In situ 4D tomography image analysis framework to follow sintering within 3D-printed glass scaffolds

Achintha I. Kondarage1,2 | Gowsihan Poolgasundarampillai3 | Amy Nommeots-Nomm2 | Peter D. Lee4,5 | Thilina D. Lalitharatne6 | Nuwan D. Nanayakkara7 | Julian R. Jones2 | Angelo Karunaratne1

1 Department of Mechanical Engineering, University of Moratuwa, Moratuwa, Sri Lanka
2 Department of Materials, Imperial College London, London, UK
3 School of Dentistry, Institute of Clinical Sciences, University of Birmingham, Birmingham, UK
4 Department of Mechanical Engineering, University College London, London, UK
5 Research Complex at Harwell, Didcot, UK
6 Dyson School of Design Engineering, Imperial College London, London, UK
7 Department of Electronic and Telecommunication Engineering, University of Moratuwa, Moratuwa, Sri Lanka

Correspondence
Angelo Karunaratne, Department of Mechanical Engineering, University of Moratuwa, Moratuwa, Sri Lanka. Email: angelok@uom.lk

Abstract
We propose a novel image analysis framework to automate analysis of X-ray microtomography images of sintering ceramics and glasses, using open-source toolkits and machine learning. Additive manufacturing (AM) of glasses and ceramics usually requires sintering of green bodies. Sintering causes shrinkage, which presents a challenge for controlling the metrology of the final architecture. Therefore, being able to monitor sintering in 3D over time (termed 4D) is important when developing new porous ceramics or glasses. Synchrotron X-ray tomographic imaging allows in situ, real-time capture of the sintering process at both micro and macro scales using a furnace rig, facilitating 4D quantitative analysis of the process. The proposed image analysis framework is capable of tracking and quantifying the densification of glass or ceramic particles within multiple volumes of interest (VOIs) along with structural changes over time using 4D image data. The framework is demonstrated by 4D quantitative analysis of bioactive glass ICIE16 within a 3D-printed scaffold. Here, densification of glass particles within 3 VOIs were tracked and quantified along with diameter change of struts and interstrut pore size over the 3D image series, delivering new insights on the sintering mechanism of ICIE16 bioactive glass particles in both micro and macro scales.

KEYWORDS
bioactive glass, bioceramics, image analysis, sintering, X-ray computed tomography
1 | INTRODUCTION

Additive manufacturing (AM) allows the fabrication of customized porous architectures with complex geometries and interconnected pores. This enhances the capabilities of porous biomaterials in the field of regenerative medicine, facilitating the development of architectures that cannot be produced by other methods or even patient-specific implants. Bioactive ceramics and glasses are promising for bone scaffolds due to their ability to bond with bone. Pore architecture plays a key role in promotion of vascularized bone ingrowth into the scaffold. Conventional methods of producing porous ceramics such as freeze casting, foaming, or foam reticulation provide a limited control over pore-architecture. With additive manufacturing techniques, pore-architectures can be predesigned using computer aided design (CAD), providing a better control over pore architecture. Various AM technologies have been used to fabricate porous ceramic structures, including selective laser sintering (SLS), selective laser melting (SLM), stereolithography (SLA), fused deposition modeling (FDM), inkjet printing (IJP), binder-based 3D-printing (3DP), and Direct Ink Writing (DIW). In most of these techniques, ceramic particles mixed with a binder are printed into a green body scaffold and then sintered to obtain a fully densified scaffold, which also results in shrinkage of the structure.

Even though AM can provide exceptional control over architecture, notable structural defects and discrepancies can occur between the final product and the design file. Sintering is an example of postprocessing that contributes to this discrepancy. The aim here was to enable a method of following the changes of particles such as rounding, coalescence, coarsening and residual pores across a 3D-printed scaffold during sintering and its influence on scaffold’s structure at macro scale.

Hot stage microscopy (HSM), dynamic scanning calorimetry (DSC), and differential thermal analysis (DTA) are commonly used to understand sintering of glasses and ceramics. HSM provides information on particle densification with respect to reduction of surface area during sintering, and DTA and DSC provide information on glass transition and crystallization kinetics. Combination of HSM with DSC or DTA deliver a good understanding on sintering kinetics of glasses. Furthermore, recently Hmood et al. presented that a 3D-printed pyrometric cone can be used for in situ monitoring of viscous flow and sintering-crystallization of glass ceramic systems, during heat treatments in a hot stage microscope. In situ micro-CT is an effective technique to analyze sintering within ceramic or glass structures, as it provides information on 3D morphological and structural changes occur during the sintering. 

While 4D in situ microcomputed tomography (micro-CT) can be used to capture the sintering process, quantifying these micro- and macro-structural changes requires automated image analysis pipelines. In literature automated thresholding algorithms are widely used to segment the solid phases in micro-CT images. Thresholding-based segmentation algorithms, such as Otsu’s thresholding, can be accurate in low noise micro-CT images, where mutually exclusive intensity level distribution can be obtained for solid phase and void. Imaging and reconstruction of a fast dynamic processes such as sintering of angular particles may generate more noise. Image segmentation conducted by thresholding is solely based on the intensity level of each pixel in the image and highly affected by noise. In such cases, advanced segmentation methodologies are required to segment the solid phase of the images. While accurate segmentation of solid phase is a key image processing challenge, implementation of an automated image analysis pipeline for the quantitative analysis of sintering within a 3D-printed scaffolds in both micro- and mesoscale produce more challenges including tracking VOIs and accurate detection of strut boundaries. In the study by Nommeots-Nomm et al., the quantitative image analysis was conducted based on a 3D image analysis pipeline implemented using Avizo 9.1.1 (FEI Company) and ImageJ. Due to the challenges of automating the pipeline over the complete image series, analysis was conducted with a limited set of scans and VOIs. Reducing the image series causes information loss and limiting VOIs reduces the accuracy of the quantification. Furthermore, manual selection of VOIs and quantification of descriptors in 3D image sets is tiresome and limits reproducibility.

To resolve this problem, we propose a new automated image analysis framework to analyze micro- and macrostructural changes that occur in AM bioceramic scaffolds using 4D image data solving image analysis challenges such as tracking VOIs and accurate segmentation of glassy phase. This image analysis framework incorporates machine learning and was developed using open-source tools. The proposed image analysis framework will allow studies to investigate dynamic processes that occur within these 3D-printed bioactive glass scaffolds and optimization of the AM process. The image analysis
framework is also applicable in analyzing a wide range of sintering processes.\textsuperscript{39,45–47}

To demonstrate the image analysis framework, the bioactive glass particles of the composition of ICIE16 (49.46 mol.% SiO\textsubscript{2}, 36.27 mol.% CaO, 6.6 mol.% Na\textsubscript{2}O, 1.07 mol.% P\textsubscript{2}O\textsubscript{5}, and 6.6 mol.% K\textsubscript{2}O) were chosen as it is a highly bioactive glass that can be fabricated into amorphous porous scaffolds using AM techniques such as DIW.\textsuperscript{26} Commercially available bioactive glasses such as the 45S5 Bioglass\textsuperscript{⃝} and S53P4 tend to crystallize on sintering, so are not appropriate for producing glass scaffolds. Moreover, ICIE16 composition shows promising osteogenic properties with respect to crystallized 45S5. Moreover, KONDARAGE et al.\textsuperscript{48} Bioglass, which should mean that 13–93 is less bioactive than 45S5 in terms of dissolution rate and apatite formation. Here, ICIE16 is of interest as it has a similar network connectivity to 45S5 but can be sintered without crystallization as the temperature difference between the glass transition temperature and the crystallization temperature is sufficient for sintering.\textsuperscript{26,41}

This is the first 4D analysis of ICIE16 bioactive glass sintering conducted using synchrotron sourced X-ray computed tomography. Previous study on four-dimensional quantitative analysis of sintering within a 3D-printed bioactive glass by Nommets-Nomm et al. was conducted for the 13–93 (54.6 SiO\textsubscript{2}, 22.1 CaO, 6.0 Na\textsubscript{2}O, 1.7 P\textsubscript{2}O\textsubscript{5}, 7.9 K\textsubscript{2}O, and 7.7 MgO, in mol%) composition.\textsuperscript{41} The 13–93 composition has a higher network connectivity (mean number of bridging oxygen bonds per silicon atom) than 45S5 Bioglass, which should mean that 13–93 is less bioactive than 45S5 in terms of dissolution rate and apatite formation. Here, ICIE16 is of interest as it has a similar network connectivity to 45S5 but can be sintered without crystallization as the temperature difference between the glass transition temperature and the crystallization temperature is sufficient for sintering.\textsuperscript{26,41}

In this study, quantitative analysis conducted using the implemented image processing pipeline delivers new insights about the sintering of ICIE16 bioactive glass related to particle densification, sintering stages, shrinkage, and the morphological changes happening within strut of 3D-printed scaffolds during sintering. Importantly, the proposed image analysis framework facilitates implementing automated, validated, and reproducible pipelines to analyze 3D-printed porous glasses and ceramics in 4D.

2 | EXPERIMENTAL PROCEDURE

2.1 | Scaffold preparation

ICIE16 bioactive glass scaffolds were prepared using the protocol presented by Nommets-Nomm et al.\textsuperscript{26} Glass particles were manufactured via melt quenching using high purity silica (SiO\textsubscript{2}) (High Purity, Prince Minerals, Stoke-on-Trent), phosphorous pentoxide (P\textsubscript{2}O\textsubscript{5}), and the carbonate equivalent of the required modifying oxides. The glass was melted at 1400°C for 2 h, in a 95% platinum 5% gold crucible, quenched into deionized water, and dried at 100°C. Glass frit was ground in a Ball mill to produce particles with a size distribution ranging from 3.3 to 30.5 µm and D\textsubscript{50} = 10.8 µm was used (measured by a Malvern Mastersizer 2000, Malvern Instruments Ltd. UK). Inks for 3D printing were produced with a 25 wt% Pluronic F-127 solution (CAS: 9003-11-6) and mixed with glass particles (47.5 vol% glass to a Pluronic solution, calculated using the relative glass densities) using a Thinky ARE-100 mixer until homogenized. Finally, scaffolds were printed using DIW, by a 3D Robocaster (RoboCAD 3.0, 3-D Inks, Still-water) with a 250 µm diameter conical nozzle, during the printing a z layer spacing of 200 µm was used (80% of the nozzle diameter). It was found in previous work that 80% is key to insuring layer-on-layer adhesion of struts, and adherence of the first layer to the printing substrate.\textsuperscript{26,49}

During the printing, the humidity was controlled at 60–80% with a 23°C temperature. Prior to sintering, the size of the printed green body scaffold was 1.8 mm × 1.8 mm × 1.8 mm.

2.2 | Imaging and image reconstruction

Viscous flow sintering of the glass particles within 3D-printed scaffolds were imaged in situ using synchrotron sourced X-ray computed tomography at the Diamond-Manchester Imaging Branchline I13-2 of Diamond Light Source. The optimum sintering temperature for ICIE16 was fixed at 690°C based on previous studies carried out for similar particle sizes.\textsuperscript{26} Sintering was performed in the bespoke proportional-integral-derivative-controlled “Laura” furnace.\textsuperscript{41,50–53} Here, the glass scaffolds were glued on an alumina sample holder using a high-temperature glue (OMEGABOND 600; Omega LTD, UK) and alumina holder was mounted on to a rotating spindle on the sample stage. Sintering was carried out in two phases: printed samples were heated up to 500°C at a rate of 3°C per minute and held at 500°C for a dwell 1 h to ensure complete removal of binder (Phase I); then samples were heated at a rate of 3°C per minute up to the sintering temperature 690°C and held for 2 h (Phase II). Any remaining binder would be removed quickly during the ramp up to the sintering temperature. Images were captured throughout the sintering process, and those captured during Phase II were analyzed in this study. The X-ray microtomography was performed with a filtered pink polychromatic beam in the energy range of 8 to 30 keV. The emitted beam was captured using a CMOS detector with a resolution of 2560×2160, which was positioned 75 mm behind the sample stage. All scans were performed with a total magnification of 8x, resulting in an effective isotropic pixel size of 0.81 µm. With this setting, projections were taken with an angular step size of 0.09 (2001 projections
per 180° for 70 min from the start of Phase II of the sintering, and the rest of the projections were acquired with an angular step size of 0.15° (1201 projections per 180°). Projections were acquired with an exposure time of 45 ms. All the projections were reconstructed using filtered back projection (FBP), incorporating dark and flat field correction and ring artifact suppression.

2.3 Development of image analysis framework

The image analysis framework was developed using ImageJ API and Skimage, Sklearn python libraries. The complete set of python scripts for the image analysis framework is available in the provided online repository (https://github.com/AchinthaIroshan/Sintering-within-3D-printed-ceramics). An image analysis pipeline was implemented using the proposed framework to understand the sintering of ICIE16 bioactive within a 3D-printed scaffold. Here, 93 3D-reconstructed images captured during the sintering process were analyzed.

2.3.1 Preprocessing, tracking volumes of interest (VOIs), and denoising of images

Initially, all the slices of 3D-reconstructed images (example: Figure 1A) were converted to 8 bits. Then initial preprocessing was conducted by rotating and cropping each 3D image (Figure 1B). Here, the rotation was conducted such that the struts of the scaffold were aligned with \( y \) and \( z \) directions of the image. Struts can be considered as the building element of the scaffold (Figure 2). Therefore, cropped 3D slices of struts with a thickness of 57.6 \( \mu \text{m} \) (example: Figure 2B) were taken as volumes of interest (VOIs) to analyze sintering.

VOIs of each image were detected using an algorithm propose with the image analysis framework. Here, the VOIs were detected by automated identification of centers of those volumes (centers of selected strut slices). The main steps of the developed algorithm are described below.

- **Resampling of the 3D image with a factor of 0.25** (Figure 1C): Here the size of the image is reduced to 1/4th of the original size, which makes it easier to compute macroscale features to detect VOIs.

- **Detecting image slices, which include VOIs using a Support Vector Machine (SVM) classifier** (Figure 1D): The 3D image can be represented as a stack of 2D images where the 2D image is called as an image slice. A set of image slices labeled with two classes (Class 1—Slices that include VOIs, Class 2—Slices that do not include VOIs) were used as training data to train a Support Vector Machine classifier, which is a machine learning classification model. The trained model was used to detect slices that include VOIs in each 3D image.

- **Segmenting regions with strut cross-sections in each detected slice using “Level-set” algorithm** (Figure 1E, F): In “level-set” segmentation algorithm, an implicit contour is fitted to boundaries of regions of interest (Figure 2E).

- **Labeling each connected component in the 3D image** (Figure 1G): Here, each connected component is considered as a VOI in the resampled 3D image. Each connected component is labeled to obtain their centers separately.

- **Calculating the center coordinates of each labeled 3D-connected component:** regionprops of Skimage was used to calculate the center.

- **Converting the calculated coordinates in coordinates of the original size image:** As volumes of interests in the original images are considered for the analysis, center coordinates of VOIs in the resampled 3D image were converted to the coordinates of the original size image.

- **Obtaining VOIs based on the center coordinate** (Figure 2H, I): For each obtained center coordinate, volumes of dimensions 150 \( \times \) 140 \( \times \) 70 (Figure 2H) and volumes of dimensions 150 \( \times \) 140 \( \times \) 70 (Figure 2I) were generated at each time point to quantify diameter and densification, respectively.

4D hyperstacks (3D volume series) of each VOI were generated by combining 3D volumes based on Euclidian distance between center coordinates of volumes in consecutive time points. Based on the obtained center information, 27 VOIs can be obtained from each 3D image, and 27 4D hyperstacks can be generated from the complete scan set captured during the sintering. Here 3 VOIs from each 3D image were taken for the analysis.

Nonlocal mean filtering was applied to further noise reduction, which uses redundant information of the image to reduce the noise by performing a weighted average of pixel values considering spatial and intensity similarities between pixels. Following the noise-reduction contrast of images was enhanced using histogram equalization.

2.3.2 Segmentation of strut and glassy phase in VOIs

The implemented image analysis pipeline to characterize sintering in the microscale consists of two main segmentation components. One of those components is segmenting the glassy phase within a strut to analyze densification during sintering. A random forest classifier was
applied to segment the glass phase until glass densification reached about 90%. The fast random forest classification is a machine learning method, which can be used for image segmentation. In this work, it was implemented incorporating the Trainable Weka Segmentation module of Fiji\textsuperscript{59} and trained to classify every voxel to glassy class or void class. A total of 162 792 labeled voxels were used in training the model, where 96 323 voxels were labeled into pore class and 66 469 into glassy class. During the model training, each labeled voxel was represented using 77 attributes based on Gaussian blur, Hessian matrix, Difference of Gaussian, Sobel filters, and directional filtering. Here, a random forest classifier with 200 trees was constructed. Once the densification of the glassy phase reached around 90%, images were segmented using Otsu’s thresholding method.\textsuperscript{60} The second segmentation component of the image analyzing pipeline is segmenting the strut without considering intrastrut porosity to analyze the diameter change of struts during the sintering. Here, a child volume of size $330 \times 306 \times 10$ was obtained from the VOIs of each 3D image without changing the center of the VOI. Then, strut regions of each image slice were segmented using the level-set algorithm to evaluate the change of strut diameter during the sintering.
The performance of image segmentation was evaluated by calculating the accuracy, precision, and Intersect over Union (IoU) values of each segmentation method. These performance measures were calculated based on a set of manually segmented images that were taken as ground truth segmentations.

2.3.3 Quantitative analysis of ICIE16 bioactive glass sintering

Three parameters were quantified for the quantitative analysis. Those are the relative densification of the glassy phase, change of diameter of the struts, and change of longest diagonal length of the interstrut pores during the sintering. Here, a complete scan set with a size of 93, captured during the second stage of the sintering, was analyzed.

To quantify the relative densification of glass at the time of a 3D scan was acquired, segmented binary volumes of $150 \times 140 \times 70$ (Figure 1J) were obtained from the detected VOIs of $330 \times 306 \times 70$, without changing the center of the volumes. Glass volume and pore volume in each VOI were calculated by counting corresponding voxels. The intrastrut porosity and relative density of the glassy phase were calculated using Equations (1) and (2).

$$\text{Porosity} \,(\%) = \frac{\text{Pore Volume}}{\text{Pore Volume} + \text{Glass Volume}} \times 100, \quad (1)$$

$$\text{Relative Density of Glassy Phase} \,(\%) = 100 – \text{Porosity}. \quad (2)$$

The interstrut pore diagonal distance (PDD) at each time point of a scan was calculated using Equation (4),

$$\text{Diameter} = 2 \times \sqrt{\frac{\text{Area}}{\pi}}. \quad (3)$$

$$\text{PDD} = \text{Distance between strut centers diagonally}$$

$$- \text{Diameter of struts}, \quad (4)$$

where the distance between strut centers placed in directions of pore diagonals was obtained as illustrated in Figure 1L.

The strut diameter of each VOI was obtained (3 VOIs) and pore diagonal distance was obtained considering 3 interstrut pore channels in each 3D image. The averages of those automated measurements were obtained as an estimator for the strut diameter and pore diagonal distance of the scaffold at the time point of the scan. As the calculation are obtained in pixels, measurements were rounded to the nearest integer and multiplied by 0.81 (the isotropic pixel size) to convert into micrometers.

Finally, strut diameter values, the relative densification of glass phase, and the longest diagonal length of
interstrut pores were plotted against temperature and time for the complete 3D image series representing the sintering of ICIE16 bioactive glass within a 3D-printed scaffold.

3 | RESULTS AND DISCUSSION

3.1 | Image analysis framework

The accuracy of 4D image quantification is highly reliant on the image analysis pipeline. Therefore, image analysis plays a vital role in the characterization of dynamic processes using 4D imaging. In this study, the relative density of the glassy phase, the diameter of the strut and interstrut pore diagonal were quantified as descriptors to analyze the sintering within a 3D-printed scaffold both micro- and macroscales.

3.1.1 | Detection of VOIs and tracking

Based on the visualization of the series of 93 3D images (Video S1), it was identified that VOIs considered for the analysis were not static over time due to the shrinkage of the scaffold. Manual identification of VOIs in each 3D image is a challenging and time-consuming task. Therefore, the proposed algorithm to detect and track VOIs is a vital component of the image analysis pipeline. Dynamic visualization of VOIs obtained from the VOI detection and tracking algorithm is included in Video S2.

3.1.2 | Noise reduction

Noise reduction of the detected child volume highly affects the accuracy of the image segmentation. Notable noise reduction of VOIs was achieved by applying nonlocal means filter. Resulting images (Figure 3A, B), histograms (Figure 3C, D), and intensity profiles of lines drawn on top of original and resulting images (Figure 3E, F) provide clear evidence of noise reduction. As shown in the histograms (Figure 3C, D), even after the noise reduction, intensity values of glass and pore voxels could not be observed as mutually exclusive sets.

3.1.3 | Image segmentation and quantification

As mutual exclusive intensity distribution was not obtained with the noise reduction, random forest classification, based on machine learning, was used to segment the glassy phase of VOIs. In random forest classification, several image features are used for the image segmentation. The performance Fast Random Forrest classification methods to segment glassy phase was evaluated with respect to Otsu’s thresholding method, using intersection over union (IoU), accuracy, and precision (Table 1). Fast random forest delivered a better performance over Otsu’s thresholding in segmentation of glassy phase for the images captured within the first 1 h and 15 min of sintering (early sintering stages) where mutually exclusive intensity level distribution was not obtained for glass and void. Figure 4A–C visualizes the results of segmentation obtained with fast random forest classifier. The out-of-bag error of the fast random forest classifier was 4.88%. After 1 h and 15 min of sintering, the densification of the glassy phase reached about 90% (Figure 4D). At this stage, mutually exclusive intensity levels can be observed for the glassy phase and void (Figure 4E). Therefore, Otsu’s thresholding provided good results for glassy phase segmentation (Figure 4F) of those images with high IoU, accuracy, and precision (Table 1).

To quantify the diameter of struts, boundaries of the struts should be accurately detected. Due to the noise and the intrastrut porosity (Figure 4G), accurate identification of strut boundaries is a challenging task. “Level-set” segmentation was identified as the most suitable algorithm.
TABLE 1  Performance evaluation of the image segmentation

| Volume of interest                                | Segmentation algorithm | Intersection over union (IoU) | Accuracy | Precision |
|---------------------------------------------------|------------------------|-------------------------------|----------|-----------|
| Glassy phase within a strut (relative densification ≤90%) | Fast random forest     | 0.83                          | 0.93     | 0.91      |
|                                                   | Otsu’s thresholding    | 0.78                          | 0.88     | 0.79      |
| Glassy phase within a strut (Relative densification ≥90%) | Otsu’s thresholding    | 0.98                          | 0.98     | 0.99      |
| Strut slice, excluding pores within the strut     | Level-set segmentation | 0.96                          | 0.97     | 0.99      |

FIGURE 4  Image segmentation of micro-CT images. (A) Preprocessed image slice of a VOI obtained to quantify relative densification where individual glass particles are visible. (B) Segmented image of glassy phase using fast random forest voxel classification. (C) Segmented image volume considered to calculate the relative densification of the glassy phase. (D) Section of a preprocessed image captured after reaching a higher densification (around 90%). (E) Histogram of the image (D). (F) Image of the glassy phase segmented using Otsu’s thresholding. (G) Preprocessed image slice of a VOI, obtained to quantify diameter change of struts. (H) Contour of level-set function fitted to the boundaries of the strut cross-section. (I) Segmented strut cross-section obtained using the level-set segmentation.
to detect the boundaries of the strut accurately. “Level-set” segmentation is carried out by fitting an implicit contour to strut boundaries (Figure 4H), resulting in a good segmentation (Figure 4I) with high IoU, accuracy, and precision (Table 1).

In the implemented image analysis pipeline, the segmentation is followed by the quantification descriptors to analyze sintering. The results of quantification and insight delivered on sintering are discussed in the next section. Furthermore, the presented image analyzing methodologies are not limited to analyzing the sintering within 3D-printed scaffolds. These techniques can also be used to analyze the structural changes, which occur in scaffolds designed using woodpile architecture and to analyze the densification of different particles within solid structures, incorporating 4D imaging.

3.2 Sintering analysis

In this work, changes in relative density of the glassy phase during the sintering were quantified as a descriptor for the microscale analysis of sintering, and both diameter and the interstrut pore diagonal distance were quantified as descriptors for macroscale analysis of sintering.

According to the classical sintering theory on densification, powder compaction is described with three defined stages. During the initial stage of the sintering, neck formation occurs between particles resulting around 10% of densification. During the intermediate stage, bulk densification occurs, collapsing interconnected pores into isolated pores reaching approximately 90% relative densification. During the final stage, size of closed pores reaches toward zero resulting 0% porosity and complete densification.61,62

The graph (Figure 5) shows the change of relative density of the glassy phase over time and temperature. As presented in the graph, three stages of densification were visible with the ICIE16 bioactive glass sintering cycle, confirming the classical sintering theory. Here, ~5% of densification occurred during the initial sintering stage, ~35% of fast densification happened during the intermediate sintering stage, and ~5% of slow densification during the fast sintering was observed. Furthermore, microstructural changes within a strut were observed by visualization of segmented child volumes of VOIs at different stages of sintering (Figure 6). Closely packed glass particles were observed within struts of the scaffold after the removal of binder at the temperature of 500°C (Figure 6A, Time 00:00:00). No notable morphological changes of glass particles were observed for the first 45 min with gradual increase of temperature to 636.8°C (Figure 6B, Time 00:45:21). Once the temperature reached 667.2°C, neck growth between glass particles was observed (Figure 6C, Time 00:55:26), which is a feature of the initial stage of sintering according to classical sintering theory. During the intermediate sintering stage, interconnected pores were present (Figure 6D, Time 01:15:21). Within a very short time (12 min), these interconnected pores transformed into isolated pores (Figure 6E, Time 01:15:30). During the final sintering stage, the isolated pores were further reduced. A set of tiny (less than 20 µm in size) isolated pores were observed at the end of the 3-h sintering cycle (Figure 6F, Time 03:02:15). These observations confirm that the densification of ICIE16 bioactive glass follows classical sintering theory. Nommets-Nomm et al. previously showed that the sintering of glass particles of the 13–93 composition within a 3D-printed scaffold also followed classical sintering theory.41

Figure 7 visualizes changes in a slice or section of a strut (VOI) within the scaffolds during the different sintering stages, which indicate the nature of a strut at the start of the sintering (Figure 7A, Time 00:00:00), showing individual particles and cylindrical morphology of the strut. At the intermediate stage of the sintering (Figure 7B, Time: 01:00:29), shrinkage of the strut occurred, and at the end of sintering (Figure 7C, Time: 03:02:15), a densified strut with a smooth surface was observed at the end of the sintering. Here (Figure 7), a notable reduction of strut diameter (15%) can also be observed.

The glass particles are packed within the strut of the scaffold. If we assume that the shrinkage of the strut due to the sintering of glass particle is isotropic, the relationship between the volume shrinkage and the linear shrinkage is as follows:63

\[
\text{Volume Shrinkage} = 1 - (1 - \text{linear shrinkage})^3.
\]  

The graph (Figure 8A) shows the change of the strut diameter during the sintering. Initial diameter of a strut was
241.4 μm. Once the temperature reached 660°C, the diameter rapidly reduced from 239.8 to 207.4 μm within 23 min as a result of bulk densification during the intermediate sintering stage. The diameter further reduced at 690°C and reached a minimum of 203.3 μm. The graph shows that the strut diameter remained at 204.1 ± 0.8 μm during the final stages of sintering. The change of pore diagonal distance also shows a pattern similar diameter reduction and shows a continuous decreasing (Figure 8B). The pore diagonal distance of the printed scaffold was 294.0 μm. Same as with the strut diameter, once the temperature reached 660°C, pore diagonal distance rapidly reduced from 288.4 to 239.0 μm within 23 min. The diagonal distance continues to decrease at 690°C, reaching a minimum of 226.0 μm.

The reduction of the diameter was 15.8%. If we consider the reduction of the diameter as linear shrinkage, the volume shrinkage can be calculated using Equation 5, giving a value of 40.4%. To evaluate whether the shrinkage of the scaffold was isotropic, volume shrinkage was calculated based on the child volumes shown in Figure 9. The shrinkage of the considered child volume was 46.7%, which proves that the volume shrinkage was not isotropic. This confirms the shrinkage results published by Nommeots-Nomm et al. using bulk measurements, which showed that linear shrinkage in the z direction was less than that of the scaffold in x and y directions. To develop patient-specific scaffolds, the shrinkage should be predetermined and incorporated into initial design. As this analysis suggests that shrinkage is not isotropic, more advanced models should be implemented to predict the shrinkage.

The global scale sintering within a 3D-printed scaffold occurs due to the coarsening of struts of the scaffold that appears at stage 3 of particle sintering. This was initially observed by Nommeots-Nomm et al. during the sintering of 13–93 bioactive glass 3D-printed scaffold. In their work, they identified a steady increase of strut diameter and a steady decrease of the longest pore diagonal length during stage 3 of sintering. This happened as a result of the coarsening of individual struts, which can be identified as a sintering at a global level. However, this change was not clearly observed (Figure 8) with ICIE16 bioactive glass composition. Furthermore, the z-layer spacing was set as 200 μm in the initial printing parameters and it was maintained in the printed green body scaffold (Figure 9Aii), resulting in a strut overlap of ~41 μm (~17% of strut diameter), and it was observed that z-layer spacing reduced to 160 μm (Figure 9Bii), resulting in a strut overlap of ~49 μm (~21% of strut diameter). The strut overlap was calculated based on the measurement taken from images shown in Figure 9Aii and Bii. This ~4% decrease of strut overlap is also not a strong indication for the coarsening of struts due...
FIGURE 7  VOIs of micro-CT images obtained at different stages of sintering: (A) (i) Segmented slice of a strut considered in sintering analysis, captured after the complete burn out of the binder. (B) (i) Segmented strut slice during the fast sintering stage. (C) (i) Segmented strut slice after the completion of the sintering process (A) (ii), (B) (ii), and (C) (ii) are cross-sections of (A) (i), (B) (i), and (C) (i) sequentially.

FIGURE 8  Quantification of changes in the global-level 3D-printed scaffold architecture as a function of sintering time, by image analysis: (A) Change of strut diameter. (B) Change of pore diagonal distance.

to global scale sintering. Yet, we cannot avoid conclude that there is no global scale sintering in ICIE16 bioactive glass.

4 | CONCLUSIONS

We developed an image analysis framework incorporating machine learning to investigate the viscous sintering of particles with randomly distributed particle size (nonidealized model) within 3D-printed scaffolds using synchrotron X-ray tomography. The image analysis framework was validated and able to automate four-dimensional analysis of sintering within 3D-printed structures. To demonstrate the developed image analysis framework, we carried out a 4D automated micro- and macroscale analysis of ICIE16 bioactive glass sintering within a 3D-printed scaffold.
designed for bone tissue engineering for the first time. The densification of bioactive glass particles was quantified during the sintering considering multiple VOIs over time and its influence on scaffold’s structure was identified. Shrinkage was found to be anisotropic and the degree of sintering must be included in the 3D design files for the scaffolds in the future, especially for patient-specific scaffolds. Sintering of the ICIE16 composition follows classical sintering theory and left some residual spherical pores within the struts. The findings of this research will be helpful in the implementation of a computational model to predict the shrinkage of bioactive glass sintering, which will enable producing patient-specific scaffolds with designable geometries. Furthermore, the proposed image analysis framework is not limited to synchrotron tomography data and can be applied with other 4D imaging modalities which can capture microscale dynamics.

ACKNOWLEDGMENTS
This research was funded by the NIHR Global Health Research (grant number 1613745; NIHR Group on PoS T Conflict Trauma in Sri Lanka; PrOTeCT). The 3D printer used in this work was funded via an EPSRC Grant for Graphene 3D networks (EP/K01658X/1). Funding for A.N.-N. was provided by RCaH-EP/I02249X/1, and the “Laura” furnace was developed under the grant NE/M013561/. GP acknowledges funding from the EPSRC grant EP/M023877/1 and PDL acknowledges funding from the MRC ImagingBioPro grant (MR/R025673/1), the Research Complex at Harwell and Royal Academy of Engineering (CiET1819/10). The experiment was performed on the Branchline I13-2 of the Diamond Light Source synchrotron in Oxfordshire, UK, and partly funded by the MT13241: Collaboration Proposal 1809.

ORCID
Achintha I. Kondarage https://orcid.org/0000-0003-2992-8980
Julian R. Jones https://orcid.org/0000-0002-2647-8024
Angelo Karunaratne https://orcid.org/0000-0001-7117-0670

REFERENCES
1. Du X, Fu S, Zhu Y. 3D printing of ceramic-based scaffolds for bone tissue engineering: an overview. J Mater Chem B. 2018;6(27):4397–412.
2. Jones JR. Review of bioactive glass: from Hench to hybrids. Acta Biomater. 2013;9(1):4457–86.
3. Baino F, Novajra G, Vitale-Brovarone C. Bioceramics and scaffolds: a winning combination for tissue engineering. Front Bioeng Biotechnol. 2015;3(3):1–17.

4. Shi X, Nommets-Nomm A, Todd NM, Devlin-Mullin A, Geng H, Lee PD, et al. Bioactive glass scaffold architectures regulate patterning of bone regeneration in vivo. Appl Mater Today. 2020;20:100770.

5. Cahill S, Lohfeld S, McHugh PE. Finite element predictions compared to experimental results for the effective modulus of bone tissue engineering scaffolds fabricated by selective laser sintering. J Mater Sci Mater Med. 2009;20(6):1255–62.

6. Liu X, Rahaman MN, Fu Q, Tomsia AP. Porous and strong bioactive glass (13-93) scaffolds prepared by unidirectional freezing of camphene-based suspensions. Acta Biomater [Internet]. 2012;8(1):415–23. Available from: http://doi.org/10.1016/j.actbio.2011.07.034

7. Jones JR, Ehrenfried LM, Hench LL. Optimising bioactive glass scaffolds for bone tissue engineering. Biomaterials. 2006;27(7):964–73.

8. Chen QZ, Thompson ID, Boccaccini AR. 45S5 Bioglass®-derived glass-ceramic scaffolds for bone tissue engineering. Biomaterials. 2006;27(11):2414–25.

9. Hwa LC, Rajo S, Noor AM, Ahmad N, Uday MB. Recent advances in 3D printing of porous ceramics: a review. Curr Opin Solid State Mater Sci. 2017;21(6):323–47.

10. Champion E, Magnaudeix A, Pascaud-Mathieu P, Chartier T. Chapter 14—advanced processing techniques for customized ceramic medical devices. In: Palermo P, Cambier F, De Barra E, editors. Advances in ceramic biomaterials. Woodhead Publishing; 2017. p. 433–68. Available from: https://www.sciencedirect.com/science/article/pii/B9780081008812000154

11. Shahzad K, Deckers J, Boury S, Neirincck B, Kruth JP, Vleugels J. Preparation and indirect selective laser sintering of alumina/PA microspheres. Ceram Int. 2012;38(2):1241–7.

12. Yves-Christian H, Jan W, Wilhelm M, Konrad W, Reinhart P. Net shaped high performance oxide ceramic parts by selective laser melting. Phys Procedia. 2010;5(2):878–94.

13. Wilkes J, Hagedorn YC, Ocylok S, Meiners W, Wissenbach K. Rapid manufacturing of ceramic parts by selective laser melting. Ceram Eng Sci Proc. 2010;31(8):137–48.

14. Altun AA, Prochaska T, Konegger T, Schwentenwein M. Dense, strong, and precise silicon nitride-based ceramic parts by lithography-based ceramic manufacturing. Appl Sci. 2020;10(3).

15. Zanchetta E, Cattaldo M, Franchin G, Schwentenwein M, Homa J, Brusatin G, et al. Stereolithography of SiOC ceramic micro-components. Adv Mater. 2016;28(2):370–6.

16. Xu N, Ye X, Wei D, Zhong J, Chen Y, Xu G, et al. 3D artificial bones for bone repair prepared by computed tomography-guided fused deposition modeling for bone repair. ACS Appl Mater Interfaces. 2014;6(17):14952–63.

17. Yang H, Yang S, Chi X, Evans JRG. Fine ceramic lattices prepared by extrusion freeforming. J Biomed Mater Res—Part B Appl Biomater. 2006;79(1):116–21.

18. Darsell J, Bose S, Hosick HL, Bandyopadhyay A. From CT scan to ceramic bone graft. J Am Ceram Soc. 2003;86(7):1076–80.

19. Bose S, Suguiria S, Bandyopadhyay A. Processing of controlled porosity ceramic structures via fused deposition. Scr Mater. 1999;41(9):1009–14.

20. Shi Y, Wang W. 3D inkjet printing of the zirconia ceramic implanted teeth. Mater Lett. 2020;261:127131. Available from: https://doi.org/10.1016/j.mattlet.2019.127131

21. Nogueira R, Lejeune M, Chartier T. 3D fine scale ceramic components formed by ink-jet prototyping process. J Eur Ceram Soc. 2005;25(12):2055–9.

22. Brunello G, Sivolella S, Meneghello R, Ferroni L, Gardin C, Pietelli A, et al. Powder-based 3D printing for bone tissue engineering. Biotechnol Adv. 2016;34(5):740–53.

23. Butscher A, Bohner M, Hofmann S, Gauckler L, Müller R. Structural and material approaches to bone tissue engineering in powder-based three-dimensional printing. Acta Biomater [Internet]. 2011;7(3):907–20. Available from: http://doi.org/10.1016/j.actbio.2010.09.039

24. Tarafder S, Balla VK, Davies NM, Bandyopadhyay A, Bose S. Microwave-sintered 3D printed tricalcium phosphate scaffolds for bone tissue engineering Solaiman. J Tissue Eng Regen Med. 2013;7:631–41.

25. Fu Q, Saiz E, Tomsia AP. Direct ink writing of highly porous and strong glass scaffolds for load-bearing bone defects repair and regeneration. Acta Biomater [Internet]. 2011;7(10):3547–54. Available from: http://doi.org/10.1016/j.actbio.2011.06.030

26. Nommets-Nomm A, Lee PD, Jones JR. Direct ink writing of highly bioactive glasses. J Eur Ceram Soc. 2018;38(3):837–44.

27. Tang S, Yang L, Liu X, Li G, Jiang W, Fan Z. Direct ink writing additive manufacturing of porous alumina-based ceramic cores modified with nanosized MgO. J Eur Ceram Soc. 2020;40(15):5758–66.

28. Souza MT, Crovace MC, Schröder C, Eckert H, Peitl O, Zanotto ED. Effect of magnesium ion incorporation on the thermal stability, dissolution behavior and bioactivity in Bioglass-derived glasses. J Non Cryst Solids [Internet]. 2013;382:57–65. Available from: http://doi.org/10.1016/j.jnoncrystol.2013.10.001

29. Massera J, Mayran M, Rogerelle J, Hupa L. Crystallization behavior of phosphate glasses and its impact on the glasses’ bioactivity. J Mater Sci. 2015;50(8):3091–102.

30. Panna W, Wyszomirski P, Kohut P. Application of hot-stage microscopy to evaluating sample morphology changes on heating. J Therm Anal Calorim. 2016;125(3):1053–9.

31. Goel A, Shaaban ER, Oliveira JB, Sá MA, Pascual MJ, Ferreira JMF. Sintering behavior and devitrification kinetics of iron containing clinopyroxene based magnetic glass-ceramics. Solid State Ionics. 2011;186(1):59–68.

32. Goel A, Kapoor S, Rajagopal RR, Pascual MJ, Kim HW, Ferreira JMF. Alkali-free bioactive glasses for bone tissue engineering: a preliminary investigation. Acta Biomater [Internet]. 2012;8(1):361–72. Available from: http://doi.org/10.1016/j.actbio.2011.08.026

33. Silva LD, Rodrigues AM, Rodrigues ACM, Pascual MJ, Durán A, Cabral AA. Sintering and crystallization of SrO-CaO-B2O3-SiO2 glass-ceramics with different TiO2 contents. J Non Cryst Solids. 2017;473(May):33–40.

34. Hmood FJ, Schmidt F, Goerke O, Günster J. Investigation of chemically modified ICIE16 bioactive glass, Part II. J Ceram Sci Technol. 2020;11(1):1–9.

35. Hmood FJ, Wilbij G, Nicolaides D, Zocca A, Günster J. An approach to monitor the real-time deformation during heat treatment of 3D-printed glass. Ceram Int [Internet].
36. Xu F, Hu XF, Miao H, Zhao JH. In situ investigation of ceramic sintering by synchrotron radiation X-ray computed tomography. Opt Lasers Eng. 2010;48(11):1082–8.

37. Bernard D, Gendron D, Heintz JM, Bordère S, Etourneau J. First direct 3D visualisation of microstructural evolutions during sintering through X-ray computed microtomography. Acta Mater. 2005;53(1):121–8.

38. Bouttes D, Lambert O, Claireaux C, Woelffel W, Dalmas D, Gouillart E, et al. Hydrodynamic coarsening in phase-separated silicate melts. Acta Mater. 2015;92:233–42.

39. Lhuissier P, Jauffre D, Lou S, Martin CL, Laboure S. Fast in situ 3D nanoimaging: a new tool for dynamic characterization in materials science. Mater Today. 2017;00(00):6–11.

40. Bernard D, Guillon O, Combaret N, Plougonven E. Constrained sintering of glass films: microstructure evolution assessed through synchrotron computed microtomography. Acta Mater [Internet]. 2011;59(16):6228–38. Available from: http://doi.org/10.1016/j.actamat.2011.06.022

41. Nommeots-Nomm A, Ligorio C, Bodey AJ, Cai B, Jones JR, Lee PD, et al. Four-dimensional imaging and quantification of viscous flow sintering within a 3D printed bioactive glass scaffold using synchrotron X-ray tomography. Mater Today Adv. 2019;2:100011.

42. Chaijaruwanich A, Dashwood RJ, Lee PD, Nagaumi H. Pore evolution in a direct chill cast Al-6 wt.% Mg alloy during hot rolling. Acta Mater. 2006;54(19):5185–94.

43. Jones JR, Poologasundarampillai G, Atwood RC, Bernard D, Lee PD. Non-destructive quantitative 3D analysis for the optimisation of tissue scaffolds. Biomaterials. 2007;28(7):1404–13.

44. Lee PD, Hunt JD. Hydrogen porosity in directional solidified aluminium-copper alloys: in situ observation. Acta Mater. 1997;45(10):4155–69.

45. Wadsworth FB, Vasseur J, Schauroth J, Llewellin EW, Dobson KJ, Havard T, et al. A general model for welding of ash particles in volcanic systems validated in situ in X-ray tomography. Earth Planet Sci Lett [Internet]. 2019;525:115726. Available from: https://doi.org/10.1016/j.epsl.2019.115726

46. Wadsworth FB, Vasseur J, Llewellin EW, Dobson KJ, Colombier M, Von Aulock FW, et al. Topological inversions in coalescing granular media control fluid-flow regimes. Phys Rev E. 2017;96(3):1–6.

47. McDonald SA, Holzner C, Laursen EM, Reischig P, Merkle AP, Withers PJ. Microstructural evolution during sintering of copper particles studied by laboratory diffraction contrast tomography (LabDCT). Sci Rep. 2017;7(1):1–11.

48. Westhauser F, Hohenfeld F, Arango-Ospina M, Schmitz SI, Wilkesmann S, Hupa L, et al. Bioactive glass (BG) ICIE16 shows promising osteogenic properties compared to crystallized 4S5S-BG. Int J Mol Sci. 2020;21(5):1–12.

49. Feilden E, Blanca EGT, Giuliani F, Saiz E, Vandeperre L. Robocasting of structural ceramic parts with hydrogel inks. J Eur Ceram Soc. 2016;36(10):2525–33.

50. Puncerebutr C, Lee PD, Hamilton RW, Phillion AB. Quantitative 3D characterization of solidification structure and defect evolution in Al alloys. JOM. 2012;64(1):89–95.

51. Kareh KM, Lee PD, Atwood RC, Connolley T, Gourlay CM. Revealing the micromechanisms behind semi-solid metal deformation with time-resolved X-ray tomography. Nat Commun. 2014;5:1–7.

52. Karagadde S, Lee PD, Cai B, Fife JL, Azeem MA, Kareh KM, et al. Transgranular liquation cracking of grains in the semi-solid state. Nat Commun. 2015;6.

53. Cai B, Lee PD, Karagadde S, Marrow TJ, Connolley T. Time-resolved synchrotron tomographic quantification of deformation during indentation of an equiaxed semi-solid granular alloy. Acta Mater [Internet]. 2016;105:338–46. Available from: http://doi.org/10.1016/j.actamat.2015.11.028

54. Chang C-C, Lin C-J. LIBSVM: a library for support vector machines. ACM Trans Intell Syst Technol. 2011;2(3).

55. Malladi R, Sethian JA, Vemuri BC. Shape modeling with front propagation: a level set approach. IEEE Trans Pattern Anal Mach Intell. 1995;17(2):158–75.

56. van der Walt S, Schönberger JL, Nunez-Iglesias J, Boulogne F, Warner JD, Yager N, et al. scikit-image: image processing in Python. PeerJ. 2014;2:e453.

57. Batchelor BG, Waltz F. Principles of digital image processing. In: Batchelor BG, Waltz F, Interactive image processing for machine vision. London: Springer London, 1993. p. 17–45. https://doi.org/10.1007/978-1-4471-0393-6_2

58. Buades A, Coll B, Morel J. Non-local means denoising. Image Process Line. 2011;1:208–12.

59. Arganda-Carreras I, Kaynig V, Rueden C, Eliceiri KW, Schindelin J, Cardona A, et al. Trainable Weka Segmentation: a machine learning tool for microscopy pixel classification. Bioinformatics. 2017;33(15):2424–6.

60. Arganda-Carreras I, Kaynig V, Rueden C, Eliceiri KW, Schindelin J, Cardona A, et al. Trainable Weka Segmentation: a machine learning tool for microscopy pixel classification. Bioinformatics. 2017;33(15):2424–6.

61. Johnson JL, German RM. Theoretical modeling of densification during activated solid-state sintering. Metall Mater Trans A Phys Metall Mater Sci. 1996;27(2):441–50.

62. Coble RL. Sintering crystalline solids. I. intermediate and final state diffusion models. J Appl Phys. 1961;32(5):787–92.

63. Francis LF. Chapter 5—powder processes. In: Francis LF, Materials processing: a unified approach to processing of metals, ceramics and polymers. 2016.

**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of the article at the publisher’s website.