition from Cu0 to Cu1+. Figure S9B shows CV curves of the CuO-U film. Notably, the oxidation peaks were observed at −0.02 and 0.05 V, which were attributed to the transition from Cu1+ to Cu2+ and from Cu2+ to Cu3+, respectively. The reduction current featured a peak at −0.60 V, which were due to the transition from Cu3+ to Cu2+ and from Cu3+ to Cu1+, respectively. The presence of glucose significantly improved the oxidation and reduction of Cu2+. The chemical formulas for the involved reactions are presented in Figure S10A.

As the glucose molecules mainly yielded an oxidative behavior on the electrode materials, we studied the oxidation current in detail to characterize the sensing behavior. The cyclic voltammograms of Cu-MS and CuO-U electrodes are presented in Figure 5A,B, respectively. The oxidation peak current increased with an increase in the scanning rate for both the electrodes, due to the fast oxygen transfer to the working electrode promoting an uncompensated solution ohmic drop. Figure 5C shows an increase of the oxidation current proportional to the square root of the scanning rate and shows a linear dependence, indicating only diffusive behavior in the electrocatalytic glucose oxidation, which is advantageous for amperometric sensing. However, Cu-MS film exhibited a steeper slope, supporting the improved oxidation rate from Cu0 compared to the CuO counterpart. As depicted in Figure 5D, the electrochemical studies showed that Cu-MS and CuO-

Figure 6. (A) Temporal amperometric response by a stepwise increase of the glucose concentration in a 100 mM NaOH electrolyte. (B) The extracted calibration curves of a current response versus the glucose concentration (same color map as for (A)). (C) Amperometric response of the composite films by a consecutive addition of 1 mM of each of the indicated analytes. (D) Amperometric signal reduction after multiple uses.
U films exhibited significantly superior performance while compared to laser carbon and the supporting copper tape. With the presence of 1 mM glucose in the electrolyte, the total area of the CV curve and the peak current increased for all of the studied films.

Figure 6A shows the real-time amperometric response of the Cu-MS and Cu-O-U electrodes, after consequent injection of glucose of different molarity starting from 10 nM to 10 mM in the 100 mM NaOH electrolyte at 0.5 V. The potential at 0.5 V was selected because a significant improvement of the current response in the CV was observed for the CuO film in the presence of glucose (Figure 5D). For comparison, the amperometric response of the laser carbon film and the supporting copper tape is also presented, each of which exhibited only a minor contribution to the total signal during the amperometric measurement. The calibration curves for both test samples are plotted in Figure 6B and follow a linear dependency on the glucose concentration up to 3 mM. Notably, the Cu-U electrode exhibited a more uniform step-like signal at a low concentration and featured a saturation of the current density at a glucose concentration of over 10 mM. The linear range of the amperometric response for both films was fit from 1 μM to 3 mM, with an R² of 0.9996 and 0.9982 for the CuO-U and Cu-MS, respectively. The nonlinear (hyperbolic) function as well can be used for fitting improvement for concentrations starting from minimal at 1 μM and allows extension of the sensitivity range to 10 mM. The sensitivity of the Cu-O-U and Cu-MS electrodes was 0.25 and 0.32 mA cm⁻² mM⁻¹, respectively, whereas the lower limits of detection (LODs) of these electrodes were 1.74 and 7.56 μM, respectively. Some increase in amperometric signal was observed after injecting 100 mM glucose, but because the high signal-to-noise ratio cannot reliably derive, all fits were initiated from 1 mM. The comparison to other reported values for carbon-based sensors is presented in Table 2. Figure 6C shows the selective current response of our electrode materials upon addition of 1 mM glucose, sucrose, acetic acid, urea, and uric acid, finalized with a control glucose addition. Cu-MS were additionally verified by a complementary addition of sucrose and glucose due to the observed response in the presence of the sucrose. In comparison, the Cu-O-U electrode exhibited specific selectivity only to glucose.

| electrode                  | detection potential (V) | sensitivity (mA cm⁻² mM⁻¹) | linear range (mM) | LOD (μM) | ref          |
|----------------------------|-------------------------|-----------------------------|-------------------|----------|--------------|
| Cu-MS/LC                   | 0.50                    | 0.25                        | 0.001–3.3         | 1.75     | this work    |
| CuO-U/LC                   | 0.50                    | 0.32                        | 0.001–3.3         | 7.56     | this work    |
| CuO/graphene               | 0.55                    | 1.36                        | 0.002–4           | 0.70     | 24           |
| Cu nanowire/graphene       | 0.60                    | 1.10                        | 0.005–6           | 1.60     | 25           |
| CuO/MWCNT                  | 0.55                    | 2.19                        | 0.001–1           | 0.80     | 26           |
| CuO NW/SWCNT               | 0.60                    | 2.19                        | 0.001–0.034       | 0.05     | 27           |
| Nafton/CuO/GC              | 0.60                    | 0.41                        | 0.05–2.55         | 1.00     | 28           |
| Cu/MnO₂/GC                 | 0.70                    | 0.38                        | 0.25–1.02         | 0.10     | 29           |
| Pt–NiO/rGO/GC              | 0.50                    | 0.83                        | 0.008–14.5        | 2.67     | 30           |
| Ni–SnO₂/PANI/CuO           | 0.65                    | 1.63; 1.33                  | 0.001–1; 1–10     | 0.13     | 37           |

The sensitivity of the CuO-U and Cu-MS electrodes was 0.25 μA cm⁻², which can be attributed to the oxidation of Cu²⁺ to Cu³⁺ (CuOOH). Such an oxidation reaction leads to the hydrolysis of sucrose to fructose and glucose in an alkaline solution. The glucose converts to a gluconolactone, and further, a gluconic acid, as shown in Figure S10B. Figure S11 shows the detailed electrochemical analysis of the Cu-MS electrode in the presence of the sucrose. The amperometry on the Cu-MS was performed by consecutive addition of sucrose. The results are presented in Figure S11C, which exhibited deterioration of the signal with an increase in concentration. This resulted in a shorter linear range from 1 to 30 μM with R² = 0.969.

The CuO-U film exhibited the presence of the majority Cu²⁺ oxide species and shows its retention after the amperometry tests, without any chemical modification, as was observed earlier via CV tests in Figures 5A and S9B. Due to this factor, the CuO-U film was selected for the test involving multiple repetitions, followed by rinsing in DI water after each cycle, and for the complementary bending tests of the composite electrode. Figure 6D shows retention of the amperometric response after multiple repetitions, whereas the signal shows a deviation of around 15% after five cycles.

Such a low-cost glucose sensor was additionally tested after flexing at ±90° and is presented in Figure S12. However, it suffered from significant deterioration of the signal at a low concentration and exhibited signal response from 4 to ~60 mM, which was fit with linear function, attributing reduction of the sensitivity to 0.072 mA cm⁻² mM⁻¹. All of these changes can be attributed to the reduction of the sensing area and the poor electrical connection between the electrode and electrodeposited copper. Further improvement for flexible sensors can be performed by implementing conductive adhesive or providing additional plating on the back side of the electrode, which lays out of the scope of the current paper.

The surface morphology and chemistry after the last amperometric measurement were studied with the SEM depicted in Figure S13 and the EDX in Figure S14, respectively. The population of Cu-MS and Cu-O-U nanoparticles decreased ~7 and ~15%, respectively. Notably, the nanometric features, nanowires, and particles from both substrates almost completely detached after amperometric testing. Due to such reduction, the chemical composition of copper significantly decreased in both cases, as depicted in Figure S15. The substrate does not feature significant oxidation and most of the oxygen is bonded with the microparticles. Figure S15B features estimation of the copper to oxygen ratio, showing an increase in the oxidation in Cu-MS and a decrease in Cu-O-U. The latter corresponds to the nanowire detachment and the exposition of the Cu₂O core below the Cu shell of the urchin.
4. DISCUSSION

Laser-induced carbonization is a rapid process for the thermochemical conversion of a polymer to a carbon material, which is induced by photochemical and photothermal mechanisms analogous to conventional thermal pyrolysis in a furnace. The accumulated heat over the polymer surface provides a sufficiently high-temperature gradient for carbonization at a focal spot and a surrounding heat-affected zone, involving a rapid expulsion of byproducts such as CNH, CO, CO₂, H₂O, and complex aromatic hydrocarbons. It has been reported that Nomex films exhibit a high carbon yield for LIC and require multiple passes of lasing for improvement of crystallinity and conductivity. Lattice parameters of the Nomex-derived laser carbon indicate a high degree of graphitization with an interlayer distance and crystalline sizes comparable to commercial graphite and superior to that obtained from the traditional furnace-based pyrolysis of Nomex. The XRD pattern of the Nomex-derived laser carbon (Figure 3A) exhibited a high degree of graphitization with a crystalline size of 56 nm. SAED, collected from laser carbon, confirmed the high graphitization by the appearance of high-order diffraction patterns, with families of {211} and {214} planes, which were not observed with XRD. TEM imaging supported the presence of thick stacks of long graphene sheets randomly distributed within a crystal comparable to glassy carbon, which is depicted in Figures 3C and S5. In support of the XRD pattern and TEM imaging, Raman spectra of the carbon film featured a narrow width of the D and G bands (Figure S7). After annealing of the electroplated films, the bandwidth decreased and I_D/I_G was improved. This can be attributed to the reduction of the heteroatom species and the alignment of the graphene sheets.

The high degree of graphitization of the laser carbon film is further confirmed by XPS, where the C 1s spectral line is centered at 284.6 eV, corresponding to an sp²-hybridized state. The tail in the C 1s peak indicates the presence of multiple functional groups with nitrogen and oxygen heteroatoms. In pairs, the deconvoluted N 1s peak is attributed to multiple configurations. Within the graphene sheets, the nitrogen heteroatoms are incorporated as N–Q and at the edge of the graphene sheet, the heteroatoms form pyridinic and pyrrolic rings and an additional N–Q species corresponds to amides. The amide species propagate after LIC by bond dissociation along the C–N bond, due to the lowest enthalpy in the chain at this site, which transforms and leaves some residue in the form of amide. The processing in an ambient environment (air) influences oxidation of dissociated species and promotes the incorporation of nitrogen into the graphene lattice. The oxygen conjugated species were observed in the O 1s line, which was present in multiple conformations with some preference at the aromatic carboxyl site in the graphene sheet. Other species were reduced but not observed in an equivalent rate to each other. This suggests an initial scission along the C–N bonds, with further bonding into a hexagonal graphene-like lattice. Across all samples, the carbon base from the C 1s line does not significantly change, indicating relatively close contribution to each functional group.

A laser carbon film serves as a preferable seed layer for the electroplating of copper. The presence of heteroatoms in the graphene sheets bolsters charge accumulation at their sites, in the case of both oxygen and nitrogen conjugated species. The site provides a higher negative potential on the Allen electronegativity scale compared to carbon, thus inducing preferred oxidation of Cu ions during electroplating and passivating the heteroatom charge. This was confirmed by the reduction of heteroatom species observed in XPS C 1s peaks in Figure 4D and by the shift of the O 1s spectrum toward CuO and Cu₂O species reducing contribution from a hydroxyl functional group (Figure 4E). As was observed from Cu 2p in Figure 4F, the surface featured not only a metallic Cu but also additional Cu₂O, CuO, and Cu(OH)₂ species, due to the immediate oxidation of the film in air, and some contamination from the CuSO₄ electrolyte in a minor concentration, fit with a standard spectrum for each species.

Annealing of the copper/carbon composite film showed persistent low diffusion of the carbon into the copper, due to its low solubility, as is indicated in the specific phase diagrams. The annealing of thin copper films at 1000 °C promoted melting and led to aggregation as droplets (microspheroids) on laser carbon, induced by heterogeneous melting with reduced Gibbs free energy for the thin films. The surface energies of copper and carbon are 1.79 and 0.26 J cm⁻² respectively, which also support the aggregation of the molten copper into microspheroids by a reduction of the contact area, due to the significantly higher surface energy, causing nonwetting of the copper on the laser carbon. The surface roughness further promoted the contact area reduction by copper.

The reduced surface coverage due to the molten copper further yielded a reduced signal intensity in XRD and XPS. Figure 4G indicates improvement in the areal contribution and narrowing the width of the C≡C–C–C/ C–H peak, caused by redistribution and cleavage of heteroatoms in the lattice. This was confirmed by the observed O 1s peaks, where the carboxyl groups are present mainly in their aliphatic form (Figure 4H).

The introduction of air during cooling at 450 °C induced a copper nanowire growth, which was promoted by a vapor–solid growth mechanism. With an excess of oxygen, a Cu₂O phase formed, fostering hillock aggregation due to relief of compressive stresses. At temperatures above 400 °C, the Cu₂O phase is not thermodynamically stable. Therefore, with further oxygen supply, the nanowires of CuO started growing, with the hillocks as the nucleation sites. At the nucleation site and in the growing nanowire, the crystal structure attained a stacked layer matching unit cell size. At the nucleation site and in the growing nanowire, the crystal structure attained a stacked layer matching unit cell size. This mechanism of nanowire growth was further confirmed with TEM (Figure 3E–G). Within the top 5 nm of the urchin shell, crystallites with layered or random orientation were found, characteristic for a monoclinic CuO lattice or an intermediate amorphous phase. As is seen in Figures 3G and S6, above 5 nm within the shell, the structure featured an initial cubic lattice specific to CuO with a matching unit cell size. At the nucleation site and in the grown nanowire, the crystal structure attained a stacked layer structure, and further along the length of the crystal featured the moiré fringes due to an overlapping of the different crystal orientations. The core nanowire crystallite was circularly grown nanowire, the crystal structure attained a stacked layer matching unit cell size.
The detailed XPS spectra in Figure 4K,L, and summary Table S5, confirmed the oxidation of the film with a shift to the dominant CuO and Cu(OH)$_2$ functional groups, with a minor contribution from Cu$_2$O, due to the short duration of the annealing step. The contribution from chemical species could be tailored with temperature and the duration of annealing.15,54

Both types of composite films were further utilized for enzyme-free glucose detection. Unlike Cu-MS, CuO retained its chemical composition after a number of cycles, which was observed by the redox reactions during CV (Figure S9). After the injection of glucose, the oxidation current significantly increased, as is seen in Figure 5D. Such an increase occurred in micro- and nanostructured copper, which provided sufficient surface area for the oxidation of the metal via the formation of gluconolactone and, furthermore, gluconic acid in the presence of the analyte in the alkaline electrolyte80 (see the equations in Figure S10A).

Despite the presumed higher surface-to-volume ratio of the CuO-U compared to Cu-MS, both materials attained insignificant differences in the amperometric response (Figure 6A,B). The equivalent responses of both electrodes were supported by the dispersion of Cu nanoparticles in addition to the microspheres on laser carbon film after annealing, which provide an improvement of the total sensing area. The presence of transition-metal nanoparticles induces the glucose-sensing ability of the electrode due to an increase in the sensing surface area.10,17 However, both films exhibited saturation in the amperometric signal for a glucose concentration over ~10 mM. As we hypothesized, this might have occurred due to constraining of the metal oxidation and a fast saturation at the active sites with hydroxyl species. On the other hand, the Cu-MS electrode featured a difference in the oxidation kinetics for the microspheres and the nanoparticles and in line with previously reported values for the metal nanoparticles.18,19 Nevertheless, the CuO-U electrode exhibited the highest sensitivity due to the superior electron transfer for the material at the selected overpotential,11 supported by a nanostructured morphology, as is observed from CV (Figure 5D). Both electrodes tested here exhibited similar performance in terms of linear range, sensitivity, but suffered from a higher LOD of glucose, compared to carbon-based glucose sensors reported by other authors, as shown in Table 2. Limited and uneven distribution of the nanostructured active sites responsible for the glucose oxidation results in some deterioration of the total acquired signal, compared to films of purely native transitional metal and its oxide.26,27,28,37

However, simplicity of preparation, utilization of low-cost materials, and preparation technologies provide great opportunity in the large-scale manufacturing of this type of sensors for adequate glucose detection levels, which requires distinguishing analyte levels from 100 to 10 mM according to reports of the Diabetes Association.94

Glucose detection in blood, sweat, and urine requires high selectivity in sensing, due to the presence of other endogenous compounds such as urea, uric acid, acetic acid, and other sugars (sucrose, fructose), which constrain the functionality of the enzyme-free sensors.13 As is shown in Figure 6C, the fabricated electrodes showed no response to other analytes. However, Cu-MS exhibited response to sucrose. The response attenuated to the baseline level after ~1000 s and further did not show any significant contribution after multiple sucrose additions. However, sucrose is not present in the human body in the free form, and after digestion, converts to fructose and glucose, thus not directly contributing to the measurement of glucose levels. The activity to the sucrose occurs due to the oxidation of the micro- and nanostructured copper.80 The sucrose was hydrolyzed to fructose and glucose in the presence of an alkaline electrolyte, further promoting the generation of gluconic acid through a similar reaction chain, and is depicted in Figure S10B.13,80 At the selected potential, only Cu-MS indicated an increase in the amperometric signal, whereas CuO-U was active only to glucose, and the reduction of the signal was observed via CV in Figure S9B.

The strong adhesion of microparticles to laser carbon substrate and the minor chemical variation of the amperometric response after multiple repetitions are depicted in Figure 6D. The detachment of the nanoparticles slightly hinders further amperometric response. The nanoparticles show faster oxidation to the highest copper valence state, which is more brittle than an initial chemical modification.25 Notably, oxidation occurs at a significantly faster rate for Cu-MS. The revealing of the rough CuO-U surface with reduced nanowire presence and underlying microparticles is still sufficient to provide a sufficient oxidation rate for amperometric signal within relatively close limits and show an ~5% signal deterioration after each cycle. Despite EDX showing a significant change in the Cu/O ratio, we speculate that the surface remains predominantly in the CuO phase due to the depth penetration of the electron beam within microns, a contribution from the core Cu$_2$O. During the amperometric test, only the soluble hydroxyl species were adsorbed on CuO-U; thus, the electrode could be reused after their removal by rinsing with DI water.11

A new generation of wearable smart devices (watches, bands, rings) has secured its position in the noninvasive, permanent monitoring of the heartbeat and pressure with reliable output, comparable to professional medicinal devices. Currently, the industry is expanding to cover direct real-time health risk monitoring of sweat, saliva, and blood.95 Inclusion of such sensors in the final device relies on their performance durability, selectivity of the detection, mechanical flexibility, and ease of manufacturing. However, the sensing area is currently constrained due to design requirements and the number of sensors is limited to the output voltage of the device for functional use throughout the day.55,96 New sensors can be included within movable parts, which require flexibility and bendability and are complemented with operation at a low power input (~100 mW).96 Thus, the selection of the material and engineering of the sensor must meet multiple parameters for successful commercial utilization. The glucose sensor reported here meets several of these requirements, such as selectivity of detection, operation during bending, and simplicity of the fabrication based on rapid manufacturing technologies. Further development in the engineering and packaging can provide improvement in the output currents by implementing conductive glue between the sensor substrate and the reading electrode, which can be fabricated by printing directly on the bendable films. Additionally, the complete sensor requires the realization of the detection electrode together with the counter electrode and the reference electrode in one system for real-time analyte monitoring, which can be easily implemented with printing and LIC technology. The sensor based on CuO-U shows good repeatability after washing in DI water, and due to selectivity only to glucose, it is feasible to rinse with the running water.
5. CONCLUSIONS

We have demonstrated the fabrication of Cu-MS- and CuO-U-based enzyme-free glucose sensors on a freestanding laser carbon film. They were fabricated on the initial substrate by the laser-induced carbonization from a Nomex sheet using optimized laser parameters, followed by the electroplating of copper for 10 min. The electroplated copper was further annealed at 1000 °C in a nitrogen atmosphere to obtain Cu-MS, which is caused by melting and wrapping of the initial copper sheets. The formation of Cu-MS depended on the annealing temperature and duration of electroplating. Exposure to air at 450 °C during the cooling process resulted in the formation of urchin-like CuO seeded with nanowires, featuring a diameter starting from 15 nm and a length up to 5 μm. The performance of the composite electrodes toward the detection of glucose was investigated by cyclic voltammetry and real-time amperometric tests. Both electrode types exhibited amperometric activity in the presence of glucose with a superior sensitivity compared to other carbon-based glucose sensor materials. The sensitivities of the Cu-MS and CuO-U electrodes were 0.25 and 0.32 mA cm⁻² mM⁻¹, respectively, within a linear range between 1 μM and 3 mM. The limit of detection was estimated at 1.75 and 7.56 μM for the Cu-MS and CuO-U electrodes, respectively, which are one of the lowest among carbon-based glucose sensors. Furthermore, the fabricated electrodes exhibited excellent selectivity toward glucose. However, the Cu-MS electrode also exhibited some sensing activity in the presence of sucrose. The composite electrodes were also flexible and retained their sensing capabilities after multiple bending repetitions. The saturation sensitivity of the electrodes was determined at 10 mM, which is at the risk level for healthy humans from urine, sweat, and blood. These fabricated substrates are suitable for personalized, low-cost, point-of-care glucose monitoring and for the determination of sugar concentration in the food industry.

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acsanm.1c03149.

Cu-MS and CuO-U evolution with variation in the duration of initial electroplating and annealing temperature, including evaluation of their sizes; assessment of the most suitable materials from sets prepared under different conditions; detailed study with TEM from selected CuO-U and its substrate and additional characterization with Raman spectroscopy; XPS survey spectra of selected specimens and their compositional assessment; deconvolution from detailed XPS spectra; full cyclic voltammogram of the studied amperometric glucose biosensors in the presence of an electrolyte with added glucose or sucrose; and chemical formulas for the involved reactions (PDF)

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Notes

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