Porous Poly Ether Ether Ketone (PEEK) manufactured by a novel powder route using near-spherical salt bead porogens: Characterisation and mechanical properties

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Abstract

Porous PEEK structures with approximately 85% open porosity have been made using PEEK-OPTIMA® powder and a particulate leaching technique using porous, near-spherical, sodium chloride beads. A novel manufacturing approach is presented and compared with a traditional dry mixing method. Irrespective of the method used, the use of near-spherical beads with a fairly narrow size range results in uniform pore structures. However the integration, by tapping, of fine PEEK into a pre-existing network salt beads, followed by compaction and “sintering”, produces porous structures with excellent repeatability and homogeneity of density; more uniform pore and strut sizes; an improved and predictable level of connectivity via the formation of “windows” between the cells; faster salt removal rates and lower levels of residual salt. Although tapped samples show a compressive yield stress > 1 MPa and stiffness > 30 MPa for samples with 84% porosity, the presence of windows in the cell walls means that tapped structures show lower strengths and lower stiffnesses than equivalent structures made by mixing.

Keywords

PEEK; porosity; powder processing; characterisation; compressive strength
1. Introduction

Permanent, porous biomaterial structures have the ability to provide a transitional space between bone and a biomaterial substrate (which provides the main structural support) and an appropriate level and geometry of porosity will enable bone in-growth and hence enhanced integration between the bone and the biomaterial structure. Poly-ether-ether ketone (PEEK) has attracted wide interest as a material from which a porous medical device could be made [1]. The benefits of PEEK include; exceptional strength and stiffness for a thermoplastic polymer, excellent chemical resistance and bio-passive behaviour, X-ray translucence and excellent wear properties which match with medical implant requirements [2-8]. Despite the fact that PEEK is not bioactive, direct bone, cartilage and fibrous contacts with PEEK implants have been observed in ovine models, although the bonding is expected to be via mechanical interlocking rather than chemical [9]. Although porous PEEK structures show better implant-cortical bone contact than solid, it is recognised that enabling processes may be required for PEEK to compete directly with porous metals for chemical ossiointegration [1].

With appropriate consideration during the design of a medical device incorporating porous features, for example for an inter-body cage for spinal fusion, the porous structure need not support the inter-lumbar pressure (which is of the order of 0.5-1.0 MPa depending upon activity [10,11]). That said; the porous element must be sufficiently robust to be handled and implanted without plastic deformation or fragmentation. The strength and stiffness of porous structures follow so-called “scaling laws” [12] where the relative stiffness and strength are related to the relative density (or solid fraction) through a constant and a power law exponent. Whilst specific targets for optimum pore fractions and sizes for implant fixation via bone in-growth remain largely undefined [13], previous studies focussing on the same target application as in this work, have sought to mimic trabecular bone, with porosity close to 80% and a mean pore diameter of roughly 700 \( \mu m \) [1]. Design of an appropriate porous structure, however, necessitates a balance between achieving sufficient strength and adequate pore spaces for bone in-growth.

The production of porous structures via the removal of a pore forming or space holding (porogen) phase within a solid structure offers the greatest potential to control pore size and porosity whilst achieving a high level of pore homogeneity [14-17]. A number of researchers have used salt (NaCl) porogens in combination with several different processing methods to produce porous polymer structures: for instance, gas foaming [18]; concurrent electro-spinning [19], gas anti-solvent precipitation [20] and phase-separation [21], that tend to exhibit a wide range of irregularly-sized and shaped pores. Whilst good progress has been made, problems still persist with this method, owing to the demand for high quality and reproducible parts associated with the target application. Central to this is the difficulty in mixing or interspersion of large, irregularly-shaped salt particles, with a wide distribution in sizes, with liquids or fine powders. The homogeneity of the resulting mixture is thus difficult to control both within a part and from part-to-part.

The most promising structures, in terms of pore homogeneity and interconnectivity, have been achieved using rounded or spherical porogens [22,23], in one case using additional sintering of the packed bed of porogen to increase the connectivity between particles and hence between pores in the resultant porous structure [23]. Examples of typical porous structures resulting from these techniques are presented in Figure 1, but unfortunately there is no indication of how repeatable the
processes are, how uniform or homogenous the porosity and structures are, or quantification of the connectivity of the structure.

In both of the aforementioned studies [22,23], interspersion of the polymer into the porous preform was affected by infiltration with a polymer solution. Unfortunately the limited solubility of PEEK in suitable solvents means that porous PEEK scaffolds cannot be made in the same way and an alternative approach is required. Although porous PEEK samples have been made by a number of different methods (reviewed in [1]), particulate leaching methods are commonly preferred. Irrespective of the process used, achieving structures that mimic trabecular bone, with an appropriate level of pore interconnectivity and adequate mechanical strength has been difficult [1] and is additionally problematic from a processing perspective given the high melting point of PEEK (345°C [24]) and the lack of suitable solvents. This study aims to demonstrate a process that can greatly enhance the quality and reproducibility of interspersion of PEEK powder into a bed of NaCl beads and hence of the density and interconnectivity of the porous product. Some preliminary mechanical tests have also been undertaken to determine their suitability as implantable structures.

![Figure 1: SEM micrographs of (a) PLGA scaffolds made using 355–450 μm diameter paraffin spheres [22] and (b) PLLA-reinforced apatite scaffolds manufactured using 300–500 μm rounded salt particles [23]](image)

2. **Experimental procedure**

Sodium chloride (NaCl or salt) powder with 99.5% purity was processed into near-spherical beads (with an aspect ratio of 1.14±0.02) using a method outlined in [25, 26]. Beads with diameters between 1.0 and 1.4 mm were selected by sieving. PEEK-OPTIMA® LT1 PEEK powder was used and was supplied by Invibio Ltd. The morphologies of the beads and the PEEK powder are shown in Figure 2. The PEEK particle size distribution was measured using a Malvern Mastersizer 2000 instrument, yielding an average diameter ($D_{50}$) of 54 μm, with a range ($D_{10}$ and $D_{90}$) of 21 and 110 μm respectively. The densities for the two components were $1.3 \text{ g cc}^{-1}$ for the PEEK and $1.65 \pm 0.05 \text{ g cc}^{-1}$ for the (23% porous) NaCl beads (as measured in [26]). The packing behaviour for the powders was studied using a Quantachrome Autotap machine, by tapping a known mass of powder in a
graduated measuring cylinder. The apparent and tap densities for the powders were 0.90 and 0.97 g cc\(^{-1}\) for the salt beads (packing fractions of 0.55 and 0.59 respectively) and 0.37 and 0.42 g cc\(^{-1}\) for the PEEK (packing fractions of 0.28 and 0.32 respectively).

![Figure 2 (a) Near-spherical salt beads with diameters between 1.0 – 1.4 mm and (b) PEEK powder](image)

In order to achieve porosity in excess of 80%, consideration of the powder densities led to a 1:6 mass ratio of PEEK to salt beads being used (leading to an estimated 83% porosity in the PEEK scaffold after salt removal). Two distinct processing methods were used, both involving a “mixing” process followed by compaction, sintering (or melting), salt dissolution and drying. The “mixing” methods differed, namely dry mixing of the PEEK and salt or tapping of the PEEK through a bed of salt beads (a schematic of the process steps is shown in Figure 3).
A dry mixing process was used to intersperse the salt beads and the PEEK powder. 42 g of salt beads were mixed with 7 g of PEEK powder in a Turbula mixer for 15 min. 4.9 g of the mixed powders was then poured into a die for subsequent compaction. Difficulties in achieving a homogeneous mixture of the dry powders were evident, as was their capacity to segregate during handling and transfer into the die. Every effort was made to minimise segregation through careful mixing, handling and transfer methods.

PEEK and NaCl beads were also combined using a tapping-based process previously applied to metal powders [25]. In this process, 4.2 g of salt beads were poured into the cavity of a 22 mm diameter die (with the bottom punch in place) and packing was enhanced by brief tapping using the same Autotap machine as described earlier. 0.7 g of PEEK powder was then placed on top of the bed, where after the top punch was inserted. The whole assembly was then tapped, using the same machine, which employs a simple lifting and dropping action, stopping after a pre-determined number of cycles (approximately 6000). An image of the tooling and a schematic of the layered structure and the tapping direction (marked with an arrow) are given in Figure 4.

After both interspersion methods, uniaxial die compaction was performed at a pressure of roughly 200 MPa to produce a robust PEEK-NaCl “composite” pellet. The pelleted samples were sintered at 380°C for 30 min and, when cool, were immersed in warm (40-50°C) water to dissolve the salt for a minimum of 4 hr and subsequently dried at a maximum of 90°C for a minimum of 3 hr.
Figure 4: (a) Tapping tooling and (b) schematic of the powder layers (the arrow shows the tapping direction, A, B and C are the lower and upper punch and die respectively)

Scanning electron microscopy (SEM) was used to analyse the PEEK scaffold morphology. Sectioned PEEK scaffolds were sputter coated with platinum prior to examination. The pore morphology was quantified using X-ray micro-computed tomography (CT) on a Scanco Medical µCT 40 instrument. Compressive testing of the foams was performed using an Instron Universal Testing machine, compressing samples 22 mm in diameter and 5 mm in height. Samples were deformed at a rate of 0.5 mm min\(^{-1}\), testing a minimum of 6 samples per type. In order to measure the stiffness, twin extensometers were used to accurately measure the compressive strain during multiple unloading and reloading cycles after initial loading to 75% of the compressive yield point.

3. Results and discussion

3.1 General Observations

For the tapping process, loading the beads and tapping creates a packed bed with height of roughly 11.3 mm (the same as that predicted from the powder mass and the tapped packing fraction). An additional approximately 5 mm thick layer of PEEK was then applied on top. After the tapping was completed (defined by when further tapping produces no further detectable migration of the PEEK – manifested by no change in the position of the top punch in the assembly) the final height of the combined powder bed was roughly 11.8 mm, indicating that nearly all the PEEK had been tapped into the void space in the bead structure and that an above-optimum mass ratio (of PEEK to NaCl) was used. It was calculated that the PEEK filled the void space created by the packed beads with a packing fraction of 0.29, lower than the 0.32 for tapped PEEK in an “infinite” vessel, no doubt affected by the scale and morphology of the pores relative to the size of the PEEK powder.

During compaction the void spaces between the PEEK particles and between the NaCl beads are removed, creating, for both cases, a compact with a density of 1.85 g cc\(^{-1}\), albeit in the case of the tapped sample with a PEEK layer on the surface. For the simpler case of the dry mixed sample,
without a PEEK layer, the fact that the compact density exceeds the theoretical density for the mixture (1.59 g cc\(^{-1}\)), indicates some densification of the porous salt beads (to an estimated 1.97 g cc\(^{-1}\) and 8% porosity). It has been observed that for beads alone, compacted at the same pressure, a density of 2.02 g cc\(^{-1}\) is achieved; indicating that bead densification is possible.

The bottom surfaces of compacted pellets made by both mixing methods are shown in Figure 5, from which it is clear that PEEK powder has migrated through the packed bed of NaCl beads. It is also apparent from these images that the distribution of salt beads is much more uniform for the tapped structure, with bead-rich and bead-depleted regions being much more prominent for compacts made by dry mixing and compaction. In both cases there was no evidence for cracking or fragmentation of the beads as a result of the compaction process.

Despite “sintering” of the samples above the melting point for PEEK, there was little to no distortion of the pellets, with only a very slight (on average <0.5%) decrease in diameter. Although the PEEK melts, it is likely to be held in the gaps between the salt beads by capillary forces. The high fraction of NaCl beads lends rigidity to the compact, without them, significant slumping is observed. Measurements of the time required for removal of the salt from the sintered pellets, revealed faster removal from tapped samples as well as lower levels of residual salt, suggesting improved interconnectivity for tapped structures.

3.2 Porous structures

It was found that both methods were able to produce a porous structure that was sufficiently robust to be handled. Photograph and CT images of typical porous pellets made by the methods described are given in Figure 6 (in the case of the sample made by tapping the thin PEEK layer on the surface was removed by manual grinding so as not to affect imaging and measurement of the porosity and other structural features). It is clear that in both cases the use of a narrow size range of spherical
beads, in conjunction with the simple processing route, gives rise to structures with far more uniform pore sizes and structures compared with those presented in Figure 1 for alternative leaching-based processes. For the structures made by tapping, it is also apparent that there are multiple small windows linking the pores.

The mean porosity for a batch of 10 samples made by dry mixing was 79%, with a standard deviation of 1.3%. For tapping, the mean porosity was 84% with a standard deviation of 0.5%. The lower standard deviation for the tapped structures demonstrates the capability for improved reproducibility in porosity for this process. The targeted porosity, based on the 1/6 weight ratio was 83%, the discrepancy between actual and target densities in the case of the dry mixed samples is due to a decrease in bead volume owing to their densification during compaction. In the case of the tapped samples, although the same bead compaction behaviour is relevant, the higher porosity is a result of incomplete interspersion of PEEK and the NaCl (manifested by the PEEK surface layer).

Figure 6: Porous PEEK samples manufactured by dry mixing (a, b) and tapping (c, d)
Figure 7 presents the porosity distribution for typical porous samples taken from area fraction data from circa 150 2D sections obtained by CT. To enable close comparison, the porosity is normalised with respect to the mean value. The improved homogeneity of porosity within the structure made by tapping is clear. The variation in porosity with height is very uniform for the tapped sample, with a standard deviation of only 1.2%. For the sample made by dry mixing, the porosity level is much more variable, with a standard deviation of 7.0%, and shows a clear tendency for a PEEK-rich layer to form at the top surface and for salt bead segregation to the base, a reflection of the difficulty in achieving homogeneous mixing of the two phases, which have a large size ratio (22:1) and the difficulty in delivering a homogeneous mixture of powders to the die.

Figure 7: The variation in normalised porosity with sample height

Figure 8 shows, more clearly, that samples produced by the tapping process exhibit windows linking adjacent pores and that this connectivity, in terms of the size and number of windows, is similar from pore-to-pore, throughout the structure. The same connectivity is not apparent in the samples made by dry mixing but, nonetheless, must be present in some form since close to all the salt is removed from the sample during the dissolution step. The “window” connections between pores are formed due to the inability of the PEEK powder (even after melting) to penetrate the regions between and in the vicinity of contacting salt beads. The number of connections is dictated by the coordination number for packing of the beads, which is between 5 and 6 for loose and dense random packing [27-28]. Typically 2 to 3 such windows can be seen in sectioned pores. The mean diameter of these windows, which is difficult to measure when observed away from normal to the viewing axis, is estimated to be between 250-350 μm, but with a large standard deviation. Although
the porous structures made by tapping clearly have large “primary” pores, broadly defined by the
glumetry of the salt beads, the windows which control the interconnectivity of this structure are
within the broad range expected for enabling cell growth and migration [29] and resultant bone in-
growth [13].

A simple geometrical model can be used to estimate the “window size” between contacting spheres
based on the closest proximity that a smaller sphere can pack between two larger ones. As scatter
in the range of possible hole sizes will result from the range in bead sizes and the range in PEEK
powder size. Taking the median for both, a window diameter of 314 µm is predicted (although the
scatter for the full range of possibilities is 190 – 456 µm).

![Figure 8: Optical micrographs of the top surface of porous PEEK structures fabricated by (a) dry
mixing and (b) tapping.](image)

Figure 9 compares the microstructures for the porous PEEK made by the two routes. It is evident in
both cases that the pores have been flattened by the compaction process to become elongated
perpendicular to the compaction direction. At this higher level of magnification, connections
between the pores in the structure made by dry mixing can be observed, some are very small, and
others very large, a result of the inhomogeneous mixing behaviour for the two components and the
potential for PEEK powder to surround the salt beads. In both cases, fusion of the PEEK particles has
occurred resulting in dense struts such that the porosity measured will predominantly be in the form
of large pores, controlled by the bead size, rather than small open or closed pores within the struts
and cell walls. The pore structures (their uniformity and connectivity) resemble those shown in
Figure 1a, that were also made using spherical porogens, but it should be noted that the struts in
porous structures made by tapping appear to be significantly more dense than those in Figure 1b, in
particular, produced by drying of a polymer solution.

The pore size distributions for the porous PEEK, measured using CT, are presented in Figure 10.
These measurements, which are also summarised in Table 1 along with other structural parameters
measured by CT, clearly show that the mean (and modal) pore sizes for the two processes are
similar, which is to be expected given the same beads sizes were used. There does, however, appear
to be a larger fraction of smaller pores for the structures made by dry mixing, which is reflected in
the higher standard deviation in the mean value. It is likely that small unfilled regions may exist
between clustered NaCl beads produced by dry mixing that may not be penetrated by the molten
PEEK, skewing the pore size distribution to lower values. The mean and maximum strut thickness, and standard deviation in the mean, indicates that a higher level of structural inhomogeneity exists for samples made by dry mixing, owing to the greater variability in the inter-bead spacing for a randomly mixed powder system.

It should be noted, however, that these pore size measurements do not correspond closely with the original bead sizes. The pore size is determined by the CT instrument by filling the pores with the largest sphere possible, which for the compacted beads form pores with a prolate spheroid shape, will approximate to the minor axis (parallel to the axis of compaction). The degree of anisotropy, measured by CT is close to 1.5 for both processes. The origin of the elongation is of course the compaction process which is similar for both processes. It should also be noted that the porous salt spheres are compressible and are able to reduce in volume (by an estimated 15%) during the compaction process. Inspection of the lower magnification images in Figure 9 support a reduction in the bead size during processing.

Figure 9: SEM images of vertical cross sections of porous PEEK structures fabricated by (a,c) dry mixing and (b,d) tapping. The arrows in the image show the compaction direction for all cases.

The connectivity density, which represents the ratio of the connecting holes, vertices (end points), and struts (bridges) in one unit volume, is also presented, which for dry and tapping processes are close to 9 and 6 per mm³, respectively. Thomsen et al [30] stated that human bone has a
connectivity density less than 12 per mm$^3$, and Kabel et al [31] found that in samples of humans between 14 and 91 years old, the connectivity density varied between 1 and 9 with a mean close to 3. The connectivity density measurements, in particular for the tapped structure, are in keeping with those expected for a material designed to resemble the structure of trabecular bone and suggested targets of 7 per mm$^3$ [1].

![Figure 10: Pore size distribution comparison for dry and tapped structures](image)

Table 1: Structural parameters for porous PEEK measured by CT

<table>
<thead>
<tr>
<th>Manufacturing method</th>
<th>DRY</th>
<th>TAPPED</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mode of pore size (mm)</td>
<td>0.63</td>
<td>0.69</td>
</tr>
<tr>
<td>Mean pore size (mm)</td>
<td>0.54</td>
<td>0.58</td>
</tr>
<tr>
<td>Standard deviation of the mean pore size (mm)</td>
<td>0.20</td>
<td>0.16</td>
</tr>
<tr>
<td>Mean strut thickness (mm)</td>
<td>0.24</td>
<td>0.17</td>
</tr>
<tr>
<td>Standard deviation of the mean strut thickness (mm)</td>
<td>0.16</td>
<td>0.07</td>
</tr>
<tr>
<td>Maximum strut thickness (mm)</td>
<td>1.17</td>
<td>0.63</td>
</tr>
<tr>
<td>Degree of anisotropy</td>
<td>1.52</td>
<td>1.50</td>
</tr>
<tr>
<td>Connectivity density (1/mm$^3$)</td>
<td>9.22</td>
<td>6.43</td>
</tr>
</tbody>
</table>

3.3 Benefits of the tapping process

The tapping process creates a stable and fairly densely packed NaCl bead structure before the PEEK is introduced and then facilitates the migration of the (relatively) fine PEEK powder through the network of channels between the beads. Using spherical beads with a reasonably tight size range results in a homogenous network that is unlikely to have inaccessible areas of clustered porogen (as is likely to be the case for angular particles with a wide size distribution). The tapping process also pre-establishes the inter-particle contacts which are well-defined due to the repeatable and predictable loose or dense random packing characteristics for near-monosized spherical particles.
The level of interconnectivity achieved by tapping is sufficient, as measured by the connectivity density, to not require an additional sintering step.

This is in stark contrast to the highly variable quality of mixing of powders with a large size difference and the segregation problems associated with flow and transfer of these mixtures. Although the use of binders or compounding can create improved distributions of the two powders and may overcome segregation of the fine powder, it will prevent direct bead-bead contact and hence greatly reduce the interconnectivity. The benefit of pre-establishing the network of NaCl beads is borne out in improved repeatability and homogeneity of density and pore size, an improved and predictable level of connectivity, faster NaCl removal rates and lower levels of residual salt. Figure 11 shows 3D X-ray tomographic reconstructions of the full sample thickness, highlighting the residual salt (bright). Not only is the salt level much lower in the sample made by tapping (estimated from image analysis to be < 0.1% of the pore volume) but the high levels of salt in the mixed samples (estimated to be > 12% of the pore volume) remain there after more than 72 hr of immersion (compared with 4 hr for the tapped sample).

It is clear that the tapping route, whilst less convenient, produces structures with the not only the characteristic structures, but also with the quality and reproducibility demanded. For the bead size used (1.0 to 1.4 mm diameter), consolidation and elongation of the beads has produced porosity that is close to, but slightly larger than the target size (700 μm) achieving a good compromise in terms of the size of the windows between the pores, strut thickness and connectivity density. If required, pore sizes and windows could be further tailored by altering the size and size distribution for the beads used, with a requisite reduction in the size of the “filler” powder as the salt bead size decreases. The ability to vary the pore fraction is more limited. Since the process “fills” the available space within the packed bead structure (a reduced number of tapping cycles only part-fills the structure and excess PEEK simply remains on the top surface), porosity is dictated by the packing behaviour of the beads and of the PEEK. If the porosity level needed to be increased (for the same bead size and type) an increase in the PEEK powder size would affect this through less efficient filling of the void spaces.

Figure 11: 3D views through the porous structures highlighting the residual salt (bright) in structures made by (a) tapping and (b) dry mixing.
3.4 Mechanical testing

Figure 12 presents stress – strain curves for a dry mixed sample, with a porosity of 79.7% (relative density, RD, of 0.20) and a tapped sample with a porosity of 84.5% (RD = 0.16). The stress-strain curves are typical of those for the samples tested, showing an initial peak, taken as the compressive strength, followed by a plateau region which is flatter for samples with lower densities. Densification of the porous structures commences at strains of approximately 60%. Table 2 summarises the mechanical properties for porous structures made by the two methods, expressing them as averages over the batch of samples tested. The samples produced by dry mixing, with the higher density, show higher compressive yield strengths and significantly higher stiffnesses. Whilst samples made by both methods demonstrate average compressive yield stresses close to or slightly above those for typical lumbar loads, the strength and stiffness are well below those for trabecular bone (36 MPa and 1-10 GPa respectively [1,13]). As a consequence, it is expected that these structures will be designed to be non-load bearing elements within hybrid structures or complex devices.

<table>
<thead>
<tr>
<th>Sample</th>
<th>Porosity / %</th>
<th>RD</th>
<th>E / MPa</th>
<th>Yield Strength / MPa</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dry mix</td>
<td>79.2 ± 1.3</td>
<td>0.21</td>
<td>65.6 ± 8.9</td>
<td>1.85 ± 0.3</td>
</tr>
<tr>
<td>Tapped</td>
<td>83.9 ± 0.5</td>
<td>0.16</td>
<td>39.2 ± 4.0</td>
<td>1.22 ± 0.2</td>
</tr>
</tbody>
</table>

Figure 12: Typical stress-stain curves for mixed (RD = 0.21) and tapped (RD = 0.16) structures

Within the samples tested for each manufacturing method there is variability in the density. Although the range of densities explored is rather limited and extrapolation should be taken with some caution, plots of normalised strength or stiffness against relative density can be produced and, through fitting
to simple models, a comparison between equivalent samples (those with the same relative density) made by the two processing routes can be made. Scaling laws for stiffness and yield strength, with respect to the relative density and properties of the solid, $s$, are given in equations 1 and 2, for open cell foams [12], where only the strengthening or stiffening due to deformation of the cell edges is considered.

$$\frac{E}{E_s} = C_1 \left(\frac{\rho}{\rho_s}\right)^2$$

Eq 1

$$\frac{\sigma}{\sigma_s} = C_2 \left(\frac{\rho}{\rho_s}\right)^{3/2}$$

Eq 2

Figures 13 and 14 plot the stiffness and strength data as a function of the relative density. Fit lines to a power law of the type shown in equations 1 and 2 are also shown, as are the fit values. The power exponents are close to the model values for the samples made by dry mixing (1.9 and 1.6 for stiffness and strength respectively), but is significantly higher in both cases for the tapped structures (3.1 and 4.2 for stiffness and strength respectively), indicating a stronger reduction in the stiffness and strength with increasing porosity. This difference is thought to be due to structural differences, principally as a result of the large windows in the cell walls that are present in the tapped structures (although it should be noted that residual salt in the mixed samples may contribute to the strength and stiffness). The fit to the open cell model is better for samples made by tapping and for the stiffness data, supporting the increased contribution of the cell walls to the strength for samples made by mixing (with no large windows in the cell walls) and the more harmful effect of cell wall porosity on stiffness [32].

Although the study presents a fair measurement of the compressive properties for samples that are typical of the scale required in a medical device and a fair comparison between the two manufacturing processes, caution should be taken to extrapolate these trends beyond the data reported here, given not only the small interval of densities studied, but additionally that the sample thickness is below the minimum recommended to show independence of properties with thickness (typically 8 times the cell diameter [33]).
4 Conclusions

Porous PEEK structures were produced by two methods, both using near-spherical salt beads. The use of near-spherical beads with a fairly narrow size range results in uniform pore structures.

The integration, by tapping, of fine PEEK into a pre-existing network of 1.0-1.4 mm diameter salt beads, followed by compaction and “sintering”, produces porous structures typified by slightly
elliptical pores, elongated normal to the compaction direction, with 5 to 6, approximately 300 μm diameter windows connecting each pore with its neighbours.

Pre-establishing the contacts between the salt beads, before addition of the PEEK powder, results in: structures with improved repeatability and homogeneity of density, more uniform pore and strut sizes, an improved and predictable level of connectivity, faster NaCl removal rates and lower levels of residual salt.

Porous structures produced by tapping and with 84% porosity, show a compressive yield stress > 1 MPa and stiffness > 30 MPa. The presence of windows in the cell walls, however, means that for samples with the same relative density, tapped structures will show lower strengths and stiffnesses compared with those made by dry mixing.

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References


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