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Nonlinear oscillatory dynamics of the hardening of calcium phosphate bone cements

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Here we report on the nonlinear, oscillatory dynamics detected in the evolution of phase composition during the setting of different calcium phosphate cements, two of which evolved toward brushite and one toward hydroxyapatite as the final product. Whereas both brushite-forming cements contained iondoped β -tricalcium phosphate as the initial phase, the zinc-containing one yielded scholzite as an additional phase during setting and the oscillations between these two products were pronounced throughout the entire 80 h setting period, long after the hardening processes was over from the mechanical standpoint. Oscillations in the copper-containing system involved the amount of brushite as the main product of the hardening reaction and they progressed faster toward an equilibrium point than in the zinc-containing system. Initially detected with the use of in situ energy-dispersive X-ray diffractometry, the oscillations were confirmed with a sufficient level of temporal matching in an in situ Fourier transform infrared spectroscopic analysis. The kinetic reaction analysis based on the Johnson-Mehl-Avrami-Kolmogorov model indicated an edge-controlled nucleation mechanism for brushite. The hydroxyapatite-forming cement comprised gelatin as an additional phase with a role of slowing down diffusion and allowing the detection of otherwise rapid oscillations in crystallinity and in the amount of the apatitic phase on the timescale of minutes. A number of possible causes for these dynamic instabilities were discussed. The classical chemical oscillatory model should not apply to these systems unless in combination with less exotic mechanisms of physicochemical nature. One possibility is that the variations in viscosity, directly affecting diffusion and nucleation rates and accompanying growth and transformation from the lower to the higher interfacial energy per the Ostwald-Lussac rule, are responsible for the oscillatory dynamics. The conception of bone replacement materials and tissue engineering constructs capable of engaging in the dynamics of integration with the natural tissues in compliance with this oscillatory nature may open a new avenue for the future of this type of medical devices. To succeed in this goal, the mechanism of these and similar instabilities must be better understood.

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1. Introduction

Life is pervaded with nonlinear, far-from-equilibrium dynamics to such an extent that linear phenomena present a comparative rarity.¹⁻³ To promote solvability of physical models and predictability of physical phenomena, naturally nonlinear processes are linearized using specific mathematical and phenomenological approximations, such as the invocation of small and harmonic oscillations, small amplitude waves propagating through homogenous environments, infinitesimal temperature changes, depiction of atoms as ideal, noncompressible spheres, and so on. This habit to linearize the nonlinear is, in fact, so pervasive that the models are frequently confused with the reality and it has become customary to perceive a plethora of inherently nonlinear processes as ostensibly linear.⁴ It is being neglected that the linearity of quantum mechanics and relativity theory are thanks to the assumption of dimensional infinity and the neglect of the infinite number of terms of the energy-mass relation series, respectively. In contrast to the nonlinear physical reality, synthetic materials and devices have been traditionally built around the concept of equilibrium dynamics. For example, the linear, zero-order kinetics of drug release is still most sought in controlled drug delivery, while the oscillatory response is a nightmare of most sensory device developers. Yet, without embodying nonlinear responses, the memory effect, the hierarchical superstructures, emergence properties and other advanced features cannot be created in materials. From today's perspective, it is doubtful if the methods based on thermodynamic equilibrium can yield some of the most sophisticated features that endow life, from creative intelligence to self-regeneration to autopoiesis. Moreover, to advance the concepts of biomimetics and biomimicry

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forward, it may be necessary to conceive materials and processes that integrate the principles of nonlinearity therein.

Oscillatory behavior under steady state has been traditionally associated with complex chemical reactions in solution,⁵⁻⁷ but is commonly detected at the quantum scale too.8-10 More seldom it has been observed involving gas catalysis on solid surfaces.¹¹⁻¹³ Its theoretical descriptions have been relatively rudimentary and there is a consensus that the understanding of the underlying mechanisms is generally poor.¹⁴ Oscillatory behavior involving material transport phenomena and phase transitions at the solid/liquid interface is detected even more rarely. Traditionally, chemical reactions are depicted as proceeding linearly from the reactants to the products until a state of equilibrium is reached. This pathway, however, broken down to infinitesimally small steps, is composed of alternations in the reaction direction - from the reactants to the products and from the products to the reactants - that, taken as a whole, still yield to the global direction of the reaction path. Such oscillations often proceed unnoticed, but in the presence of specific inhibitors in networked, multicomponent systems with a sufficient number of intermediaries and significantly slowed diffusion, they can be magnified and detected at timescales on the order of seconds to minutes to even hours, depending on the nature and the conditions of the reaction. Here we report on the detection of such oscillations in the setting reaction of three different self-setting calcium phosphate (CP) cements.

Developed in late 1980s and approved by the FDA in 1996,15 self-setting CP cements remain state-of-the-art inorganic bone fillers. They are typified by excellent osteoconductivity; perfect fit to the bone defect geometry; intimate bone interface and union; and good flow properties and injectability for a pure inorganic material. Given that their viscosity depends on the shear rate and deformation history, self-setting CP cements are non-Newtonian fluids characterized by a nonlinear rheology. Their main downside comes from setting in relatively dense forms in the absence of additives acting as porogens.16,17 These nonporous forms are nowadays being challenged by tissue engineering constructs with precisely designed porosities that allow for cell infiltration and superior integration with the adjacent bone.18 The setting reaction in CP cements follows one out of two possible paths, one ending with brushite as the final product and the other one ending with hydroxyapatite (HAp).¹⁹ In this study, we tested variations to both of these cement types.

2. Experimental

Zn²⁺- and Cu²⁺-substituted β-tricalcium phosphate (β-TCP, whitlockite) powders were obtained by admixing Cu(NO₃)₂ or Zn(NO₃)₂ aqueous solutions conforming to Ca_{2.917}Cu/ Zn_{0.083}(PO₄)₂ stoichiometry to an aqueous solution of Ca(NO₃)₂ and then adding (NH₄)₂HPO₄, while maintaining the solution at neutral pH conditions during precipitation. Specifically, to prepare Zn/Cu-TCP, 875 ml of 0.5 M Ca(NO₃)₂ and 125 ml of 0.1 M Cu(NO₃)₂ or Zn(NO₃)₂ × 6H₂O were mixed and 667 ml of 0.5 M (NH₄)₂HPO₄ was added dropwise over the course of 10 min. The pH of the solution was kept at around 7 throughout the reaction by adding concentrated ammonia solution. This was done to minimize the alkalinity, but ensure the precipitation of apatite and not brushite, thus minimizing the chances for the formation of pyrophosphate as the secondary phase during calcination.²⁰ After 30 min of aging, the precipitate was filtered, washed with distilled water, dried at 80 °C and then calcined at 900 °C to remove the traces of NH₄NO₃ and to form the partially Zn²⁺/Cu²⁺-substituted whitlockite structure.

Zn²⁺- and Cu²⁺-substituted β -TCP cements were obtained by mixing the given β -TCP powders with monocalcium phosphate monohydrate (Ca(H₂PO₄)₂ × H₂O, MCPM) and carbonated HAp at the weight ratio of 1 : 0.764 : 0.091, and then adding a 0.45 M aqueous solution of citric acid. The powder-to-liquid ratio was 4.08 g ml⁻¹. The main product of both setting reactions was dicalcium phosphate dihydrate (DCPD, brushite) per the following equation:

$$Ca_{3}(PO_{4})_{2} + Ca(H_{2}PO_{4})_{2} \cdot H_{2}O + 7H_{2}O \rightarrow 4CaHPO_{4} \cdot 2H_{2}O (1)$$

Gelatin-containing HAp cement was fabricated by mixing 25 ml of 5 wt% porcine-skin-derived type A gelatin (Sigma) containing 500 µl of tetraethylorthosilicate (Sigma) with a HAp suspension prepared by adding 3 ml of 1 M NH₄H₂PO₄ (aq.) (Fisher Scientific) dropwise to 2.5 ml of 2 M $Ca(NO_3)_2$ (aq.) (Fisher Scientific) and then adding 9 ml of concentrated, 28% NH4OH (Sigma Aldrich) and 85 ml of deionized, Millipore water and bringing to boil. Right after the boiling point was reached, the two solutions were mixed and stirred at 400 rpm. The colloid was sedimented by centrifugation and washed with deionized water. The resulting precipitate comprising poorly crystalline HAp embedded in a gelatinous matrix was dried overnight in vacuum (-25 mmHg, Savant SPD1010 Speedvac concentrator). The dried powders crushed in a mortar down to fine particle size were mixed with 2 wt% NaH₂PO₄ (Fisher Scientific) as the liquid phase in 0.33 mg μ l⁻¹ powder-to-liquid ratio to yield the cements. Vortex-mixing was performed in almost full Eppendorf tubes to minimize the exposure to air and the dissipation of the paste on the tube walls.

The control, amorphous CP powder was prepared by abruptly adding a solution containing 100 ml 0.5 M Ca(NO₃)₂ and 7 ml 28% NH₄OH into a solution comprising 100 ml 0.2 M NH₄H₂PO₄ and 4 ml 28% NH₄OH. The fine precipitate formed upon mixing was aged for 15 s before it was collected, centrifuged, washed with 0.14 w/v% NH₄OH, centrifuged again, and dried overnight at low pressure (p = -20 mmHg) and room temperature. The pH of the supernatant following the precipitation reaction was 9.3. The powder was turned into a cement by mixing it with a 2 wt% aqueous solution of Na₂HPO₄ at the 1.782 g ml⁻¹ solid-to-liquid ratio using a digital vortex mixer (Fisher Scientific) for 5 min at 2000 rpm.

X-ray diffractograms (XRDs) were recorded *in situ*, at different time points of the setting reaction, either on a Rigaku D/Max-2500 or a Bruker D2 Phaser diffractometer. Time-resolved XRD characterization of the setting and hardening of Zn- and Cu-TCP cements was carried out by means of a home-made energy-dispersive XRD (EDXRD) instrumentation²¹⁻²³ equipped with a W X-ray tube operating at 50 kV, 30 mA, 200 µm

wide apertures and $\theta^{\circ} = 5^{\circ}$. After manual mixing with a putty knife, a small quantity of the cement was placed on a low scattering glass sample holder and quickly transferred to the optical center of the diffractometer. The semi-quantitative method based on relative intensity ratios (RIR) compared to the corundum standard²⁴ was applied to estimate the weight ratio between crystalline phases at each time point. The following equation was used to calculate the weight content of phase α (W_{α}) in a mixture of *n* phases, with RIR values taken as 1.10 for hydroxyapatite (HAp), 1.42 for brushite (DCPD), 0.71 for monetite (DCPA), 0.45 for α -TCP and 1.20 for β -TCP (PDF, ICDD) for Cu K α irradiation source, and I_{α} being the intensity of the most intense reflection of the phase α :

$$W_{\alpha} = \left(\frac{I_{\alpha}}{\mathrm{RIR}_{\alpha}}\right) / \sum_{\alpha=0}^{n} I_{\alpha} / \mathrm{RIR}_{\alpha}$$
(2)

The degree of crystallinity, X_c , in HAp was calculated in relation to the absolute intensity of the (300) reflection, I_{300} , and the absolute intensity of the hollow between (112) and (300)

reflections, *V*₁₁₂₋₃₀₀, which disappears in non-crystalline samples.

$$X_{\rm c} \approx 1 - \left(\frac{V_{112-300}}{I_{300}}\right)$$
 (3)

XRDs converted to compositional values were analyzed using the Johnson–Mehl–Avrami–Kolmogorov (JMAK) nucleationgrowth kinetic model.²⁵ The following relationship was used to derive the Avrami rate constant, k, for crystallization of DCPD and the Avrami exponent, n, related to nucleation mechanism, where the extent of the reaction, α , was determined as the ratio of a selected diffraction peak at time t and at its maximal value during the reaction:

$$\ln[-\ln(1-\alpha)] = n\ln(k) + n\ln(t) \tag{4}$$

Absolute intensities of the diffraction peaks were measured as their maximal height, while integrated intensities were measured by calculating a background function and deconvoluted peak profile using an automated Gaussian-fitting routine (OriginPro 2016). Correlations between the diffraction



Fig. 1 Oscillations of the absolute intensities of the most intense reflections of the three different crystalline phases detected during the setting of the Zn-TCP cement. CaHPO₄ is brushite (DCPD), Ca₃(PO₄)₂ is whitlockite (β -TCP), and CaZn₂(PO₄)₂ is scholzite (ZCP). Green rectangles denote regions in which oscillations between the two phases – *i.e.*, a decrease in the intensity of one phase paralleled by an increase in the intensity of the other phase – were detected.

intensities and compositional variables are semi-quantitative and the pronounced scale and repeatability of the appearance of detected oscillations, significantly minimized in control solid samples and in cements at later stages of the setting reaction, presents the basis for arguments that follow. Fourier transform infrared spectroscopy (FT-IR) was used to additionally follow the setting reaction over time, and the analysis was done on a Bruker Alpha Platinum ATR spectrometer.

Degrees of saturation (DS) for different CP phases were calculated using an algorithm based on Debye–Hückel equation:

$$DS = pK_{sp} - pQ$$
(5)

$$Q = \{ Ca^{2+} \}^{x} \{ PO_{4}^{3-} \}^{y} \{ H^{+} \}^{z} \{ OH^{-} \}^{w}$$
(6)

Q is the ionic activity product of the solution, and $pK_{\rm sp}$ is the negative logarithm of the solubility product of a given CP phase, equaling 117.3 for HAp, 6.6 for DCPD, 29.5 for β -TCP, and 48.4 for octacalcium phosphate (OCP). Activity coefficients were calculated through log $\gamma = -Az_i^2 I^{1/2}$, where z_i is the charge number of ion species i, *I* is the ionic strength of the solution,

and *A* is the constant depending on the dielectric constant of the solution, temperature and Debye screening length.

3. Results

3.1. Zn-TCP cement

Fig. 1–3 display the diffraction intensity and compositional oscillations between the three different crystalline phases detected during the setting of the Zn-TCP cement: brushite (DCPD, CaHPO₄ × 2H₂O), partially zinc-substituted β-tricalcium phosphate (β-TCP, Ca₃(PO₄)₂/Ca_{3-x}Zn_x(PO₄)₂) and zinc calcium phosphate (ZCP, CaZn₂(PO₄)₂ × 2H₂O) a.k.a. scholzite, a dihydrous compound crystallizing in the dipyramidal orthorhombic *Pbmm/Pbc2*₁ space group symmetry (*mm*² point group). Oscillations between two phases are noted within any given time range whenever the presence of one phase increased on the expense of the diminished presence of another. The most pervasive oscillations of this type were observed between the two major final phases of the setting reaction, absent in the initial composition: DCPD and ZCP. These oscillations were present in 75% of the first 80 h of the reaction for the diffraction



Fig. 2 Oscillations of the weight ratios of three different crystalline phases detected during the setting of the Zn-TCP cement. CaHPO₄ is brushite (DCPD), $Ca_3(PO_4)_2$ is whitlockite (β -TCP), and $CaZn_2(PO_4)_2$ is scholzite (ZCP). Green rectangles denote regions in which oscillations between the two phases – *i.e.*, a decrease in the weight percentage of one phase paralleled by an increase in the weight percentage of the other phase – were detected.

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intensity (Fig. 1c) and 88% for the calculated composition, including the entire time span from 35 to 80 h (Fig. 2c). Oscillations involving either of the two products, DCPD or ZCP, and the initial phase of the reaction, partially Zn-substituted β -TCP, excluded each other, as can be seen by comparing Fig. 1b and d. Namely, in whatever the time range the oscillations between β -TCP and DCPD were present, they were absent between β -TCP and ZCP. Added to one another, however, they occur in 72% of the total reaction time. This suggests that the two reaction products, DCPD and ZCP, form in separate reaction paths and are connected by the initial and central component, Znsubstituted β -TCP. Per eqn (7), any increase in the amount of DCPD forming from β -TCP would rebalance the reaction per Le Chatelier's principle in such a manner that some of the ZCP, lying on the opposite end of the reaction, would transform back to β -TCP. Similar depletion of DCPD in the product would entail the formation of ZCP per eqn (7). This explains how the oscillations between β -TCP and ZCP exclude those between β -TCP and DCPD, while oscillations between DCPD and ZCP dominate the reaction.

$$DCPD \rightleftharpoons \beta\text{-}TCP \rightleftharpoons ZCP$$
 (7)

The fact that the initial phase, β -TCP, remains in the system (Fig. 1a) and steadies at around 30 wt% of the product mass (Fig. 2a) facilitates the extended duration of oscillations, which would have been shorter had the phase transformation been more complete. As a result, oscillations between the two phases of the product, DCPD and ZCP, were sustained throughout the entire 80 h period following the onset of the setting reaction.

Between each of the 19 data points in the 35–80 h time range, the trend was the same: one of these two phases increased in abundance at the expense of another (Fig. 3c). Although the cement set within a minute²⁶ (Fig. 4), as confirmed by the attainment of the right mechanical integrity (ISO9917E), it is apparent that the internal phase transformations extend far beyond the scale of the mechanical setting. On the other hand, oscillations in compressive strength previously observed in a chitosan–DCPD cement similarly extended beyond the end of



Fig. 4 The initial (60 s) and the final (80 h) EDXRD patterns measured during the *in situ* analysis of the phase composition transformations accompanying the hardening of the Zn-TCP cement. The scattering vector, *q*, is related to the Bragg spacing, *d*, as $q = 2\pi/d$, and to the scattering angle, 2θ , as $q = (4\pi/\lambda)\sin(2\theta/2)$.



Fig. 3 Oscillations between the weight ratio of the two products of the setting reaction, brushite (DCPD, CaHPO₄) and scholzite (ZCP, $CaZn_2(PO_4)_2$) magnified down to scales of 1–30 h (a) and 35–80 h (b). Green rectangles in (a) denote regions in which oscillations between the two phases – *i.e.*, a decrease in the weight percentage of one phase paralleled by an increase in the weight percentage of the other phase – were detected. Blue and red arrows in (a) denote the direction of change in the weight ratios of DCPD and ZCP, respectively, within a given time window. Blue and red arrows in (b) denote the direction of change of the weight ratios of DCPD and ZCP, respectively, between two data points, demonstrating the "seesaw" effect, *i.e.*, an increase in the abundance of one phase at the expense of another in the entire 35–80 h time range.

the minutes-long setting time, into the 10-60 h time range, and correlated well with a change in the intensity of diffraction from the (120) plane,²⁷ suggesting that even the mechanical properties of the bulk, hardened cement are subject to change even after the hardening process is considered over per the standard, Vicat or Gilbert needle tests. As determined by neutron Compton scattering,28 such fluctuations in mechanical properties, specifically fracture toughness, extending over 62 h of the setting of a glass ionomer cement are paralleled by oscillations of atomic events, including hydration, fluorination, calcium binding and aluminium coordination change due to chelation by the polymeric phase, with the significant involvement of the amorphous matrix, whose inherent far-from equilibrium structure due to the trapping of the excess enthalpy during the quenching of the melt may be a possible factor predisposing cements for the displays of such nonlinear dynamics.

3.2. Cu-TCP cement

Unlike in the case of the Zn-TCP cement, where the initial reactant produced two different products in opposite paths, yielding an oscillation between the diffraction intensities and weight proportions of the two, in the case of the Cu-TCP cement, the initial reactant yielded a single reaction product.

Oscillations were observed, but in a more trivial manner, involving the diffraction intensity and concentration of the main product: DCPD. Fig. 5 thus shows the oscillation in the absolute intensity of the most intense, (041) reflection of the product. This intensity reaches a maximum on the 30th minute of the setting reaction, but then drops and eventually settles, after a series of oscillations, at 35% of this maximal value (Fig. 5a), at which point the oscillations have decreased in the intensity and fallen below the detection limit of the device. Similar oscillations, progressing toward an equilibrium point, were observed for two other major diffraction intensity peaks: (-242) and (241)/(062).²⁹ The magnitude of oscillations of the concentration of the product in the setting cement decreases over time (Fig. 5c), while the periodicity between the peaks in diffraction intensity increases (Fig. 5d). This periodicity increase follows an exponential route in the first 16 min of the setting reaction, before plateauing until the 35th minute, when it began to follow the exponential dependence until the last time point of the reaction (100 h). To exclude the possibility that the detected oscillations are of an instrumental error origin, it should be noted that they were absent in control solid samples (silicon, corundum) and that the absolute intensities varied minimally in the later stages of the setting reaction, e.g., from the 30th to the 100th hour (Fig. 6a), the standard deviation was



Fig. 5 Oscillations of the absolute intensity of the (041) reflection in the diffractogram of DCPD in the first 24 h (a) and 1 h (b) of the setting reaction time, oscillations of the rate of change of this intensity (c), and a gradual increase in the periodicity between the adjacent peaks in intensity over the first 80 h of the setting reaction time (d).



Fig. 6 Negligible oscillations in the absolute intensity of the (041) reflection in the diffractogram of DCPD between 23 h and 100 h of the Cu-TCP cement setting reaction time (a) and a negligible difference between XRD patterns of a control, amorphous CP cement hardening into HAp in the amorphous, $68-70^{\circ} 2\theta$ range at 5, 30 and 60 min setting timepoints (b). Only the 5 min pattern corresponds to intensity count values on the *y*-axis, whereas 30 and 50 min patterns are shifted by 10 and 20 count units upwards to avoid the overlap and accentuate the pattern similarity. A difference between the subtracted diffractograms taken at 60 and 30 min timepoints in the setting of the control, amorphous CP cement (c) and of the oscillatory, Cu-TCP cement (d). Diffractograms at 30 and 60 min timepoints are plotted in black and red, respectively, while the subtracted patterns are plotted in blue. The scattering vector, *q*, is related to the Bragg spacing, *d*, as $q = 2\pi/d$, and to the scattering angle, 2θ , as $q = (4\pi/\lambda) \sin(2\theta/2)$.

only 0.80% of the mean, as compared to 81.87% of the mean of the intensities in the first hour of the reaction. Simultaneously, a control cement comprising only an amorphous CP phase displayed a negligible difference between XRD patterns at some of the adjacent timepoints (Fig. 6b) during the 2 h long process of hardening into HAp. The comparison between the flat pattern obtained by subtracting two proximal timepoints (60-30 min) in this control (Fig. 6c) and the fluctuant corresponding pattern (Fig. 6d) in the Cu-TCP cement presents another evidence in favor of the veracity of oscillatory changes detected in the latter material. Following the linear portion of the evolution of the (-242)diffraction peak of DCPD in this early time range allowed for the determination of the Avrami exponent, n, and the reaction rate constant, k (Fig. 7). The more intensely nonlinear, oscillatory evolution of other analyzed reflections, including (041) and (241)/ (062), did not give satisfactory correlations. The Avrami exponent value of 2 is indicative of the edge-controlled nucleation mechanism, as opposed to 1 for surface and 3 for point sites, and agrees

with the earlier scanning probe microscopy studies demonstrating that step edges, not facets, are the docking sites for ionic growth units and molecular growth modifiers in DCPD.³⁰ The initial and the final XRD patterns of the Cu-TCP cement during the hardening reaction, captured after 60 seconds and 100 hours, respectively, are shown in Fig. 8.

In an attempt to detect the phase oscillations in the Cu-TCP cement using a complementary technique, the hardening process was monitored *in situ* using FT-IR. The overlay of spectra recorded at different timepoints of the setting reaction are shown in Fig. 9. A characteristic HPO₄^{2–} band, a signature of DCPD,³¹ was distinguishable at 874 cm⁻¹ and used to estimate the amount of DCPD in the mixture in comparison with the ν_3 PO₄^{3–} doublet at 1024 and 999 cm⁻¹ (Fig. 10a). The PO₄^{3–} triplet appearing at 914, 932 and 965 cm⁻¹ (ν_1) and the band at 450 cm⁻¹ (ν_2) are equally indicative of the presence of TCP.³² The same calculation was carried out for two adjacent bands: ν_1 (P–O) symmetric stretch of PO₄^{3–} at 970 cm⁻¹ and ν_1 (P–O(H))



Fig. 7 Linear fit of the DCPD crystallization extent as a function of time during the 5–50 minute time window in the setting of the Cu-TCP cement, along with the selected kinetic parameters derived from the fit, including the Avrami exponent, n, and the Avrami rate constant, k. The reaction extent, α , was measured on the (–242) diffraction peak of DCPD.

symmetric stretch of HPO_4^{2-} at 954 cm⁻¹ (Fig. 10d). In both cases, an exponential rate ($R^2 = 0.98817$ and 0.98262, respectively) of the increase in the amount of DCPD, the main reaction product, normalized to the amount of the main reactant, TCP, was derived (Fig. 10b and e). However, in the first hour of the reaction, an oscillation in the concentration of DCPD was observed, marked by a drop in the given band intensity ratio between the 15th and the 45th minute of the reaction (Fig. 10c and f), coinciding with a similar drop observed in the XRD analysis (Fig. 5b).

The P–O vibrations of HPO_4^{2-} are characteristically downshifted compared to those of the PO_4^{3-} ion due to the addition of a hydrogen atom to the P–O vibrational group. Namely, at 160.8 pm, the P–O(H) bond distance in HPO_4^{2-} is considerably longer than those of the remaining three P–O groups: 150.6, 151.6, and 153.2 pm.³³ Therefore, the appearance of the band at 954 cm⁻¹, originating from $\nu_1(\text{P–O}(\text{H}))$ symmetric stretch of the



Fig. 8 The initial (60 s) and the final (100 h) EDXRD patterns measured during the *in situ* analysis of the phase composition transformations accompanying the hardening of the Cu-TCP cement. The scattering vector, *q*, is related to the Bragg spacing, *d*, as $q = 2\pi/d$, and to the scattering angle, 2θ , as $q = (4\pi/\lambda)\sin(2\theta/2)$.

HPO₄^{2−}, on the account of the disappearance of the adjacent v_1 (P–O) symmetric stretch of PO₄^{3−} at 970 cm⁻¹ is indicative of the TCP → DCPD transformation (Fig. 10d). The similar, although less pronounced effect can be observed in relation to 916/931 cm⁻¹ v_1 P–O symmetric stretch (Fig. 11a) and 1168/1185 cm⁻¹ v_3 P–O antisymmetric stretch (Fig. 11b). Typifying all these transitions is the spectral reversal in the first 45 minutes of the setting time, suggesting oscillations in this early phase of the reaction, as in agreement with the findings of the XRD analysis.

As indicated by the transition of an initially broad OH⁻ stretch in the 2500–3500 cm⁻¹ region into resolvable multiplets (Fig. 11c), the setting reaction is accompanied by the structuration of water molecules. Given that water is a crystalline component of the DCPD stoichiometry, this effect is expected. However, as also seen from Fig. 11c, this structuration increases at each successive timepoint and does not proceed in a reverse direction between any two timepoints, thus lacking the key signature of oscillatory behavior. The multiplet structure is fully formed by the 24th hour and shows no change with the further increase in the setting time. In contrast, the $\nu(H)$ stretch in HPO₄²⁻ peaking at 2350 cm⁻¹ – albeit being a possible OH⁻ stretch combination band - transitions from a singlet to a doublet (Fig. 11d) in a way that resembles the oscillation in the amount of DCPD observed in the early time points of the setting reaction in the XRD analysis. Namely, the doublet is first seen at 1 h timepoint, after which it becomes noticeably less pronounced at 2 h and completely disappearing at 3 h, before reappearing again after 48 h (Fig. 11d). The relatively early timepoints at which the reaction reversibility is evidenced in FT-IR coincide with the peaking of the DCPD concentration at 0.6 h timepoint before its subsequent drop during the next hour, recovery for the next half an hour and then steadying around an equilibrium point (Fig. 5a and b). The origin of this doublet may be found in the fact that the hydrogen bonding of two water molecules of DCPD (W(1) and W(2) in CaHPO₄ \times 2H₂O) differs in strength and geometry.³⁴ An additional hydrogen bond in the DCPD crystal structure exists within the Ca²⁺-HPO₄²⁻ layer and



Fig. 9 FT-IR spectra of the setting Cu-TCP cement over the first 48 h of the setting reaction.

strong hydrogen bonding is expected to be responsible for the downshift of the OH⁻ stretch to below 3300 cm⁻¹. The absence of oscillations in this OH⁻ stretching vibration band suggests that water molecules *per se* are not central to oscillations, whereas the hydrogen bonded network of HPO₄²⁻ ions is.

3.3. Gelatin-HAp cement

Unlike the Cu- and Zn-TCP cements, which both transform to DCPD upon setting, gelatin-containing HAp cement undergoes a transformation to an amorphous CP phase during gelation and back to poorly crystalline HAp upon hardening (Fig. 12). In

this case, gelation partially dissolves and amorphizes the HAp crystals and initiates their setting back into the hardened, solid HAp form. As seen from Fig. 12, beyond the endpoint of the setting reaction, 8 hours after the onset of gelation, the phase composition of the product is identical to that of the solid reactant, comprising HAp only, except for the more prominent presence of the amorphous hump centered at $2\theta = 23^{\circ}$ in a generally noisier pattern and a lower average crystallite size estimated from the (211) peak halfwidth and the Scherrer equation (7.4 vs. 9.6 nm), suggesting the partially incomplete transition of the reactant back to its original crystalline state and the retention of some of the amorphousness of the gelled phase. The analysis of diffraction patterns measured after the initial setting time was over, *i.e.*, after $\sim 10^{\text{th}}$ minute of the reaction time, indicated a finite degree of oscillatory changes in crystallinity. Unlike in the case of Zn-TCP, where the oscillations lasted for over 80 h, i.e., for the entire duration of the experiment, or the case of Cu-TCP, where the oscillations wound down after \sim 24 h of the reaction time, the oscillations in the gelatin-HAp system were more short-lived and came to standstill in a matter of hours. This may be caused by the lack of aforementioned disparity between the solubility products of the product and the reactants, which allows for the transformation of the product back to the reactant to occur, along with the resulting oscillations in the phase content. Still, the oscillations were intense, particularly between the 10th and the 120th minute of the reaction, as seen in Fig. 13. The same extension of the period between the adjacent peaks in intensity over time as that observed during the setting of Cu-TCP cements (Fig. 5d) was present. Fluctuations in the diffraction peak intensity (Fig. 14a) entailed those in crystallinity (Fig. 14b). No significant



Fig. 10 FT-IR *in situ* phase composition analysis of the setting of the Cu-TCP cement over the first 48 h, showing the $v_3(PO_4^{3-})$ (a), $v_1(PO_4^{3-})$ (d) and different $v_1(HPO_4^{2-})$ (a, d) stretching vibration bands changing over time as well as the intensity ratios between the 874 cm⁻¹ $v_1(HPO_4^{2-})$ and the 1024 cm⁻¹ $v_3(PO_4^{3-})$ bands (b and c) and between the neighboring 954 cm⁻¹ $v_1(HPO_4^{2-})$ and the 968 cm⁻¹ $v_1(PO_4^{3-})$ bands (e and f). Green arrows denote the direction of change (increase or decrease) of the given intensity ratios as function of time.



Fig. 11 FT-IR spectra of the setting Cu-TCP cement over the first 48 h of the setting reaction, showing adjacent 916 and 931 cm⁻¹ ν_1 P–O symmetric stretches of HPO₄²⁻ and PO₄³⁻ ions, respectively (a) and 1168 and 1185 cm⁻¹ ν_3 P–O antisymmetric stretches of HPO₄²⁻ and PO₄³⁻ ions, respectively (b), along with the OH⁻ stretch in the 300–3700 cm⁻¹ spectral region (c) and the H stretch in HPO₄²⁻ with the doublet at 2341 and 2359 cm⁻¹ (d).

difference was observed between absolute and integrated (211) peak intensities (Fig. 14a), justifying the interchangeable use of the two in determining the phase composition. After transiently peaking only 2 minutes into the setting reaction and reaching 20% per eqn (3), crystallinity drops down to 5% on the 10th minute of the reaction, paradoxically coinciding with the first peak in the amount of the solid phase. Crystallinity then increases, but does not exceed 17% before setting, after 90 min,



Fig. 12 XRD patterns of hardened HAp-gelatin cement (top) and of its powder precursor (bottom). Reflections derived from HAp are marked with Δ .

at ${\sim}10\%,$ which is the typical value observed for apatites synthesized by precipitation under ambient conditions. 35

4. Discussion

In this study, we report on the nonlinear, oscillatory dynamics detected in the evolution of phase composition during the setting of different CP cements, two of which evolved toward DCPD and one of which evolved toward HAp as the final product. Whereas both DCPD-forming cements contained iondoped β-TCP as the initial phase, the Zn-TCP one yielded ZCP as an additional phase during setting and the oscillations between these two products were pronounced throughout the entire 80 h setting period. The physicochemical cause of the oscillatory dynamics in each of these three distinct cases is difficult to pinpoint. In the case of the oscillations observed between DCPD and ZCP, a first-order reaction model given in eqn (7) cannot oscillate without the addition of a finite number of sub-processes feeding into this global pathway. As a result, no matching with the classical chemical oscillatory models could be found. Although the evolution of the β -TCP systems does resemble the Brusselator dynamics approaching a fixed point in a stable regime, no fit with the model was observed based on the given assumptions. Another argument in disfavor of the "chemical clock" mechanism of oscillation is that such



Fig. 13 Time-lapse diffractometric analysis of the setting gelatin–HAp cement.

oscillations are limited to intermediary species, whereas in this case the final phases are observed to oscillate. Notwithstanding the possibility that phases more stable per the Ostwald–Lussac rule may present the truer final phases, albeit minorly present and undetectable using XRD, gravitation of oscillations toward an equilibrium point is suggestive of a different, less exotic mechanism.

Oscillations between different equilibrium states in a liquidto-solid phase transition were previously noticed and attributed in origin to diffusive fluxes at the interface between phases.³⁶ However, surface diffusion anisotropy can only modify the existing oscillations, but cannot cause them. To arise, oscillations require two diametrically opposite and competing processes, *e.g.*, dissolution and crystallization. Synchronization enabled by the local variations in the chemical potential³⁷ due to competing crystallization and dissolution reactions may be responsible for sustaining the nonlinear, oscillatory dynamics at the solid/liquid interface and can lead to pattern formation, but, again, cannot be a key thermodynamic driver behind them. Oscillations were also observed at the particle–solution interface during precipitation and dissolution reactions and were correlated with autocatalysis or reaction inhibition by the

product formation.³⁸ However, the reaching of the equilibrium state and the absence of autocatalytic behavior are hallmarks of solid-state oscillators compared to the molecular ones. In all three cases reported here, the oscillatory reaction eventually settles in an equilibrium state, suggesting the absence of a constant influx of the driving energy for the phase transitions between reactants, products and possible intermediates.³⁹ Still, the compositional oscillations leading to this equilibrium, caused by the relatively high reverse reaction rates, were pronounced to the point of disallowing any correlations in isothermal kinetic models commonly used to study the crystallization kinetics in cements, e.g., JMAK⁴⁰ or Šesták-Berggren. The only exception was the evolution of the (-242) DCPD reflection in Cu-TCP cement in the first hour of the reaction time, allowing for the extrapolation of the Avrami exponent and estimation of the DCPD crystallization rate and mechanism (edge-controlled growth). A direct correlation between (i) a gradual decrease in the oscillation intensity and (ii) a gradual drop in the concentration of both the amorphous solid phase and the liquid phase, present only in the initial stages of the reaction, suggests that the amorphous phase, both liquid and solid, must be a key intermediate and that the process is



Fig. 14 Variations in the absolute (\bullet) and integrated (\blacktriangle) intensities of the most intense, (211) reflection (a) in the diffractogram of gelatin–HAp cement and of its crystallinity (b) in the first 2 h of the setting reaction.

primarily entropically driven. Slowing down of the kinetics of crystallization in an Fe–Co alloy was similarly associated with a reduced presence of the amorphous matrix, which was proven to be necessary for the grain growth.⁴¹ The fact that even in the final, hardened form, the materials do not exhibit exceptional crystallinity (~10%) suggests that the amorphous content plays a crucial role in ensuring these compositional oscillations by providing both a source from which individual phases gain their content and an end to which they lose it.

Numerous kinetic effects can be invoked as candidates for roles in limiting the diffusion and inhibiting the phase transformation reactions. One role of additives - Zn²⁺, Cu²⁺ and gelatin - may have been to decelerate the phase transformations and make them detectable under regular measurement conditions. While Zn^{2+} and Cu^{2+} exerted this effect by entering the crystal lattice of β -TCP, gelatin did so by coating the CP grains and partially blocking the growth and recrystallization via dissolution/reprecipitation mechanism. A similar blocking effect was previously observed when HAp nanoparticles were interspersed in a chitosan matrix.42 Oscillations at the scale observed in gelatin-HAp system were not observed in pure HAp cements,43 suggesting that either an organic additive in a monophasic CP product system or the complex entwinement of product formation pathways are needed to promote such oscillations. This raises expectations that CP remodeling process and interaction with cells and tissues, naturally interfaced by organic layers, is naturally oscillatory at fine spatiotemporal scales.

Zinc, for one, is a cation that decreases the solubility⁴⁴ and inhibits the crystal growth⁴⁵ of CPs. Both of these effects effectively slow down the phase transformations in the system and may extend the timescale of oscillations. The mechanism of structural stabilization caused by doping with Zn²⁺ may be related to the formation of intracrystalline O-Zn-O bridges.46 However, the role of Zn²⁺ as a cationic catalyst of phase transformations causative of these oscillations cannot be excluded either. Given that $Cu^{2+} \rightarrow Ca^{2+}$ appears to have the opposite effect on solubility,47 it can be concluded that ionic substitution may enhance, but not cause the oscillatory dynamics. Next, both final phases of the reaction, DCPD and ZCP, are dihydrates, unlike β -TCP, the initial reaction component, which is an effect lowering the entropic barrier that prevents the dissolution of the product and its partial transformation back to the reactant. The solubility products of these two phases are also in the similar range: $-\log K_{sp} = 9.46$ for ZCP⁴⁸ and $-\log K_{sp} = 6.59$ (ref. 49) for DCPD. In contrast, the solubility of β -TCP $(-\log K_{\rm sp} = 29.50 \text{ (ref. 50)})$ is lower and this uncommon property of the system (*i.e.*, the products being more soluble than the reactant) may be another factor that favors the oscillatory behavior, which feeds on the ability of the product to transform back to the reactant in relatively copious amounts. The effect of citrates present in the setting mixture on this disparity, which potentially favors oscillatory dynamics, is difficult to assess. Namely, on one hand, based on the ionic and solubility product calculation using the Debye-Hückel limiting law, the acidification effect due to citrates is such that it would make TCP undersaturated at pH < 2.7 and DCPD the only supersaturated

CP phase at pH < 2.15. On the other hand, in the viscous cement where protonation is hindered, this effect may be negligible and the citrate may predominantly act as a nucleation inhibitor that slows down the crystallization dynamics and sets off the oscillatory behavior.

The interfacial energy fluctuation argument offers another teleological perspective at the origin of these dynamic instabilities. Variations in the surface tension previously paralleled the periodic movement of ionic species across the biphasic interface consisting of an amphiphilic, cetyltrimethylammonium bromide monolayer and the resulting oscillations in electrical potential⁵¹ and pH,⁵² but their underlying cause was still atomistic in nature. Interfacial tension has also been associated with the Marangoni instability, but again being caused by local compositional fluctuations.53 Phases with high interfacial energies tend to be more unstable and more prone to dissolution than the low energy surfaces, whose tendency is to grow. Although amorphous CP and HAp may be compositionally highly similar or even identical, the former phase will have a lower interfacial energy than HAp because of the higher degree of entropic similarity with the liquid phase, and its growth potential can be assumed to be higher than the dissolution potential when compared against HAp. These interfacial energy propensities are at odds with the thermodynamic bulk propensities, which dictate that HAp would be less soluble than amorphous CP because of the lower solubility product: $-\log K_{sp} =$ 114.7 for HAp and \sim 25 for amorphous CP. This argument is, however, not specific to this system and applies to every crystallization system, requiring the addition of kinetic effects thereto. It should be noted that similar oscillations involving an alternate dominance of dissolution over precipitation and the other way around were noticed during the regular immersion of CP powders in aqueous solutions. For example, during the first 100 h, Ca²⁺ levels in the supernatant fluctuated between 80 and 30 ppm.54 Hence, aside from these thermodynamic potential causes of oscillations, kinetic factors must be crucial in terms of enabling their temporal extension and detection.

Unlike in the case of ion-doped cements, oscillations in the gelatin-HAp system occur between (i) dissolution of HAp into growth units solidifying as amorphous CP and (ii) ripening of amorphous CP into HAp. Ripening may involve the martensitic transformations of the crystal lattice in the solid state, but may also have dissolution/reprecipitation as the dominant mechanism of phase transformation. The interrelation between the three aspects of the process - dissolution, reprecipitation of the amorphous phase, and recrystallization - must be precisely tuned for the oscillations to be observed under ordinary conditions. Usually, in oscillatory processes, one of the three interrelated effects is delayed compared to the other two, which often proceed simultaneously. Even with this assumption, however, it would still not be clear which feedback effect synchronizes these oscillations and ensures than they are not averaged over the entire volume of the material. For, although ensembles of atoms forming surface islands of nuclei fluctuate in size and the chemisorption energy and reactivity oscillate as a function of size,55 the total surface area of each chemisorbed phase remains constant over time.56 Still, the same effects governing these local fluctuations can be considered as

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potential candidates for the causes of oscillations at more global scales. Thermal feedback effects have been causative of oscillatory behavior in Ti-Zr alloys, but during oxidation at elevated temperatures.57 At the same time, hydration of CP cements is an exothermic process, liberating heat estimated at \sim 200 J g⁻¹ and being inversely proportional to Ca/P molar ratio of the CP phase.⁵⁸ In general, hydrolysis of CPs is an exothermic process and the crystallization, conversely, endothermic when solute speciation is taken into account, the reason for which boiling is often added to ambient CP precipitation protocols to send the solution over - not below as for most other salts - the saturation level and precipitate ions that will block the particle surface and increase stability. In one such scenario, the transformation of amorphous to crystalline CP would, thanks to its endothermic effect, transiently undersaturate the system and promote the dissolution of the material and/or its transformation back to the amorphous state, thus potentially causing oscillations of this kind in the system. However, even though precipitation of HAp, being an endothermic process, would consume the heat and lower the cement temperature, thus increasing its solubility and causing the destabilization of the precipitate, it is not certain whether this thermal effect is large enough at room temperature to trigger and sustain the oscillations in the phase composition and crystallinity.

The selective transfer of ions as growth units is another possible factor affecting oscillations. Namely, Ca²⁺ ions are more mobile than $H_x PO_4^{x-3}$ ions, which tend to reside on surfaces longer and present the starting points for nucleation of CPs.^{59,60} Per the Ostwald-Lussac rule, the growth of a most stable phase proceeds through successive growth and dissolution of less stable ones in the direction spanning from the least to the most stable. At the same time, the solubility of CPs is mostly inversely related to Ca/P molar ratio, with solubility decreasing in the following order: Ca(H₂PO₄)₂ (MCP) > CaHPO₄ $(DCP) > Ca_3(PO_4)_2$ $(TCP) > Ca_{10}(PO_4)_6(OH)_2$ (HAp). With Ca^{2+} ions dominating the first wave of growth species arriving at the growing surface, its Ca/P molar ratio is shifted to higher values, which counteracts dissolution and favors stability. However, the high interfacial energy of this layer with respect to both the bulk region of the particle and the environment is unsustainable and favors the opposite effect, namely that of dissolution and destabilization, potentially causing alternate waves in solidification of a new phase and its dissolution. This propensity of the high interfacial energy phase, in this case HAp, to dissolve is additionally intensified by its naturally nanoparticulate form initially, given that HAp has a natural tendency to solidify as nanocrystals, which only subsequently aggregate and ripen into larger units. Because of their high surface-to-volume ratios and corresponding reactivities, nanomaterials should be intrinsically predisposed to exhibit dynamic instabilities in their formation pathways.

The most plausible argument, however, may be that pertaining to the viscosity of the liquid phase surrounding the growing grains. As the viscosity of the liquid and semisolid pockets within the setting cement increases as the setting time progresses, so does the diffusion rate drop and the oscillations become less pronounced. Therefore, it can be concluded that the

nonlinear behavior is viscosity-controlled and diffusion-limited, and a possible mechanism, at least in the case of Cu-TCP and gelatin-HAp cements, would be such that the decrease in viscosity associated with the crystallization of DCPD and HAp, respectively, slows down diffusion and growth and promotes homogeneous nucleation of the low-energy phase in the liquid portion of the cement. The alternate dissolution and recrystallization of this low-energy phase, in this case amorphous CP, accompanied by the initially intense and subsequently subtle waves of viscosity, may give rise to oscillations in the composition. In addition to the bulk diffusion, however, which must have a role in controlling the intensity of fluctuations, surface diffusion, being more prone to exhibition of kinetic effects, is another possible source of oscillatory behavior. Speciation of growth units near the surface, including Ca^{2+} , $H_rPO_4^{x-3}$, $OH^$ and Na⁺ coming from the liquid phase of the cement, could also play a role. Oscillations of ionic fluxes and electrical potential in Martin's oscillator⁶¹ – involving a hypodermic syringe filled with a physiological solution, having a plunger removed and being immersed vertically in pure water - cannot be due to interfacial tension or diffusion potential, and are thought to be due to periodic changes in charge separation in the electrical double layer caused by the mobility of its diffusive outer portion.62 However, this explanation was derived for glass capillaries, a material that is relatively smooth compared to the uneven surface of CP grains and that carries a significantly higher surface charge than CP under physiological conditions: ζ < -20 mV (ref. 63 and 64) vs. $-10 < \zeta < 10$ mV,⁶⁵ respectively. Finally, profiles of the liquid order, including hydrogen bond and dipole ordering, forming in response to the electronic and geometric features of the solid surface, are typified by fluctuations at very short, pico- and femto-second timescales. Although they are finely localized and transient, it is conceivable that their superimposition can yield periodicities at more robust scales. Notwithstanding the same limitations applying to the double layer argument, the probing of one such interfacial order at the ionic and molecular scales typified by fine localization and low free energies may be the basis for a more comprehensive model of the solid/liquid interface and the route to explaining these effects from a more quantitative and more sophisticatedly empirical perspective.

Finally, a number of possible effects causing or contributing to the oscillatory dynamics have been listed, but it is possible that, in analogy with oscillatory chemical reactions, where no less than three interrelated intermediates in a feedback-looped network must be present, a sufficient number of thermodynamic drives and kinetic variables, each affecting the oscillations while being related to all the other variables, is needed to properly explain these effects. Attempts to explain oscillations at the interface were similarly based on invoking three distinct but interrelated types of flow: the flow of fluid, the flow of solutes and the flow of current.⁶⁶ In all scenarios, however, for the oscillations to be detected at the timescales of minutes to hours, inhibitory effects through the use of right chemical additives and physical conditions that impair diffusion must be imposed. Phase transformations in CPs, in general, are known to be comparatively slow,67 while the

setting reaction itself is a relatively slow transformation, taking place over minutes and hours. Given the comparatively slow chemical reactions being a hallmark of living systems, CP are a convenient building block for biomineralization and this may be one of the reasons why CPs coevolved with the vertebrates into a major biomineral on Earth. In addition, oscillations in the composition lasting for the entire 80-100 h of the setting reaction, well over the time at which the cement visually appears to have fully hardened and gained a sufficient mechanical integrity, agree with the idea that CPs are "living" materials, exhibiting internal changes perhaps more pronouncedly than most other solids. One could hypothesize that it was for this reason that CP was evolutionarily selected as the major mineral component of vertebrate bones and teeth. Intrinsic preference for the nonlinear dynamics may indirectly explain the coevolution of CP and the higher organisms. For, there may be no better material to endow with the attribute of "living" than the one that exhibits nonlinear, oscillatory behavior, which emerges as more dominant with every higher level of complexity of living systems. If oscillatory reactions are primitive prototypes of self-organization phenomena, thanks to which inanimate matter spontaneously transformed to living matter before the evolution even began, then, observed in a material, they might be perceived as a gateway to a new generation of biomaterials, capable of engaging in a more intimate relationship with the biological milieus.

5. Conclusion

Oscillations present more of a rule than an exception in the biological domain. This is understandable in view of the fact that oscillations are, ultimately, a form of collective dynamic behavior that marks the transition from the potentiation of an entity to the potentiation of a relationship and represents the first step in the animation of an otherwise inanimate matter. To be thermodynamically open and autopoietic, the biological system must be built around the concept of causal chains, meaning that the nonlinear, oscillatory dynamics cannot be foreign to it. The very fact that feedback loops must be present in a cell, the basic unit of biological systems, to enable its sensing functions and the ability to respond to environmental stimuli in an intelligent manner implies that cyclical causalities, including oscillatory behavior, are an integral trait of the cell and of the more complex biological wholes that it constitutes. Spatiotemporal oscillations regularly figure in intracellular processes that include the dynamics of axonemes, dynein motors, microtubules and other aspects of cytoskeletal motility; signal transduction pathways involving mechanosensitive ion channels; transmembrane potential regulation; spatial structuration of mitotic cells; gene expression; energy metabolism; and others.⁶⁸ They also govern cell aggregate and organism behavior, where they range from cardiac electrical activity to ventricular pressure to optokinetic nystagmus and multiple other features of sensory perception to circadian rhythm to fertility to population dynamics and beyond.

Here we report on the detection of oscillatory behavior in three different calcium phosphate cements, each involving a mechanistically distinct pathway. Specifically, during the setting of the zinc-doped whitlockite cement, this initial phase transformed into brushite and scholzite along two different reaction paths and created an 80 hour long "seesaw" effect where the two products, brushite and scholzite, oscillated in concentration in the solid phase. Oscillations in the coppercontaining whitlockite cement were more simplistic in nature and involved the sinusoidal variations in the concentration of brushite, the main product of the hardening reaction, as it gradually progressed toward an equilibrium point. Finally, the hydroxyapatite-forming cement exhibited oscillations in crystallinity on the timescale of minutes within the timeframe between the onset of gelation and the endpoint of the setting reaction. It is conceivable that the same oscillatory character of phase transitions occurs as a part of apatite biomineralization and bone remodeling processes, in which case the systems presented here may be decent models for the study of the biological and perhaps medical benefits of such oscillations. The conception of bone replacement materials and tissue engineering constructs capable of engaging in the dynamics of integration with the natural tissues in compliance with this oscillatory nature may open a new avenue for the future of this type of medical devices. It may also be an impetus for the greater involvement of complexity science, systems biology and cybernetics in the design of biomaterials. Albeit parlous, one such exotic research direction will lead to biomaterials far more describable with the epithet of "living" and worthy of the prefix "bio-" than their current generation. Such a subtle animation of matter may bring about groundbreaking medical benefits.

Conflicts of interest

There are no conflicts to declare.

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