

Chemical imaging of buried interfaces in organic-inorganic devices using FIB-TOF-SIMS

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Abstract

Organic-inorganic hybrid materials enable the design and fabrication of new materials with enhanced properties. The interface between the organic and inorganic materials is often critical to the device's performance and therefore chemical characterization is of significant interest. Since the interfaces are often buried, milling by focused ion beams (FIB) to expose the interface is becoming increasingly popular. Chemical imaging can subsequently be obtained using secondary ion mass spectrometry. However, the FIB milling process damages the organic material. In this study, we make an organic-inorganic test structure to develop a detailed understanding of the processes involved in FIB milling and SIMS imaging. We provide an analysis methodology that involves a "clean-up" process using sputtering with an Argon gas cluster ion source to remove the FIB induced damage. The methodology is evaluated for an additive manufactured encapsulated strain sensor containing silver tracks embedded in a polymeric material. We show a polymer-silver interface with a resolution of 440 nm and that the polymer contains a low level of silver particulates.

Keywords: *FIB, ToF-SIMS, Argon cluster, milling, polymer, hybrid interfaces, additive manufacturing*

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Introduction

Organic/inorganic interfaces play an important role in many innovative technologies, including polymer electrolyte membrane fuel cells, dye-sensitized solar cells, and next-generation nanoelectronic devices which incorporate organic components alongside conventional conductors and semiconductors.^{1,2,3,4,5} Another important example is the enabling technology of additive manufacturing (AM) which is at heart an interfacial problem where multiple layers of material are consecutively deposited to build up a physical 3D object from digital data. This interface challenge with AM will be further extended with emergence of the next-generation AM that seeks to combine both structural and functional materials together in a single entity. For future 3D-printed electronics, this will necessitate the co-deposition of both dielectric (organic) materials alongside conductive inorganics that will create complex organic-inorganic interfaces as both inter and intra layers.⁶

The design, fabrication and optimisation of such devices demand methods for characterizing the organic-inorganic interfaces. Since these interfaces are usually buried, techniques are required for these inter- and intra-layers that can access depths of tens of microns into these complex structures. The integration of a focussed ion beam (FIB) to mill away material and expose these buried interfaces for high-resolution scanning electron microscopy (SEM) has revolutionised the measurement capability.⁷ Tomographic images can be created through sequential milling and imaging cycles with a spatial resolution approaching 1 nm in the plane of the SEM image and a slice thickness down to 3 nm.^{7,8} Analytical capabilities are possible when combined with energy dispersive x-ray spectroscopy (EDX) for chemical imaging of elements and back scatter electron diffraction for crystal microstructure. However, the capability is limited for light elements such as lithium used in battery technologies, hybrid organic-inorganic materials such as perovskites and for organic materials.

Secondary ion mass spectrometry (SIMS) uses a focused ion beam to probe a surface causing atoms and molecules to be emitted in a process known as sputtering. A small fraction of this sputtered material is charged and is subsequently analysed in a mass spectrometer. Traditionally, this has been a time-of-flight (ToF) analyser but, more recently, a high-resolution Orbitrap spectrometer has been used.⁹ The same approach, used successfully in dual beam FIB/SEM, may be used with SIMS. The use of FIB milling to access buried interfaces overcomes well-known limitations of traditional sputter depth profiling that result from material dependent sputtering yields and the presence of voids. This was demonstrated in early work by Satoh et al.¹⁰ and Crow et al.¹¹ In both studies, a Ga⁺ FIB was used to mill and map elemental distributions in integrated circuits. A few years later, using a serial milling approach, Tomiyasu et al.¹² and Dunn & Hull¹³ showed the potential for FIB-SIMS tomography based on 3D reconstruction from sequences of SIMS images. FIB-SIMS tomography developed concurrently with, if not slightly ahead of, FIB/SEM but the use of the methodology has been very limited in

1 comparison. This, in part, could be due to the difficulty in extracting secondary ions for mass analysis
2 from the surfaces cut at an angle to the average surface. However, in recent years ToF-SIMS instruments
3 have been equipped with FIB to access buried interfaces, and ToF analysers have been integrated in
4 SEM instruments for mass analysis.¹⁴ The use of FIB in state-of-the-art ToF-SIMS instrumentation has
5 proven powerful e.g., for multi-element tomography of solid oxide fuel cells^{15,16,17} and the analysis of
6 battery materials where ToF-SIMS imaging provided evidence of the dissolution of manganese from a
7 $\text{LiNi}_{0.5}\text{Co}_{0.2}\text{Mn}_{0.3}\text{O}_2$ electrode.¹⁸ These results are excellent but chemical degradation caused by the FIB
8 and possible beam heating effects in materials could limit the usefulness of FIBs for preparation of
9 inorganic-organic hybrid materials for SIMS imaging.¹⁹ The many existing protocols for FIB
10 preparation and tomography of organic samples are useful but, in general, do not address chemical
11 degradation as they have been developed for electron microscopy rather than molecular analysis using
12 a surface sensitive technique.

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14 The chemical degradation causes the FIB-SIMS analysis of organic-inorganic hybrid materials to be
15 challenging. Methods are needed to minimize or remove the FIB damage for successful FIB-SIMS
16 analysis. Recent work by Iida et al. shows how organic material that has been damaged by a FIB may
17 be removed using Ar^+ gas cluster ion beam (GCIB) sputtering.²⁰ Here, we present a study of FIB-SIMS
18 using three different FIB sources Ga^+ , Bi^+ and Bi_3^+ . A simple inorganic-organic test device has been
19 developed consisting of a regular pattern of polymer, either polystyrene (PS) or poly(methyl
20 methacrylate) (PMMA), and silicon oxide. This is used to study and optimize the FIB-SIMS parameters
21 and the FIB damage cleanup procedure. The effectiveness of the method is demonstrated in the study
22 of the organic-metal interface of a prototype encapsulated strain sensor made by 3D inkjet printing and
23 a Cu nanoparticulate deposited on poly(ethylene terephthalate) (PET) via low power IR irradiation
24 procedure.

25 **Experimental Methods**

26 **Test device:** Organic-inorganic test devices were made using a Long-Life™ MCP-10 microchannel
27 plate (MCP) (Photonis, US) that are commonly used in ToF detection systems. The MCP (Figure SI.1)
28 consists of a regular array of hexagonal close packed open tubes with 10 μm hole diameter and a centre-
29 to-centre separation of 12 μm . These holes are tilted at 12° to the surface normal. The MCP is made
30 from glass tubes sintered together with a rubidium-containing Nichrome coating for secondary electron
31 emission. The holes are subsequently filled with either PS or PMMA using a 4-step procedure described
32 by Steinhart et al.²¹

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34 The MCP was degreased in a bath of 20% nitric acid (Sigma-Aldrich, UK) for more than 24 hours and
35 subsequently rinsed with purified water and acetone (Sigma-Aldrich, UK). The dried MCP was placed
36 in a glass petri dish and the surface of the MCP was covered with powdered polystyrene (molecular

1 weight 2430, Sigma Aldrich, UK) or PMMA (molecular weight 340000, Sigma Aldrich, UK). A metal
2 weight of 500 g was placed on the polystyrene. Keeping this arrangement, the polymer was annealed to
3 200 °C for 5 h in a vacuum allowing the polymer to melt and fill the channels in the MCP (see the
4 scheme process in Figure SI.1)

5
6 **3D inkjet printed organic-inorganic sample:** A prototype encapsulated strain sensor containing silver
7 tracks embedded in a polymeric material was manufactured by 3D inkjet printing of two different inks.
8 The organic ink is made in-house and based on tri (propylene glycol) diacrylate (TPGDA) and the silver
9 nanoparticle ink was purchased from Advanced Nano Products (SilverJet DGP-40LT-15C). Both
10 polymer and silver layers were printed and cured/sintered contemporaneously by a LED-based UV
11 source connected to the print-heads of a PixDro LP50 printer. More details of the ink formulations and
12 printing process are described elsewhere.²² The analysed volume of the sample contains, from bottom
13 to top, 15 layers of polymer, 10 layers of silver and 5 layers of polymer. Based on previous studies,
14 each polymer layer is approximately 10 µm thick and each silver layer is between 300 nm and 700 nm
15 thick.

16 **Low temperature sintering of Cu nanoparticulate:** The Cu on PET sample was produced using a novel
17 printing technique that utilises a low-power, focused IR laser to selectively irradiate metallic
18 nanoparticulate loaded slurries. Using the widely known phenomena of low-temperature sintering of
19 nanoparticulates, the laser irradiation causes the nanoparticulates – Cu in this case – to consolidate
20 at a lower temperature than would be expected for the bulk materials. Though this approach shows
21 promise as 3D printing process, this particular sample demonstrates the potential for use within
22 flexible 2D printed electronics, hence the use of a flexible insulating substrate (PET). Using such a
23 method shows the potential of achieving high resolution conductive tracks in an affordable and
24 industrially acceptable material. It is therefore important to understand the inorganic/organic
25 interface to assess aspects such as adhesion and conductivity of the tracks.

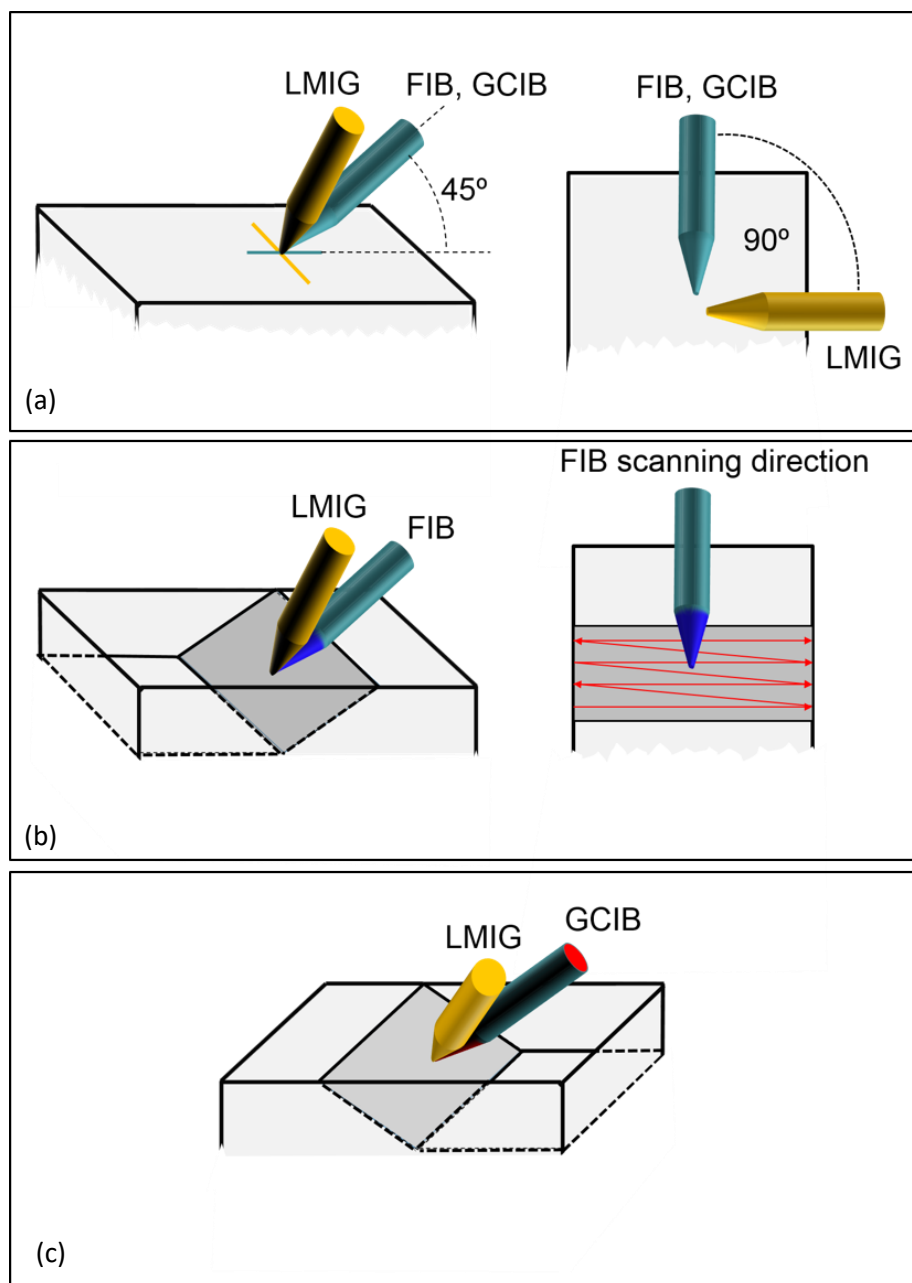
26
27
28 **FIB-ToF-SIMS:** In this study we use a TOF-SIMS 5 (ION-TOF GmbH, Germany). The ToF analyser
29 is normal to the sample surface and it is equipped with a dual source ion column mounted at 45° to the
30 sample surface. The dual source ion column provides a 20 keV Ar⁺ GCIB or a 30 keV Ga⁺ FIB ion
31 beam aligned in the same direction. A liquid metal ion gun (LMIG) provides a Bi_n⁺ analytical beam.
32 This is also at 45° to the sample surface but is located at an azimuthal angle of 90° from the Ar⁺ GCIB
33 and Ga⁺ column. Low energy electrons are used for charge neutralization and are provided by a flood
34 gun delivering a current of 5 µA with an energy of 20 eV mounted at an angle of 57° to the sample
35 normal. The secondary ion and electron imaging of the cross section are made using a mass-separated

1 30 keV Bi_3^{2+} beam in the high lateral resolution, “Fast Imaging”, mode operating with the analyser in
2 delayed extraction mode. Using this condition, a spatial resolution of approximately of 200 nm is
3 achieved.^{23,24} As already shown by Mihara et al.²⁵ this modality allows the minimization of topographic
4 effects to improve the detection of secondary ions from the crater wall.

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6 **Cross-sectioning and clean-up:** To prepare a cross section for analysis, a Ga^+ FIB is used with an energy
7 of 30 keV and a current of 20 nA to mill out a crater that is 120 μm in the x -direction and 80 μm in the
8 y -direction as shown in Figure 1 (see Table 1 for the beam parameters). The milling is monitored using
9 the Bi_3^{2+} beam with a field of view of 200 μm by 200 μm containing 512×512 pixels. This configuration
10 is shown in Figure 1a with the Ga^+ FIB beam entering from the top of each image. The crater wall
11 prepared for SIMS imaging is the top wall, closest to the Ga^+ source, at an angle of approximately 45°
12 to the MCP surface normal. The beam follows a sawtooth raster pattern with the scan direction being
13 from left to right in the images, starting from the bottom and moving to the top in each scan as showed
14 in Figure 1b.

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2 *Figure 1. Schematic representation of the FIB-ToF-SIMS geometry. (a) FIB, LMIG, GCIB and sample orientations. (b)*
 3 *Sample milling orientation and FIB scan direction. (c) Cleaning process using the GCIB with a 180° rotation of the*
 4 *sample.*

5 The sample is rotated 180° (see Figure 1c) for removal of the damaged layer on the FIB section using
 6 the Ar⁺ GCIB. This clean-up procedure leads to the recovery of characteristic organic secondary ion
 7 signals from the milled wall (at approximately 45°, as we shall see later). The Ar⁺ GCIB was an Ar₂₅₀₀⁺
 8 beam at 10 keV (4 eV/atom) with a beam current of 6 nA and was rastered over a 500 μm × 500 μm
 9 area. Secondary ion images are recorded during the Ar⁺ GCIB clean-up to follow the organic signal
 10 recovery using the Bi₃²⁺ beam with a current of 0.02 pA.

11 A further study to compare the effect of a bismuth source in the milling process is made by using the
 12 3D OrbiSIMS (based on the TOF-SIMS 5 platform from ION-TOF GmbH, Germany) equipped with

1 a 30 keV bismuth LMIG with a Wien filter so that mass selected direct current beams of Bi⁺ or Bi₃⁺ can
 2 be used.

Focused Ion Beam Parameters						
Type	Energy	Crater size	Current	Pixels in crater	Milling passes	Dwell time
Ga ⁺	30 keV	120 μm × 80 μm	20 nA	206 × 137	1, 2, 3	25 ms/pixel

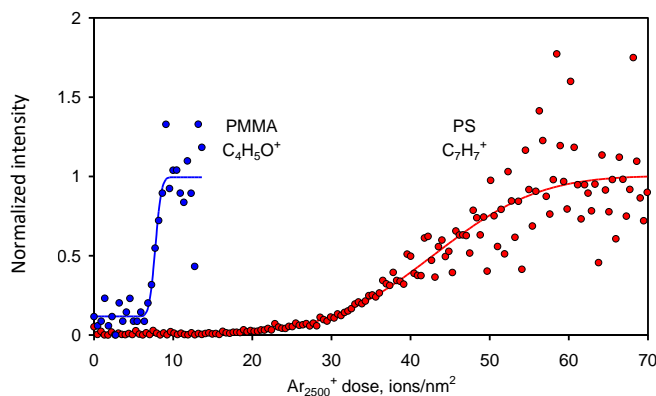
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 4 *Table 1. FIB parameters used for the milling process.*

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 6 To optimize the lateral resolution of the secondary ion image, after the cleaning procedure, the sample
 7 is further rotated 90°. In that orientation the Bi₃²⁺ beam is normal to the FIB section surface to give an
 8 unhindered view of the cut surface.

9 Results and discussion

10 The recovery of characteristic polymer fragment ion signals from the 45° milled wall of both PS and
 11 PMMA filled MCPs is shown as a function of the Ar⁺ GCIB sputter dose in Figure 2. The characteristic
 12 ion signal is seen to rise after a certain dose similar to that previously reported for ion and electron beam
 13 damage.^{26,27,28,29}

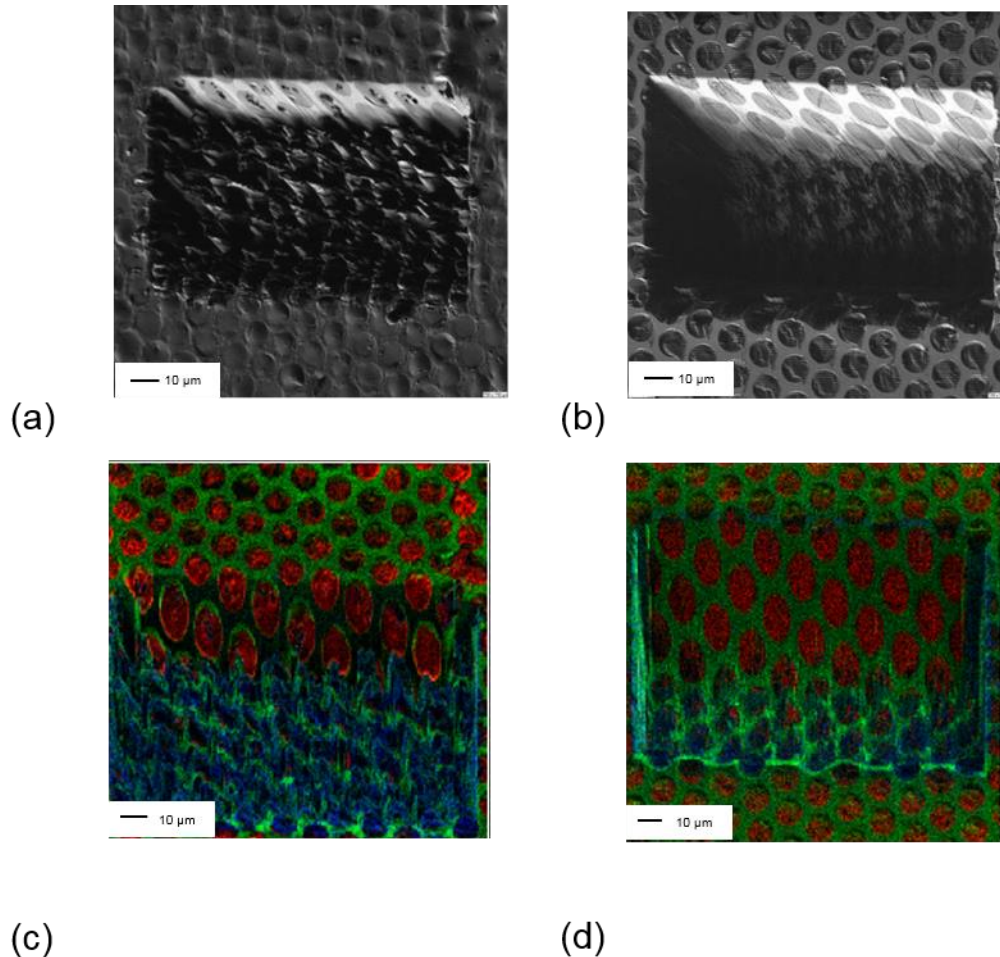
14
 15 After FIB milling, a high Ga⁺ signal is detected on the sloping surface and the organic signals are very
 16 weak, presumably, due to polymer crosslinking and carbonisation. The damage provided by a FIB dose
 17 of 4.08×10^4 ions/nm² was removed by approximately 10 ions/nm² of 10 keV Ar₂₅₀₀⁺ for PMMA and
 18 60 ions/nm² for PS as shown in Figure 2. The faster recovery of the signal from PMMA is expected as
 19 this polymer is not of the type that cross-links when exposed to ionizing radiation.³⁰



20 *Figure 2. Recovery of the intensity of characteristic fragment ions from PS (C₇H₇⁺, red) and PMMA (C₄H₅O⁺, blue)*
 21 *during Ar⁺ GCIB clean-up.*

1

2 In Figure 3a and 3b we observe, respectively, a SE image of the crater obtained providing a total dose
3 of 4.09×10^4 ions/nm² for the MCP filled with PMMA and PS. Below, in Figure 3c and 3d we show
4 the secondary ion images after the clean-up process for both filled specimens.



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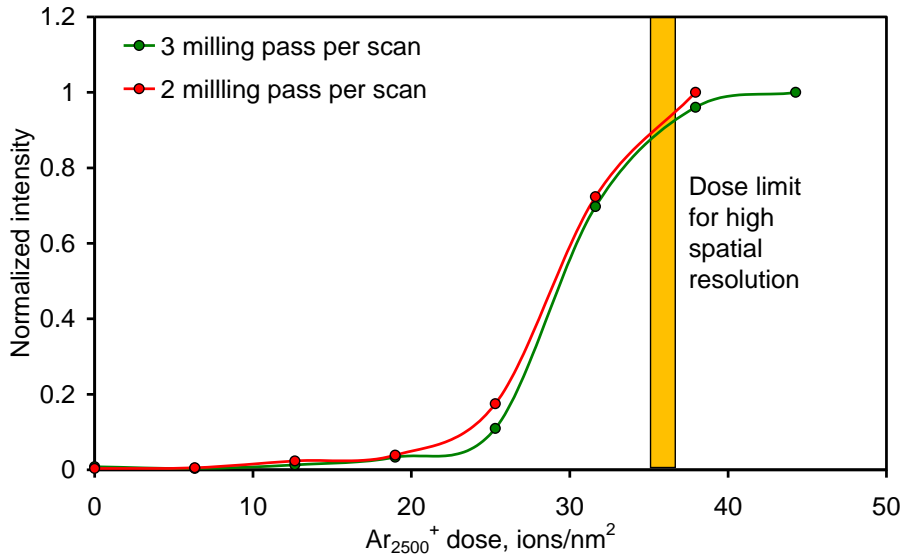
Figure 3. Secondary Electron (SE) in (a) and (b) and Secondary Ion (SI) in (c) and (d) images using the Bi₃⁺⁺ ion beam of a crater milled into MCP test devices filled with either PS or PMMA. (a) SE image of MPC filled with PS and (b) similarly for PMMA. (c) SI overlay images of Rb⁺ (green), Ga⁺ (blue) and the characteristic PS fragment C₇H₇⁺ (red) and (d) similarly for PMMA characteristic fragment C₄H₅O⁺. Note that the imaging beam is at 45° to the MCP overall surface but the imaging system arranges the scan to a smaller deflection in its tilt direction to form a true square on horizontal surfaces. This causes the left hand vertical crater wall in (a) and (b) to appear to be sloping. Its opposite wall becomes hidden. Images (c) and (d) do not show this as the sample is rotated 90° for analysis and the images are rotated back that 90° for consistent presentation with (a) and (b).

6

7 FIB milling cutting procedure with Ga⁺ using 1, 2 and 3 milling scans

8 Part of this work consisted of the observation of how the number of milling scans affect the milled depth
9 and the required cleaning. By using 1, 2 and 3 milling scans, we linearly increase the Ga⁺ beam dose.
10 As a first step, we observe by optical microscopy how the milled depth develops in the case of the
11 empty MCP matrix and in the case of the MCP matrix filled with PS (Figure SI.2). This increase of
12 depth with dose appears to be near-linear and is very similar in the polymer-filled MCPs and in the

1 empty MCP since the sputtering of the polymer occurs at a higher rate than the sputtering of the
2 inorganic MCP.³¹



3

4 *Figure 4. C₇H₇⁺ fragment signal recovery as a function of Ar⁺ GCIB dose for 2 milling pass per scan and 3 milling pass*
5 *per scan FIB craters in the PS filled MCP. The yellow band shows the dose limit for retaining the highest spatial resolution.*

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7 Figure 4 shows the clean-up at the crater wall (by selecting a region of interest centralized in the 45°
8 wall) formed by using 2 and 3 scans for the FIB milling. These craters are obtained by using the Ga⁺
9 doses of 40900 ions/nm² and 61300 ions/nm². In cleaning the MCP, Figure 4 shows that a dose of 38
10 ions/nm² largely cleans the surface. In this geometry, the Ar⁺ GCIB is normal to the surface. It is shown
11 elsewhere³² that the sputtering yield of pure PS at 0° incidence, using 10 keV Ar₂₅₀₀⁺, is 21 nm³.
12 However, in sputtering the Ga implanted PS, the yield is reduced to ~0.6 nm³, a reduction of ~50 times
13 so that the 38 ions/nm² is what is required to remove the remaining 23 nm of damaged and implanted
14 PS.³² If we were to overclean by a further 38 ions/nm², we should remove around a further 1 μm of pure
15 PS which would lead to blurring at the edges of included phases by significantly more than this figure
16 since the yield rises strongly at edges. Even if one limited the overclean dose to just 10%, the spatial
17 resolution would begin to be compromised. Different polymers are affected to different extents so that
18 the correct dose for both cleaning and retaining optimum spatial resolution will depend on the organic
19 material analysed. It is, therefore, recommended to measure the signals for the organic materials
20 anticipated in the sample and to stop the sputtering for imaging the sample at, or soon after, the signal
21 has reached 90-95% of its final value, as shown in Figure 4.

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23 Theory

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25 In the sputter shaping of materials, the angle dependence of the sputtering yield is critical. Wehner³³
26 first showed the importance of this effect in the surface topography after sputtering. In 2012, Seah³⁴

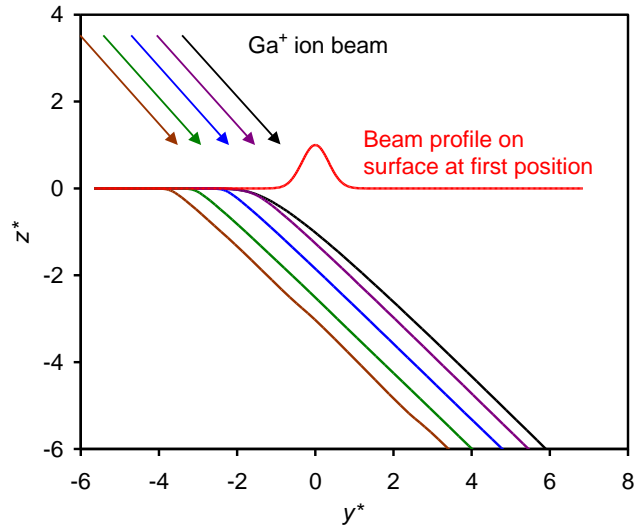
1 showed that good computations of surface form could be obtained by using a combination of Sigmund's
2 sputtering theory³⁵ and the evaluations of relevant parameters by Yamamura et al.³⁶ Further evaluations
3 were made by Seah³⁴ leading to a set of 9 equations to describe the sputtering yield and its dependence
4 on the incidence angle, θ .

5
6 In the present work, we sputter using 30 keV Ga⁺, 30 keV Bi⁺ or Bi₃⁺, or 10 keV Ar₂₀₀₀⁺. In the first
7 case, we sputter unfilled MCPs. Their geometry gives a solid fraction of 0.37 for the SiO₂ of the MCP.
8 The angle dependence of the sputtering yield for SiO₂ using 30 keV Ga⁺ primary ions using Seah's
9 equations³¹ shows that the maximum yield occurs at $\theta_{\max} = 80^\circ$ (see Figure SI.3). Similar calculations
10 for Ga⁺ and Bi⁺ and for energies in the range 10 keV to 50 keV give θ_{\max} as 77° to 81° with slightly
11 higher values for the lower mass primary ion and at the higher energies.

12
13 At 45° incidence to the MCP surface, the sputtering yield is 6.74 atoms/ion. In our 3 craters, we have
14 used doses of 20400, 40900 and 61300 ions/nm² so we would expect depths of 4.8; 9.7 and 14.6 μm
15 allowing for the 37% solid fraction. Actually, the depths measured by confocal microscopy are 13.5,
16 25.5 and 31 μm , some 2.4 times higher. The reason for this is that whilst the average MCP surface is at
17 45° to the beam, the local surface being sputtered is not at 45° to the beam. The sputtering is conducted
18 as a raster scan where the beam has a profile, approximately Gaussian with a full width at half maximum
19 of 0.8 μm on the sample surface and the pixel-to-pixel interval is 0.156 μm . Thus, the first application
20 of the beam at 45° to the MCP surface digs a small hole but at the next pixel, there is an increased slope
21 on the side being sputtered so the yield rises and a slightly deeper hole is made and so on until the angle
22 of the FIB-milled slope at the side of the hole is approximately θ_{\max} . It may locally increase beyond this
23 value as a result of changes to other pixels. At 70° to the FIB-milled surface, the predicted vertical
24 depths are 12.5, 25.1 and 37.7 μm , not very different from those observed using the confocal
25 microscopy. The average angle of the beam to the FIB-milled surface may be less than θ_{\max} since the
26 sputtering takes some time to equilibrate and also the fringes of the beam are sputtering surfaces that
27 are at lower angles, or it may be higher as discussed above. Thus, the measurements of the depth milled
28 are broadly consistent with the model proposed. In Figure SI.4, we show AFM measurements on an
29 unfilled MCP confirming that the FIB-milled slope angle starts at around 30° and rises to 40° as the
30 depth from the MCP surface increases.

31
32 Note that here we have assumed an average solid fraction of 37% which would be valid if the holes
33 were random. However, we can see in that the alignment of the holes means that, for the beam mid-way
34 between the holes, the solid fraction is 100% whereas when aligned at the centres of the holes it is only
35 17%. This causes the unevenness observed of both the crater floor and the important back wall.

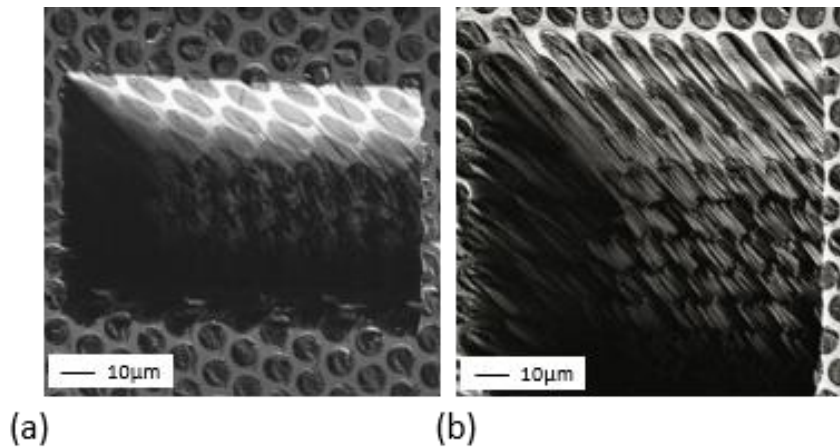
1 Calculations of the sputter profile using 30 keV Ga^+ on glass, at the final edge, are shown in Figure 5.
2 These calculations are made with the beam orientated along the z -direction with columns of material
3 along that direction and at the end, for presentation purposes the whole image is rotated 45°
4 anticlockwise. The starred co-ordinate directions are in this rotated plane.



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25 *Figure 5. Calculations of the developing profile for 30 keV Ga^+ sputtering of glass at 45° incidence. The sputtering starts*
26 *at $y^* = 0$ and proceeds with many pulses at that position before moving 156 nm to the left and repeating the process many*
27 *times. The beam profile is taken to be a Gaussian as shown but the precise shape is not important.*

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29
30 **Could FIB milling with Bi^+ and Bi_3^+ be a useful alternative to Ga^+ for milling organic-inorganic**
31 **hybrid materials?**

32
33 Gallium is the most widely used source for FIB milling. Our 3D OrbiSIMS instrument also has a 30
34 keV bismuth liquid metal ion source equipped with a Wien filter so that mass selected direct current
35 beams of Bi^+ or Bi_3^+ can be used, as mentioned in the experimental section. This provides an interesting
36 possibility since cluster beams are known to create less damage in organic materials and therefore may
37 be more suitable for inorganic-organic materials. Bi^+ and Ga^+ have much the same effect on glass in
38 terms of the theory given above. To study the efficiency of these sources, we use the PS-filled test
39 device. Figure 6 shows secondary electron images of craters with milling using Ga^+ (ToF-SIMS V) and
40 Bi_3^+ (OrbiSIMS).



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2
3 *Figure 6. Emitted electron image using Bi_3^{++} ions in the SIMS instrument of milled craters. (a) milled with Ga^+ and*
4 *milled with Bi_3^+ . The sloping face is at the top. The left wall looks sloping but is vertical and this arises since the imaging*
5 *beam comes from the right and the image surface geometry is “corrected” by expanding the x-direction by a factor of $2^{0.5}$.*
6

7 The results for Bi^+ (not shown) are very similar to, but poorer than, those for Ga^+ . The results for Bi_3^+
8 are very much poorer and this arises partly due to the very different angular dependence of the sputtering
9 of cluster ions compared with monatomic ions^{37,38}. The value of θ_{max} for Bi_3^+ is much lower than for
10 Ga^+ as the ions do not penetrate so far or so linearly. This leads to a slope closer to 30° with the ions no
11 longer at the very grazing incidence into the polymer in the MCP holes shown in Figure SI.5. This
12 allows a greater physical roughness to develop there and may explain the poor results for Bi_3^+ .

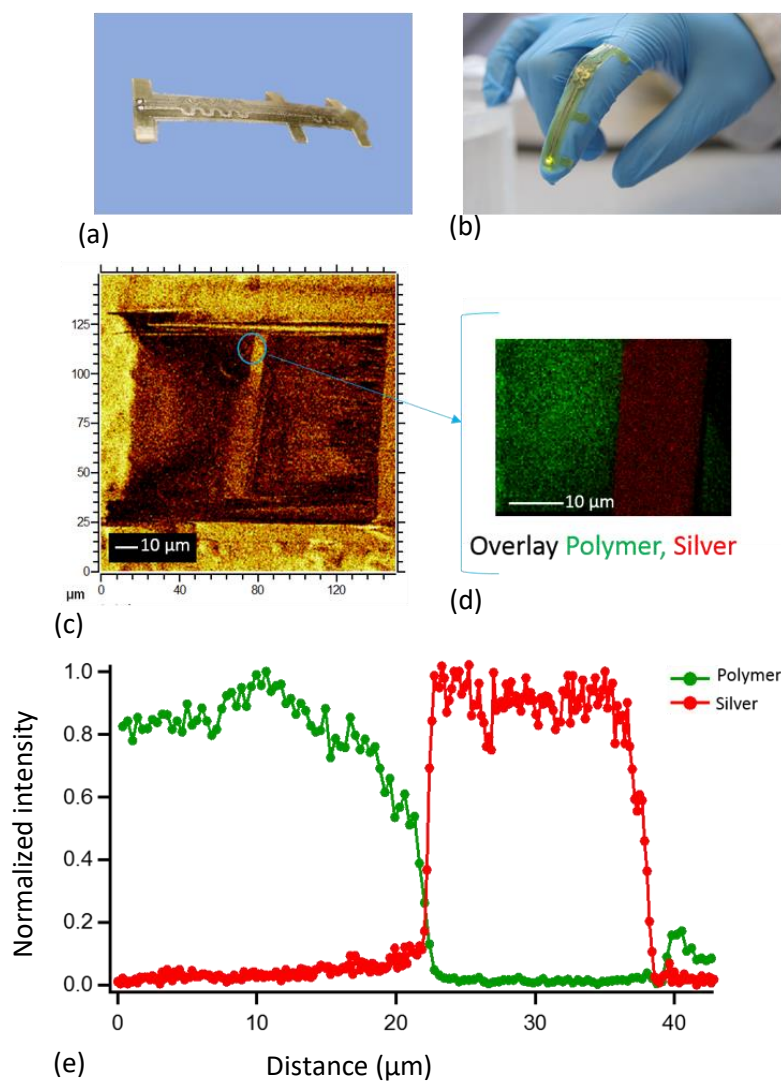
13 Observation of the columnar structure in each MCP hole using Bi_3^+ in Figure 6b would indicate that
14 when using Bi_3^+ sputtering ions, some 2 to 3 μm of organic material may need to be removed to clean
15 up the surface. If this were pure polystyrene, this would need the high dose of 70 ions/ nm^2 for 10 keV
16 Ar_{2500}^+ , but if the removal of damaged polymer is one to two orders of magnitude slower^{27,29, 32}, this
17 figure is raised by that factor and would be impossible to remove in a practical time. Clearly, the spatial
18 resolution is very poor and the cleaning of the FIB section cut using Bi_3^+ may degrade that resolution
19 further. It is therefore clear that the FIB-sectioning should be made with Ga^+ .

20
21 **Application of FIB-TOF-SIMS to a 3D inkjet printed material**

22 We applied FIB-TOF-SIMS to image the buried organic-inorganic interface of the 3D printed samples
23 described in the experimental section. The two samples has been milled from the upper polymer part
24 following the methodology described earlier.

25 Figure 7 (a, b) shows the strain sensor and the total secondary ion images of the FIB crater after GCIB
26 cleaning with a dose of 18.7 ions/ nm^2 (Fig. 7c). Figure 7d shows a 45 μm x 45 μm map with an overlay
27 of the signals for polymer ions and silver. After removal of the implanted gallium, the interface between
28 the two materials is clearly distinguishable. To improve the signal to noise, an integrated intensity
29 profile is created by summing the horizontal linescans between the regions indicated in Fig. 7d. Owing

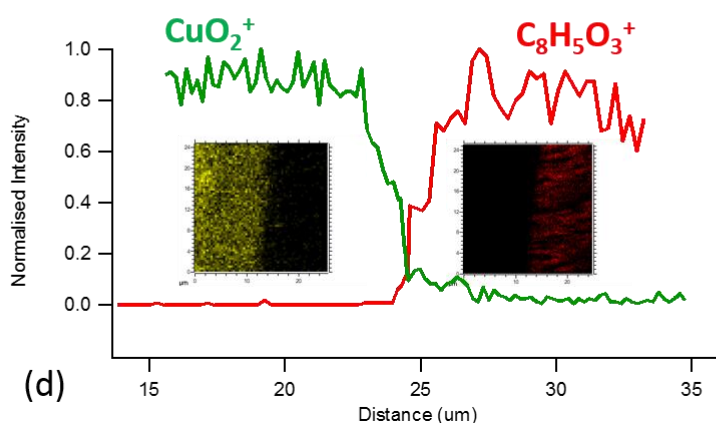
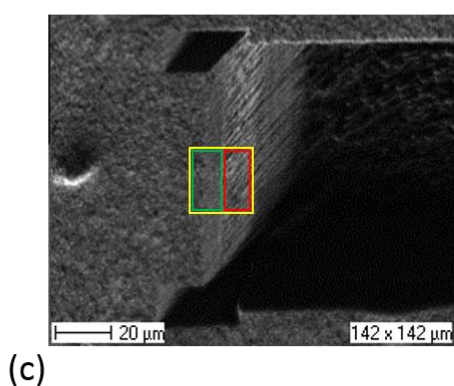
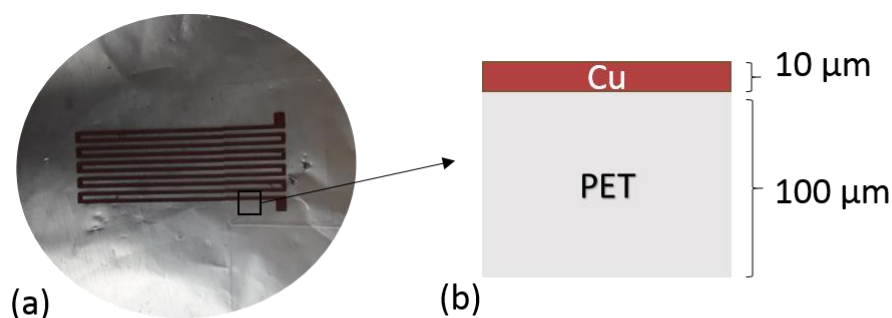
1 to a tilt in the sample, a small horizontal offset is applied to each linescan to align them. Using the 16-
 2 84 % definition of profile resolution, the silver signal intensity across the interface (Fig. 7e) gives an
 3 organic-inorganic interface width as 440 nm. Above the interface, we see a small quantity of Ag. This
 4 could be interpreted as atomic diffusion whilst the overlayers are deposited or it could be from very
 5 small particles of Ag. Analysis of the SIMS spectra shows significant populations of Ag_3^+ and Ag_5^+
 6 within the polymer, consistent with very small particulates, whilst none of the Ag adducts that would
 7 be expected for atomic or organically bonded Ag were observable. The populations are slightly smaller
 8 than for bulk Ag.



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11 **Figure 7. FIB-TOF-SIMS analysis of an additive manufactured encapsulated strain sensor. (a) Optical image of the sensor**
 12 **and (b) the sensor integrated with a disposable glove. (c) Total SI image of milled and cleaned crater. (d) High-resolution**
 13 **SI image of the interface with characteristic polymer fragment m/z 51 (green) and silver (the sum of Ag_3 and Ag_5) (red).**
 14 **Ag_1 is not used owing to mass interferences). (e) Normalized intensity profile of summed lines centred at the interfaces in**
 15 **(d).**

1 Fig. 8 (a, b) shows the Cu nanoparticulate deposited on PET substrate and the schematic of the layer
 2 thickness. The sample has been milled with a Ga dose of 40900 ions/nm² and the characteristic PET
 3 fragment C₈H₅O₃⁺ signal has been recovered with a dose of 12 ions/nm². In this specific case the
 4 interface has been determined in two steps: once the crater wall has been selected (see Fig. 8 c), the
 5 copper signal coming from CuO₂⁺ ions has been recorded. Subsequently the Ar cluster cleaning, the
 6 PET fragment signal has been recorded from the same area selected in Fig. 8 (c) and the line profile of
 7 125 summed lines centered at the interface is shown in Fig. 8 (d). This approach permitted us to define
 8 a well interface also in the case of friable material.



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10 *Figure 8. FIB-TOF-SIMS analysis of copper nanoparticulate layer on PET via low power IR irradiation. (a) Image of the*
 11 *device and (b) schematic of the layer depth.. (c) SE image of milled crater. Normalized intensity profile of summed lines*
 12 *centred at the interfaces in (d with incorporate the SI image from the characteristic signal of the Cu nanoparticulate in*
 13 *green and the PET fragment in red.*

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Conclusions

FIB-TOF-SIMS is a useful method for chemical imaging of buried interfaces. However, the FIB milling damages organic surfaces, which could limit the application for important organic-inorganic systems. We show that an Ar⁺ GCIB clean-up cycle can remove this damage and provide essential information to do this effectively. We have developed a test device based on a microchannel plate consisting of a regular honeycomb array of tubes of 10 μm diameter in glass. The tubes are filled with either PMMA or PS. Using this device, we determine that 10 keV Ar₂₅₀₀⁺ doses of 10 ions/nm² and 60 ions/nm² are required to recover the organic signal for PMMA and PS, respectively.

We postulate that since Bi₃⁺ has a higher sputtering yield than Ga⁺ for organics, that the effects of damage on organics may be less. We use the test-device to evaluate this for FIB milling with 30 keV Ga⁺, Bi⁺ and Bi₃⁺. We find, the reverse is in fact the case and explain this in terms of a theory based on the angular dependent sputtering yield. In this theory, the angle of incidence evolves from the initial 45° geometrical angle with the fresh surface to stabilize at near θ_{\max} as the milling proceeds. Since θ_{\max} differs substantially between atomic (approximately grazing) and cluster projectiles (approximately 45°) then the ability to successfully mill through materials with vastly different sputtering rates is affected. Therefore, gallium is recommended for milling organic-inorganic hybrid materials. We propose that the polymer filled MCP test device could have utility for FIB-TOF-SIMS interlaboratory comparisons to evaluate instrumentation and protocols and to improve reproducibility.

We apply the method to study the buried interface between a track of sintered silver nanoparticles and a UV-cured polymer in a 3D inkjet printed prototype encapsulated strain sensor. The two materials show a resolution at the interface of 440 nm. Above the interface is a very low level of Ag particulates.

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Supporting Information. This shows the method of preparation of the polymer-filled MCP, details of the MCP craters as well as the AFM study to check the precise orientation of the sputtered plane of the sample, optical microscope images on empty and filled MCP, angular dependence on sputtering yield

Bibliography

- (1) Urban, J. J. One Model to Rule Them All. *Nat. Mater.* **2017**, *16* (2), 157–159.
- (2) Maurin, G.; Serre, C.; Cooper, A.; Férey, G. The New Age of MOFs and of Their Porous-Related Solids. *Chem. Soc. Rev.* **2017**, *46* (11), 3104–3107.
- (3) Breitwieser, M.; Klingele, M.; Vierrath, S.; Zengerle, R.; Thiele, S. Tailoring the Membrane-Electrode Interface in PEM Fuel Cells: A Review and Perspective on Novel Engineering Approaches. *Adv. Energy Mater.* **2018**, *8* (4), 1701257.
- (4) Freitag, M.; Teuscher, J.; Saygili, Y.; Zhang, X.; Giordano, F.; Liska, P.; Hua, J.; Zakeeruddin, S. M.; Moser, J. E.; Grätzel, M.; et al. Dye-Sensitized Solar Cells for Efficient Power Generation under Ambient Lighting. *Nat. Photonics* **2017**, *11* (6), 372–378.
- (5) Ali, Z.; Shakir, I.; Kang, D. J. Highly Efficient Photoelectrochemical Response by Sea-Urchin Shaped ZnO/TiO₂ Nano/Micro Hybrid Heterostructures Co-Sensitized with CdS/CdSe. *J. Mater. Chem. A* **2014**, *2* (18), 6474–6479.
- (6) EPSRC. EPSRC Programme Grant: Enabling Next Generation Additive Manufacturing <http://gow.epsrc.ac.uk/NGBOViewGrant.aspx?GrantRef=EP/P031684/1> (accessed Aug 13, 2018).
- (7) Giannuzzi, L. A.; Stevie, F. A. *Introduction to Focused Ion Beams : Instrumentation, Theory, Techniques and Practice*; Springer, 2010.
- (8) Narayan, K.; Danielson, C. M.; Lagarec, K.; Lowekamp, B. C.; Coffman, P.; Laquerre, A.; Phaneuf, M. W.; Hope, T. J.; Subramaniam, S. Multi-Resolution Correlative Focused Ion Beam Scanning Electron Microscopy: Applications to Cell Biology. *J. Struct. Biol.* **2014**, *185* (3), 278–284.
- (9) Passarelli, M. K.; Pirkl, A.; Moellers, R.; Grinfeld, D.; Kollmer, F.; Havelund, R.; Newman, C. F.; Marshall, P. S.; Arlinghaus, H.; Alexander, M. R.; et al. The 3D OrbiSIMS—label-Free Metabolic Imaging with Subcellular Lateral Resolution and High Mass-Resolving Power. *Nat. Methods* **2017**, *14* (12), 1175–1183.
- (10) Satoh, H. Three-Dimensional Analysis of a Microstructure by Submicron Secondary Ion Mass Spectrometry. *J. Vac. Sci. Technol. B Microelectron. Nanom. Struct.* **1991**, *9* (5), 2638.
- (11) Crow, G. A.; Christman, L.; Utlaut, M. A Focused Ion Beam Secondary Ion Mass Spectroscopy System. *J. Vac. Sci. Technol. B Microelectron. Nanom. Struct.* **1995**, *13* (6), 2607.
- (12) Tomiyasu, B.; Fukuju, I.; Komatsubara, H.; Owari, M.; Nihei, Y. High Spatial Resolution 3D Analysis of Materials Using Gallium Focused Ion Beam Secondary Ion Mass Spectrometry (FIB SIMS). *Nucl. Instruments Methods Phys. Res. Sect. B Beam Interact. with Mater. Atoms* **1998**, *136–138*, 1028–1033.
- (13) Dunn, D. N.; Hull, R. Reconstruction of Three-Dimensional Chemistry and Geometry Using

- 1 Focused Ion Beam Microscopy. *Appl. Phys. Lett.* **1999**, 75 (21), 3414.
- 2 (14) Whitby, J. A.; Östlund, F.; Horvath, P.; Gabureac, M.; Riesterer, J. L.; Utke, I.; Hohl, M.;
3 Sedláček, L.; Jiruše, J.; Friedli, V.; et al. High Spatial Resolution Time-of-Flight Secondary Ion
4 Mass Spectrometry for the Masses: A Novel Orthogonal ToF FIB-SIMS Instrument with *In Situ*
5 AFM. *Adv. Mater. Sci. Eng.* **2012**, 2012, 1–13.
- 6 (15) Hammond, J. S.; Fisher, G. L.; Bryan, S. R.; Kanarbik, R.; Möller, P. FIB-TOF Tomography of
7 Solid Oxide Fuel Cells. *Microsc. Microanal.* **2013**, 19 (2), 672–673.
- 8 (16) Kanarbik, R.; Moller, P.; Kivi, I.; Lust, E. Application of FIB-TOF-SIMS and FIB-SEM-EDX
9 Methods for the Analysis of Element Mobility in Solid Oxide Fuel Cells. *ECS Trans.* **2013**, 57
10 (1), 581–587.
- 11 (17) PRIEBE, A.; GORET, G.; BLEUET, P.; AUDOIT, G.; LAURENCIN, J.; BARNES, J.-P. 3D Correlative
12 Morphological and Elemental Characterization of Materials at the Deep Submicrometre Scale.
13 *J. Microsc.* **2016**, 264 (2), 247–251.
- 14 (18) Börner, M.; Horsthemke, F.; Kollmer, F.; Haseloff, S.; Friesen, A.; Niehoff, P.; Nowak, S.;
15 Winter, M.; Schappacher, F. M. *Degradation Effects on the Surface of Commercial*
16 *LiNi0.5Co0.2Mn0.3O2 Electrodes*; 2016; Vol. 335.
- 17 (19) Seah, M. P. Cluster Ion Sputtering: Molecular Ion Yield Relationships for Different Cluster
18 Primary Ions in Static SIMS of Organic Materials. *Surf. Interface Anal.* **2007**, 39 (11), 890–897.
- 19 (20) Iida, S.; Carr, D. M.; Fisher, G. L.; Miyayama, T. Accurate and Reproducible In-Depth
20 Observation of Organic–inorganic Hybrid Materials Using FIB-TOF-SIMS. *J. Vac. Sci. Technol. B,*
21 *Nanotechnol. Microelectron. Mater. Process. Meas. Phenom.* **2018**, 36 (3), 03F107.
- 22 (21) Steinhart, M.; Wendorff, J. H.; Greiner, A.; Wehrspohn, R. B.; Nielsch, K.; Schilling, J.; Choi, J.;
23 Gösele, U. Polymer Nanotubes by Wetting of Ordered Porous Templates. *Science (80-)*.
24 **2002**, 296 (5575), 1997.
- 25 (22) Saleh, E.; Zhang, F.; He, Y.; Vaithilingam, J.; Fernandez, J. L.; Wildman, R.; Ashcroft, I.; Hague,
26 R.; Dickens, P.; Tuck, C. 3D Inkjet Printing of Electronics Using UV Conversion. *Adv. Mater.*
27 *Technol.* **2017**, 2 (10), 1700134.
- 28 (23) Vanbellinghen, Q. P.; Elie, N.; Eller, M. J.; Della-Negra, S.; Touboul, D.; Brunelle, A. Time-of-
29 Flight Secondary Ion Mass Spectrometry Imaging of Biological Samples with Delayed
30 Extraction for High Mass and High Spatial Resolutions. *Rapid Commun. Mass Spectrom.* **2015**,
31 29 (13), 1187–1195.
- 32 (24) Claus, T. K.; Richter, B.; Hahn, V.; Welle, A.; Kayser, S.; Wegener, M.; Bastmeyer, M.; Delaittre,
33 G.; Barner-Kowollik, C. Simultaneous Dual Encoding of Three-Dimensional Structures by Light-
34 Induced Modular Ligation. *Angew. Chemie Int. Ed.* **2016**, 55 (11), 3817–3822.
- 35 (25) Mihara, I.; Havelund, R.; Gilmore, I. S. Embedding-Free Method for Preparation of Cross-
36 Sections of Organic Materials for Micro Chemical Analysis Using Gas Cluster Ion Beam
37 Sputtering. *Anal. Chem.* **2017**, 89 (9), 4781–4785.
- 38 (26) Miyayama, T.; Sanada, N.; Bryan, S. R.; Hammond, J. S.; Suzuki, M. Removal of Ar⁺ Beam-
39 Induced Damaged Layers from Polyimide Surfaces with Argon Gas Cluster Ion Beams. *Surf.*
40 *Interface Anal.* **2010**, 42 (9), 1453–1457.
- 41 (27) Yamamoto, Y.; Ichiki, K.; Seki, T.; Aoki, T.; Matsuo, J. Ion-Induced Damage Evaluation with Ar
42 Cluster Ion Beams. *Surf. Interface Anal.* **2013**, 45 (1), 167–170.
- 43 (28) Havelund, R.; Seah, M. P.; Shard, A. G.; Gilmore, I. S. Electron Flood Gun Damage Effects in 3D

- 1 Secondary Ion Mass Spectrometry Imaging of Organics. *J. Am. Soc. Mass Spectrom.* **2014**, 25
2 (9), 1565–1571.
- 3 (29) Kawashima, T.; Morita, H.; Fukumoto, N.; Kurosawa, T.; Aoyagi, S. Examination of Ion Beam
4 Induced Damage on Polymer Surface Using Ar Clusters. *Surf. Interface Anal.* **2016**, 48 (11),
5 1175–1180.
- 6 (30) Moore, J. A.; Choi, J. O. Degradation of Poly(Methyl Methacrylate); 1991; pp 156–192.
- 7 (31) Seah, M. P. Universal Equation for Argon Gas Cluster Sputtering Yields. *J. Phys. Chem. C* **2013**,
8 117 (24), 12622–12632.
- 9 (32) Tiddia, M.; Seah, M. P.; Shard, A. G.; Mula, G.; Havelund, R.; Gilmore, I. S. Argon Cluster
10 Cleaning of Ga + FIB-Milled Sections of Organic and Hybrid Materials. *Surf. Interface Anal.*
11 **2018**, DOI: 10.1002/sia.6522.
- 12 (33) Wehner, G.; Gottfried. Influence of the Angle of Incidence on Sputtering Yields. *J. Appl. Phys.*
13 **1959**, 30 (11), 1762–1765.
- 14 (34) Seah, M. P. Topography Effects and Monatomic Ion Sputtering of Undulating Surfaces,
15 Particles and Large Nanoparticles: Sputtering Yields, Effective Sputter Rates and Topography
16 Evolution. *Surf. Interface Anal.* **2012**, 44 (2), 208–218.
- 17 (35) Sigmund, P. Theory of Sputtering. I. Sputtering Yield of Amorphous and Polycrystalline
18 Targets. *Phys. Rev.* **1969**, 184 (2), 383–416.
- 19 (36) Yamamura, Y.; Itikawa, Y.; Itoh, N. *Angular Dependence of Sputtering Yields of Monoatomic*
20 *Solids*; 1983.
- 21 (37) Seah, M. P.; Spencer, S. J.; Shard, A. G. Depth Resolution, Angle Dependence, and the
22 Sputtering Yield of Irganox 1010 by Coronene Primary Ions. *J. Phys. Chem. B* **2013**, 117 (39),
23 11885–11892.
- 24 (38) Seah, M. P.; Spencer, S. J.; Shard, A. G. Angle Dependence of Argon Gas Cluster Sputtering
25 Yields for Organic Materials. *J. Phys. Chem. B* **2015**, 119 (7), 3297–3303.

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