Glassy carbon manufacture using rapid photonic curing

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ABSTRACT

Photonic curing was explored as a rapid method for producing glassy carbon coatings, reducing processing time from \(
\sim \) 20 h for conventional thermal processing down to \(
\sim \) 1 min. A resole-type thermoset polymer resin coated on steel foil was used as a precursor, placed in a nitrogen purged container and exposed to high energy light (\(\sim \) 27 J/cm\(^2\) per pulse for up to 20 pulses). Comparison samples were produced at 800 °C using a conventional nitrogen purged thermal route. For both photonic and conventionally produced coatings, Raman spectroscopy and primary peak XPS data showed \(sp^2\) bonded carbon, indicative of bulk glassy carbon. This transformation evolved with increasing number of pulses, and therefore amount of energy transferred to the coating. The produced coatings were resilient, highly smooth, with no evidence of surface defects. XPS analysis indicated greater \(sp^3\) content at the immediate surface (5–10 nm) for photonic cured carbon compared with thermally cured carbon, likely due to the local environment (temperature, atmosphere) around the surface during conversion. The ability to rapidly manufacture glassy carbon coatings provides new opportunities to expand the window of applications of glassy carbons in coatings towards large-scale high volume applications.

Introduction

Glassy carbon (GC) also known as vitreous carbon, is a non-graphitizing carbon with characteristics intermediate between those of graphite and ceramics\textsuperscript{[1, 2]}. Non-graphitizing carbons are carbons that consist of graphitic nano-crystallites that are cross-linked between crystals in an isotropic fashion\textsuperscript{[1]}. This structure offers a host of important material properties including substantial hardness, high

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temperature resistance, impermeability to gases, low electrical resistance, chemical resistance and biocompatibility [1–8]. These properties make it a useful material for electrochemical, corrosion and medical applications among others. GC is commonly used as a reference electrode in electrochemistry [4, 9–14], as corrosion protection for current collectors [6, 15–17] and as a surgical implant material in both humans and animals [7, 8, 18–20].

Glassy carbon is typically produced through the carbonization of a resole-type thermoset resin precursor [21, 22] either as a bulk material or as a film [6, 16, 20, 23–26]. Common precursor resins are phenol formaldehyde and furfuryl alcohol [27]. After coating and prior to heat treatment the precursor is cured by temperature or catalyst [20, 28]. Carbonizing the precursor is achieved by heat treatment in an inert atmosphere with a minimum temperature of 600 °C although most commercially produced glassy carbon materials are carbonized between 1000 and 2000 °C [23, 25, 26]. Limitations of this production method are the time and temperature required to carbonize the precursor. Polymer carbonization is a process that generally requires several hours of high temperature thermal treatment, typically in an inert atmosphere, with additional time for ramping the furnace temperature up and down. For a high quality conversion from resin to carbon low ramp rates are used, typically 1–5 °C/min, this causes the typical conversion time to lie in the order of several hours [24, 29]. The requirement of high temperatures and long carbonization times are bottlenecks in the usability of GC in all its applications. In addition to this, if the material were to be required as a coating then the substrate itself would need to be subjected to the same thermal treatment as the glassy carbon.

Photonic curing is a recently developed alternative technology to the conventional convection methods which currently sees most use in the drying and sintering of printed or coated thin films [30–35]. Photonic curing uses xenon lamps to rapidly pulse high-intensity light, transferring a high energy density to the surface of an object [30]. Photonic curing minimizes the thermal treatment times while allowing conventional oven-comparable temperatures to be reached at the surface of the exposed object [31]. The reduction in time is achieved because there are no requirements for ramping, holding or cooling times as the energy delivery using light is almost instantaneous. An example from Cronon et al. [36] where photonic curing is used to sinter aqueous silver inks shows a reduction in curing time from 90 min to 9 ms. Photonic curing can also be integrated with roll-to-roll processing [32] and the curing speed allows for a decrease in physical space required in automated, in-line manufacturing as it would replace an otherwise large heat or radiation curing oven [33]. The localised and transient nature of photonic curing also enables the potential use of low-cost, flexible, temperature sensitive components/carrier substrates to be used for thin and thick carbon films that ordinarily could not withstand conventional thermal treatment [33]. Photonic curing pulse parameters, such as lamp power, exposure time and frequency determine the overall power density transferred onto the film [30]. Current applications of photonic curing include the manufacture of flexible printed electronics (such as RFID and photovoltaics) [31, 32], where the manufacturing process requires the thermal treatment or chemical conversion/sintering of a film on a temperature sensitive substrate. Photonic curing has been demonstrated to convert a CuO ink into a Cu thick-film rapidly on a cheap PET substrate, a process which otherwise requires long thermal treatment times in an inert atmosphere [34]. In this case, no inert atmosphere is required as the energy delivery is faster than rate of copper oxidation. Another application is sintering silver-based inks to improve the conductivity [31] or to improve the electronic properties of ZnO nano-wires [35]. This method is most suited to surface treatment and is therefore applied to coatings on a carrier material rather than bulk materials.

In this paper, photonic curing is explored as a faster means of converting films of phenol formaldehyde resin into glassy carbon. This paper proposes a novel manufacturing process for glassy carbon using a rapid photonic curing method, which has the ability to cure the coating without adversely affecting the temperature of the substrate while decreasing the conversion time from hours to seconds. The converted resin is then compared against a thermally derived glassy carbon. According to the most recently accepted models [3] the structure of GC consists of predominantly $sp^2$ bonded carbon domains that are disordered graphitic sheets, giving it a comparable structure to fullerenes. The randomly orientated graphitic sheets give GC its isotropic nature (3). Raman spectroscopy and X-ray photoelectron spectroscopy are therefore used to elucidate the
structure to focus on bulk (microns) and surface (nanometres), respectively, given their different penetration depths [37, 38], while morphology is compared using a scanning electron microscope and white light interferometry.

**Methodology**

**Materials**

The selected precursor for glassy carbon is phenol formaldehyde resin, a resole-type thermoset polymer resin [Curaphen 40-852 B60, Bitrez Limited (UK)]. The selected substrate for the precursor deposition is 316 stainless steel foil of 25 micron thickness [Advent Research Materials (UK)].

**Conversion to carbon methods**

The resin was coated onto the steel foil substrate using bar coating (PrintCoat Instruments K Control Coater). Close-wound bars sizes “0” and “2” (0.05 and 0.15 mm wire diameters, respectively) were used yielding a theoretical wet coating thicknesses of 4 and 12 microns, respectively, at a speed of 1.1 m/min. Two different coating thicknesses were used as a result of preliminary tests which revealed the structural integrity limitations of the phenol formaldehyde coatings during the heat treatment stages. The thicker coating (12 micron wet) is used in the conventional thermal conversion because that method experienced delamination with thinner coatings, while thin coatings (4 micron wet) are used for the photonic curing method which experiences delamination for thicker coatings. These limitations are specific to the materials and methodology used in this paper. Ambient temperature and humidity during coating were 22 °C and 33%, respectively. To permanently crosslink the thermoset resin the manufacturer’s suggested conditions were used, a temperature cure of 220 °C for 30 min in a convection oven (Votsch VTL 60/90). Carbonization of the cured phenolic resin was then achieved using two different methods: a conventional thermal curing method and the photonic curing method.

The conventional method of producing glassy carbon is heat treatment of the cross-linked phenol formaldehyde precursor in an inert atmosphere. For this work, the coated phenol formaldehyde resin samples were exposed to a set point temperature of 800 °C which was reached at a ramp rate of 2 °C per minute in a tube furnace (Carbolite GERO GHA 12/450) with a holding time of 2 h. An inert atmosphere in the furnace was achieved with a constant 1 L/minute flow of nitrogen gas. This tube furnace required temperatures of < 50 °C in order to be opened, hence resulting in long average cooling times of ~ 12 h. These parameters follow conventional processes as established in the literature [23–26, 29, 39]. A maximum carbonization temperature of 800 °C was selected to prevent excessive softening of the steel carrier substrate.

For the photonic curing method, the coated phenol formaldehyde resin samples were heat treated using a photonic curing system (Novacentrix PulseForge 1200, Austin TX). The selected pulse parameters for carbonization were developed via preliminary testing and were a lamp voltage of 450 V and a pulse duration of 2000 μs giving an energy output of ~ 27 J/cm² per pulse. Multiple sequential pulses were performed, up to a maximum of 20 pulses (5, 10, 15, 20 pulses), at a frequency of 0.3 Hz. The sample resided in a dedicated sample holder with a transparent glass window as seen in Fig. 1. The sample holder was continually vented with nitrogen gas to achieve an inert atmosphere and remove the evolving off-gases during treatment. The steps required for both treatments are demonstrated graphically in Fig. 2. The total heat treatment cycle of the conventional method takes > 20 h, whereas the photonic curing method requires up to 64 s (depending on the number of pulses).

**Analysis of carbonized samples**

To characterize the glassy carbon material, Raman spectroscopy, X-ray photoelectron spectroscopy (XPS) and scanning electron microscopy techniques were used. Raman spectra of the carbon coatings were compared using a Renishaw inVia Raman system (Renishaw plc., Wotton-Under-Edge, UK) in backscattering configuration with a 50× objective lens (NA: 0.50, spot size ≈ 1 μm) The spectrometer used a laser of 532 nm excitation wavelength with a maximum power output of 32 mW. The spectrum acquisition parameters were set to 5% laser power (= 1.6 mW), 10 s exposure time and a grating of 1800 lines/mm. Raman spectroscopy of this particular wavelength penetrates approximately 0.7–1.0
microns in carbon materials which aligns with the typical coating thicknesses used in this study, and therefore is representative of the bulk morphology of the coating [37, 38].

X-Ray photoelectron spectra (XPS) were obtained using a Kratos Axis Supra (Kratos Analytical, Manchester, UK) using a monochromated Al Kα source (225 W power). All spectra were recorded without using a charge neutraliser. Survey scans were recorded at a pass energy of 160 eV and the high resolution data at 40 eV with energy steps of 0.1 eV. The emission current was set at 15 mA. Peaks were identified and quantified with Shirley backgrounds and GL(50) line shapes, with carbon D parameter analysis performed as outlined by Kaciulis [40] using CasaXPS Version 2.3.22PR1.0. The photon penetration depth of XPS lies in the range of 5–10 nm [37, 38], so that XPS is primarily a surface characterisation method rather than being representative of the bulk.

Scanning electron microscopy [JEOL 7800F with Oxford Instruments Aztec EDS system XMaxN detector field emission gun scanning electron microscope (FEGSEM)] was used to determine the surface topography of the carbon coatings. White light interferometry (NT2000, Veeco Instruments, Inc., Plainview, NY, USA) was used to measure a full three-dimensional surface profile over the coated surfaces in order to evaluate surface roughness. Twenty times magnification was used, giving a measurement area of 232 μm by 310 μm (at a resolution of 640 × 480 pixels with sampling at 485 nm intervals).

**Results and discussion**

**Coating topography**

The photonic and thermal cured carbon coatings were both well adhered to the stainless steel substrate under manual handling. Both methods yielded uniform, smooth and durable carbon coatings. The coatings did not readily delaminate when exposed to bending of the coated foil, see Fig. 3.

SEM images of the photonic cured carbon and thermal cured carbon samples are presented in Fig. 4. To provide a contrast for SEM imaging, between the

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**Figure 1** Converted carbon coating (~ 3 × 3 cm surface area) placed in the photonic curing sample holder attached to the nitrogen gas supply.

**Figure 2** Breakdown of the time requirements of the iterative steps to complete both heat treatment methods.

**Glass-like Carbon Conversion Methods: Time Scale Comparison**

- **Conventional Method**
  - Increasing furnace temperature to desired set point: 6.5 hours
  - Furnace set point holding time: 2 hours
  - Furnace cooling time before samples can be extracted: >12 hours
  - Finished Carbon Coating

- **Photonic Curing**
  - Exposure to 1-20 pulses of 27 Jcm⁻² light: Time ranging between 20ms and 64 seconds dependent on the amount of pulses.
  - Finished Carbon Coating
coating and the underlying steel substrate, part of the coating was removed. The images show the photonic cured samples after different exposure times (10, 15 and 20 pulses) as well as the thermal cured material at a magnification of 30000 times. Regardless of the preparation method, the carbon coatings presented a highly smooth surface and were consistent without evidence of any defects such as pinholes, delamination or scratches. Thermal and photonic derived surfaces presented similar visually glossy appearances, showing no discernible difference between the two carbonisation methods.

White light interferometry measurements show a full three-dimensional surface profile over the coated surfaces. A total of 5 measurements are taken for each sample with average surface roughness measurements (\(S_a\)) calculated over the measured surface, the errors are calculated using standard deviation. These average surface roughness measurements are compared in Fig. 5. The untreated phenol formaldehyde resin precursor had an average surface roughness, \(S_a\), of 470 nm. Upon treatment, the roughness of the surface was reduced, and the \(S_a\) values of both the photonic cured carbon coatings and the thermal cured carbon coating were similar and in the range of \(\sim 110–140\) nm. This data demonstrate that both thermal and photonic carbonization methods resulted in similar carbon surface topographies.

**Raman spectroscopy of treated materials**

Raman spectroscopy of carbons revolves around 4 major peaks; \(D\), \(D'\), \(G\) and \(T\) band peaks (at around \(1360, 1620, 1590\) and \(1060\) cm\(^{-1}\), respectively). From the locations, shapes and relative intensities of these 4 peaks the carbon–carbon bond structure can be deduced. The two main \(sp^2\) bonded carbon peaks are the \(D\) band (carbon ring breathing mode) and the \(G\) band (\(sp^2\) bonded carbon stretching mode) at \(\sim 1360\) cm\(^{-1}\) and \(\sim 1590\) cm\(^{-1}\), respectively. The \(T\) peak (\(\sim 1060\) cm\(^{-1}\)) is an indicator for \(sp^3\) bonded carbon in the sample. The \(D'\) peak is an indicator of the grain size of the carbon material, located at \(1620\) cm\(^{-1}\) [41–44]. This \(D'\) peak becomes more prominent as the grain size increases, which in the case of glassy carbons is associated with an increase in carbonization temperature. From the amorphization trajectory in the study of disordered carbons by Ferrari et al. [45], the \(G\) band location and the relative intensity ratio between the \(D\) and \(G\) band (referred to as \(I(D/G))\) can be used to estimate the relative \(sp^2\) and \(sp^3\) content of the carbon samples.

The Raman spectra of photonic cured (5, 10, 15 and 20 pulses) and thermal treated coatings are compared in Fig. 6. Mean \(I(D/G)\) ratios are indicated together with standard deviation based on three repeat measurements at different locations on each sample. The Raman spectra of the polymer precursor without thermal treatment were not Raman active. The Raman spectrum of the conventionally thermal treated coatings showed the typical glassy carbon shape that would be produced at the low end of the typical heat treatment temperature (HTT) range. The \(D\) peak was less intense than the \(G\) peak, whereas its full-width half maximum (FWHM) was approximately 3 times that of the \(G\) peak. The secondary peaks in the
**Figure 4** Scanning electron microscopy images of photonic cured carbon coatings at different amounts of pulses and thermal cured carbon with exposed stainless steel substrate at 30000 magnification. A 10 pulses, B 15 pulses, C 20 pulses, D thermal cured carbon.

| Sample Type                     | Sa Value   |
|---------------------------------|------------|
| Untreated Resin                 | 470 ± 23 nm|
| Photonic Cured (5 Pulses)       | 116 ± 11 nm|
| Photonic Cured (10 Pulses)      | 122 ± 19 nm|
| Photonic Cured (15 Pulses)      | 122 ± 10 nm|
| Photonic Cured (20 Pulses)      | 138 ± 5 nm |
| Thermal Cured                   | 137 ± 13 nm|

**Figure 5** White light interferometry images of phenol formaldehyde resin, photonic cured carbon samples and thermal cured glassy carbon. All Images are of 200 × 200 μm dimensions.
2000–3500 cm\(^{-1}\) range were present but at low intensity and merged. All these Raman characteristics are typical of low HTT glassy carbon from the literature [38, 39, 42, 43, 46, 47].

For the photonic cured samples, progressing through the number of pulses emitted, the first Raman active sample was observed after 5 consecutive pulses, amounting to a total of 100 ms exposure time and \(\ast 135 \text{ J cm}^{-2}\) energy density exposure. A bulge was evident at \(\ast 1150 \text{ cm}^{-1}\) in the sample subjected to 5 pulses, but it steadily diminished with longer photonic curing treatment (Fig. 6). Carbon \(D\) and \(G\) band peaks were present in all samples, the \(D'\) peak was obscured by the \(G\) band peak suggesting a small grain size in the carbon coatings which is consistent with the relatively low carbonization temperatures used to produce these coatings [45, 46]. There was an increase in the intensity ratio between the \(D\) and \(G\) band as the number of pulses increased, starting with an \(I(D/G)\) ratio of 0.67 after 5 pulses, increasing to 0.83 after 20 pulses, and compared to 0.88 for conventional thermal treatment (Fig. 6). Carbon \(D\) and \(G\) band peaks were present in all samples, the \(D'\) peak was obscured by the \(G\) band peak suggesting a small grain size in the carbon coatings which is consistent with the relatively low carbonization temperatures used to produce these coatings [45, 46].

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The 20 pulses photonic cured sample was the most similar to the thermal cured carbon coating. The underlying individual Raman peaks were fitted and compared in more detail for the thermal cured and 20 pulses photonic cured samples in Fig. 7. Both of these spectra consisted of the same primary and secondary peaks, thus demonstrating fundamental similarities between the two resultant carbon coatings. The primary peaks were due to the \(D\) band (carbon ring breathing mode) and \(G\) band (\(sp^2\) bonded carbon stretching mode) at \(\sim 1360 \text{ cm}^{-1}\) and \(\sim 1590 \text{ cm}^{-1}\), respectively. The secondary peaks consisted of the \(2D\) band (\(\sim 2600 \text{ cm}^{-1}\)) and the \(D + D'\) band (\(\sim 2900 \text{ cm}^{-1}\)).

The \(G\) band in both carbonization methods is located at \(1590 \text{ cm}^{-1}\), the \(D\) bands are in similar positions, 1357 cm\(^{-1}\) for the thermal curing method and 1360 cm\(^{-1}\) for the photonic curing method. The full-width half maximum (FWHM) of the \(G\) band peak correlates directly to the measure of disorder, with an increase in FWHM demonstrating greater disorder [45]. The conventional method Raman spectrum \(G\) band had a FWHM of 102 cm\(^{-1}\) while the photonic curing \(G\) band gave a FWHM of 97 cm\(^{-1}\). This indicates that the materials produced from both methods had similar levels of disorder in their structures.

**XPS analysis of treated materials**

The XPS spectra of photonic cured (5, 10, 15 and 20 s) and thermal treated coatings are compared in Fig. 8 and show primary peaks at \(\sim 284–285\) eV and \(\sim 531–533\) eV binding energies. These two peaks are indicative of carbon–carbon (C1s) and carbon–oxygen (O1s) bonds which are expected for carbon material derived from an organic polymer precursor [40, 48, 49]. Figure 8, further, demonstrates the similarity of the chemical content distribution of the carbon coatings and allows the chemical distribution of each coating to be calculated using the relative areas under the peaks. Conventional method GC coatings contain \(\sim 97\%\) carbon content and \(\sim 3\%\) oxygen.
content, compared to the photonic cured coatings which contain ~96% carbon content and ~4% oxygen content. These values are in line with similar studies on commercially manufactured glassy carbon from Lim et al. (48) and Yi et al. (17).

The primary carbon peaks (C1s) of the carbon coatings are compared in Fig. 9 against graphite (exclusively $sp^2$ bonded carbon) and diamond (exclusively $sp^3$ bonded carbon), the reference materials that reside on either extreme of the carbon bonding spectrum. The location and shape of the peaks in Fig. 9 show that the thermal and photonic cured carbon peaks are located in the same position as the reference graphite sample ($sp^2$ only) at ~285.0 eV and that they are of similar shape, with a high binding energy tail indicative of the presence of $sp^2$ carbon. The $sp^3$ diamond peak is clearly located at a lower binding energy of ~283.5 eV and has a different shape with a bulge on the lower binding energy side of the peak. Unlike graphite or the glassy
carbon coatings, with their expected high sp² carbon content, and ability to resupply electrons, the diamond required charge neutralisation, and hence shifted to a lower binding energy. These results indicate that the photonic cured and thermal cured carbon have similar carbon–carbon bond structures to the graphite reference sample.

A further XPS test was performed on the samples at a higher binding energy (1190–1250 eV) (Fig. 10). In this range, carbon materials have another peak known as the Auger peak (C KLL). This Auger peak is the observation of an electron going through a unique transition between energy levels known as the Auger electron [49, 50]. With mathematical manipulations to the Auger peak data an empirically derived value known as the D parameter can be calculated. The D parameter is defined as the binding energy difference between the position of highest peak and the lowest trough of the Auger peak (C KLL) and is calculated from the first derivative between these positions, and has been used to empirically determine the sp²/sp³ bonded carbon ratio of carbon materials [48, 49]. Using the CasaXPS software, the C KLL peak data were smoothed and differentiated as suggested by Kaciulis et al. [40]. Adopting the assumption that diamond has 0% sp²/100% sp³ carbon bonds and graphite has 100% sp²/0% sp³ carbon bonds, a linear relationship between the D parameter and the sp² content percentage can be established as demonstrated in Fig. 10 (right). Using this empirical linear relationship, estimates for carbon materials with unknown sp²/sp³ ratios can be made, leading to thermal cured glassy carbon having an estimated sp² content of 96–100%, while photonic cured carbon on the other hand showed a more even sp²/sp³ split at 55 ± 4% sp².

The apparent sp² content differed depending on the analysis technique used. Using Raman spectroscopy, predominantly sp² content was observed both for photonic and thermally derived carbon coatings. However, XPS analysis indicated a mixed ~ 55% sp² structure for the photonic derived carbon and a much more exclusively sp² structure for the thermally derived carbon coatings. A key distinction between these two analysis techniques is the penetration depth. Raman spectroscopy has a penetration depth of approximately 1 micron for this type of material and is therefore representative of the bulk characteristics of the coating. On the other hand, XPS is focussed on morphology at approximately 5–10 nm from the surface, and is therefore primarily indicative of the surface. This suggests a difference in the morphology at the immediate surface of the photonically cured coating when compared to the bulk. This might be associated with the local environment around the transitioning samples. Conventional thermal conversion takes place in a consistent environment at 800 °C, with a slow and continuously purged carbonisation process. The photonic conversion occurred at close to ambient temperature, with heat directed at the sample surface and not the gas space around it. There are also differences in the rate...
at which the off-gases are produced by the different techniques, which might affect the chemical content of the gas space. This could potentially be explored with a larger sample enclosure and more aggressive purging. Despite this the photonic route offers a rapid means of producing glassy carbon, in seconds or minutes compared to hours, with potential for a host of new applications, even on temperature sensitive substrates which would be destroyed with conventional processing.

Conclusions

In this study, photonic curing was explored as a method to rapidly produce glassy carbon coatings. The conclusions are as follows:

- Photonic curing was able to convert a polymer resin precursor into a bulk glassy carbon structure which was \( sp^2 \) dominant, as indicated by Raman spectroscopy, and closely resembled a conventional derived glassy carbon structure produced from thermal treatment at 800 °C.
- Photonic curing was able to convert the precursor in around 1 min, a manifold reduction in processing time compared to the approximately 20 h processing time when using a conventional thermal conversion method.
- Carbon coatings produced photonically were resistant to bending, highly smooth and did not show any evidence of surface defects, such as pinholes, delamination or scratches.
- XPS analysis suggested a difference in the immediate surface chemistry of the photonic cured samples compared to thermally derived. This is likely associated with the different environmental conditions around the sample during treatment.
- The ability to rapidly photonically manufacture glassy carbon coatings expands the potential window of applications of glassy carbons for high volume applications such as coatings for energy storage, rapid manufacture of complex electrically conductive shapes, and the opportunity to use temperature sensitive substrates.

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Author contributions

BdeB Conceptualisation, Methodology, Investigation, Formal analysis, Writing—original draft, Visualisation; CP Methodology, Writing-Review and Editing, Supervision; YCL Methodology, Investigation; AA Conceptualisation; JM Formal analysis, Investigation; DD Conceptualisation, Methodology, Resources, Writing—review & editing, Supervision, Funding acquisition, Project administration.

Declarations

‘Yin Cheung Lau’.Conflict of interest The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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