The mesoscale order of nacreous pearls

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A pearl’s distinguished beauty and toughness are attributable to the periodic stacking of aragonite tablets known as nacre. Nacre has naturally occurring mesoscale periodicity that remarkably arises in the absence of discrete translational symmetry. Gleaning the inspiring biomimetic design of a pearl requires quantifying its structural coherence and understanding the stochastic processes that influence formation. By characterizing the entire structure of pearls (~3 mm) in a cross-section at high resolution, we show that nacre has medium-range mesoscale periodicity. Self-correcting growth mechanisms actively remedy disorder and topological defects of the tablets and act as a countervailing process to long-range disorder. Nacre has a correlation length of roughly 16 tablets (~5.5 μm) despite persistent fluctuations and topological defects. For longer distances (>25 tablets, ~8.5 μm), the frequency spectrum of nacre tablets follows f−15 behavior, suggesting that growth is coupled to external stochastic processes—a universality found across disparate natural phenomena, which now includes pearls.

nacre | pearls | TEM | SEM | mesoscale

The artist Jorge Méndez Blake illustrated a basic principle of structure in mesoscale periodicity and grown nanocrystalline materials when he placed a single book (i.e., a defect) in a brick wall, displacing every subsequent brick layer (1). Blake’s work demonstrates the principle of paracrystallinity: disorder from any one defective site propagates throughout the layers of the material (2–4). In the context of brick laying, skilled masons overcome this difficulty using external templates to achieve translational order—marked guideposts and facing cords align layers to prevent disorder from propagating (5, 6). Only with great calculated effort can a 10-story building ensure the same number of aligned brick layers on all sides (7). Regarding layered growth of nanomaterials, of course, external templates are limited and rare. Mesoscale periodicity—defined here as long-range translational order of mesoscopic building blocks—is thus improbable, due to natural variations in the unit sizes, without the aid of some additional countervailing mechanism. The rare instances where nature assembles mesoscale periodicity therefore merit our attention (8). Nacre in pearls and mollusk shells are one such example where mesoscale periodicity arises in an environment without discrete translational symmetry (9–11).

Pearls are renowned and coveted for their beauty; that beauty results from the diffracting iridescence of periodically stacked tablets (~500-nm-thick units) known as nacre (12–15). Nacre tablets form within interlamellar organic sheaths (~10- to 20-nm thickness) that act as the main patterning agent between layers. During formation, not-yet-calciﬁed layers grow within multiple closely spaced sheaths that separate progressively during mineral growth (16, 17). While some work has focused on describing the atomic-to-nanoscale structure of the tablets themselves as assemblies of nanoparticles of aragonite (18–21) or the atomic order within and between tablets (22), the current work is focused on how these tablets are arranged at the next length scale (hundreds of nanometers—several microns [i.e., herein, the mesoscale]).

Mesoscale periodic nacre is a highly ductile structural composite that can withstand mechanical impact and exhibits high resilience on the macro- and nanoscale (23–25). Because of its superior toughness, nacreous shells protect the mollusk’s soft body (11, 26–30) and inspire scientists designing next-generation supercomposites (31–33). Yet, despite a century of scientiﬁc fascination with nacre (9, 11, 24, 26, 34), its astonishing mesoscale periodicity has not been quantiﬁed, leaving key questions about this material unanswered: does nacre have long-range order? What is the stochastic process that governs its formation?

Here, we show nacre forms remarkable medium-range mesoscale periodicity through corrective processes that remedy disorder and topological defects. The entire nanostructure of nacreous pearls is characterized in cross-section to reveal complex stochastic processes that inﬂuence ordered nacre growth. Beginning atop an initially formed organic center, nanoparticles self-assemble into bulk aragonite, later followed by nacre deposition. The initial layers are disordered; however, this disorder attenuates within the first 200 layers through corrective growth processes that persist throughout the entire pearl. When a tablet is grown too thin, the next tends to be thicker—and vice versa. Thus, in an environment without discrete translational symmetry, there remains a fundamental question of whether long-range order exists. In this work, the stochastic growth that leads to mesoscale order in the nacreous pearl is revealed by quantifying its structural coherence across entire pearl specimens. We ﬁnd that mollusks strike a balance between preserving translational symmetry and minimizing thickness variation of layers by creating a paracrystal with medium-range order. Self-correcting growth processes allow pearls to quickly attenuate disorder, accommodate topological defects in tablet structure, and maintain order throughout a ﬂuctuating external environment. These observations were made possible by characterizing the entire structure of Akoya “keshi” pearls (~3 mm) at high resolution (~3 nm).

Significance

Despite a century of scientiﬁc fascination with nacre’s periodic mesostructure, there remains a fundamental question of whether long-range order exists. In this work, the stochastic growth that leads to mesoscale order in the nacreous pearl is revealed by quantifying its structural coherence across entire pearl specimens. We ﬁnd that mollusks strike a balance between preserving translational symmetry and minimizing thickness variation of layers by creating a paracrystal with medium-range order. Self-correcting growth processes allow pearls to quickly attenuate disorder, accommodate topological defects in tablet structure, and maintain order throughout a ﬂuctuating external environment. These observations were made possible by characterizing the entire structure of Akoya “keshi” pearls (~3 mm) at high resolution (~3 nm).

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versa—thus compensating for the initial error. Local irregularities in thickness (±15% variation of interfacial curvature) and topological defects (5.9 × 10^8 m^-2 defect density) intermittently appear, yet the pearl maintains medium-range order with a correlation length of ~16 tablets (~5.5 μm). At larger distances, the positions of tablet layers are uncorrelated. However, a pearl does not have perfect translational symmetry, and local disorder persists into subsequent layers—a paracrystalline property which ultimately limits long-range order. For longer-length scales (e.g., >25 tablets, ~8.5 μm) the aperiodic fluctuation of nacre’s tablet thicknesses follow f^-1.5 noise behavior [f = 1 / (number of tablets)], which could represent external Markov processes with longer memory attributable to cooperative environmental changes such as temperature, pH, food availability, seasonality, and tidal cycles.

Results
The Start of Nacre in Pearls. Pearls form either as a natural response to mantle tissue injury or when mantle tissue is deliberately transplanted from a donor into a host animal for pearl culturing. In both cases, the mantle epithelium develops a closed cyst—the so-called pearl sac—programmed to reproduce the structure of the shell (35). CaCO_3 is then secreted onto a manufactured bead (for bead-cultured pearls) or any available debris enclosed by the developing pearl sac (for nonbeaded pearls) within a limited space (36). The shape of the manufactured or natural nucleus usually dictates a pearl's macroscopic shape (36, 37) (SI Appendix, Figs. S1–S5). Here, we primarily study non–bead-cultured “keshi” pearls, grown for ca. 18 mo in Pinctada imbricata fucata mollusks on the Eastern shoreline of Australia (Methods) (38).

The cross-section of a nonbead “keshi” pearl is shown in Fig. 1B. The mollusk deposits calcium carbonate upon the irregular organic center (~200-μm diameter) and after roughly ~15 μm of bulk aragonite, nacre layers begin to form. Nacre layers are bonded by chitin-proteinaceous organic sheaths periodically secreted progressively in the direction toward the margin by the epithelium (39–41). Ordered nacre makes up 87% of this 2.5-mm pearl (Fig. 1B and C). However, the nacre is preceded by more disordered growth stages. Nacre and other biominers form by nonclassical crystallization via metastable transient precursor phases (42, 43) and attachment of CaCO_3 nanoparticles.

Fig. 1. Formation of non–bead-cultured Akoya “keshi” pearls produced in a P. i. fucata mollusk. (A) Optical overview of a nonbeaded keshi–cultured pearl showing iridescence due to the interplay of constructive interference at nacre tablet layers with light. (B) Cross-section showing CaCO_3 growth begins onto an organic center. (C) Mature nacre (purple box in B) showing the ordered state in their thickness and interface curvature. (D and E) Atomic-resolution ADF-STEM of mature nacre and its corresponding Fourier transform indicating highly crystalline nacre and a lattice constant that is consistent with aragonite. (F) Cross-sectional backscatter SEM at the center of the pearl (yellow box in B) showing transition from spherulitic aragonite structures to nacre. (G and H) Aggregated nanoparticles (at time of observation) form massive, structurally indistinct aragonite structure. (I) Formation of nacre begins directly on massive, structurally indistinct aragonite.
interspersed by organic macromolecules (Fig. 1 G and H and SI Appendix, Fig. S6). The nanoparticle packing intermittently increases until forming columnar aragonite with segregated regions of organic. In Fig. 1B, this occurs over roughly 200 μm of growth. Throughout this process, the pearl's volume becomes larger and rounder, before penultimately forming a bulk aragonite (Fig. 1C). In most observations, we observe bulk aragonite begins after a region reaches positive curvature (44–46).

In pearls, nacre forms abruptly on bulk aragonite. At the start of nacre, around 440 μm (±12%) from the organic center in Fig. 1, the bulk aragonite-to-nacre interface is approximately flat (curvature of ~0.002 μm⁻¹) with locally rough texture (~200-nm variation over 7 μm). About 10 to 20 μm prior to the first nacre layer, larger organic deposits appear in comparable quantity to the interlayer sheaths that appear in mature nacre—suggesting the genetic processes directing the system toward sheath deposition start earlier than the structure itself (SI Appendix, Fig. S9). This same transition observed in Pinctada fucata pearls was also found in nonbead cultured Tahitian “keshi” pearls produced by the Pinctada margaritifera mollusk (SI Appendix, Fig. S8). This direct nacre growth onto bulk aragonite in pearls is reminiscent of P. fucata, Unio pictorum, and Nautilus pompilius shells but is distinct from nacre formation reported in Pinna nobilis, where the assembly process is driven by aggregation of nanoparticles within a several-micrometers-thick organic matrix (11, 47, 48).

**Mesoscale Periodicity of Nacre.** The first nacre tablets are non-uniform, with substantial thickness variation and a rough interface with the preceding bulk aragonite. The Fourier transform of early nacre layers imaged by backscattered electron scanning electron microscopy (BSE-SEM) reveals ±15° angular variation in their interface curvature (Fig. 2A, Inset). This angular broadening decreases quickly to ±6° after 100 layers and further down to ±5° after 200 layers (mature nacre). This smoothing of tablet interfaces is clearly visible (Fig. 2A and SI Appendix, Fig. S10). At the same time, tablet thickness decreases by 30%, reducing from 500 ± 300 nm for early nacre to 340 ± 120 nm (a ~40% reduction of variance) for mature nacre. In mature nacre (Fig. 2A), peaks visible in the Fourier transform (first peak at (~340 nm)⁻¹) provide the signature presence of mesoscale periodicity (i.e., translational symmetry of tablets). However, these peaks in Fourier space do not quantify the long-range order nor describe the stochastic processes that create the mesoscale periodicity.

Pair-correlation functions quantify the mesoscale periodicity of pearls in nacre along the growth direction (Fig. 2B) by measuring the probability of tablet spacings (Methods). Subsequent peaks indicate the distance to the first nearest neighbor (NN), second NN, third NN, etc. These plots were calculated based on all pairs of tablets in each stage of nacre (Fig. 2A). The first peak represents the average distance between adjacent tablets, and subsequent peaks describe longer-range periodic order.

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**Fig. 2.** Quantification of mesoscale periodicity in nacre. (A) Cross-sectional backscatter SEM of the early, middle, and mature stages of nacre growth shows ordering through reduced variation in the interface curvature and tablet thickness. Fourier transforms (Right) of nacre (Left) imaged by BSE-SEM show that angular broadening decreases from ±15° to ±5°. (B) Pair-correlation functions of nacreous layers represent the probability of finding tablets spaced a given number of unit cells apart. Sharpening of peaks in later nacre indicates increasing long-range order. (C) Correlation of the thickness of tablets with nearest neighbors shows a negative correlation. If one tablet is thick, the next one tends to be thin. (D) Cumulative disorder in nacre described by a real paracrystalline model, demonstrating that nacre has order between that of a crystal and a paracrystal model.
For ideal crystals, every peak will appear equally sharp. However, in nacre, peaks in the pair-correlation function increasingly broaden—indicating propagation of disorder, as described by the paracrystal model (4, 49, 50).

Nacre is well-described by an ordered paracrystal or a medium-ranged mesoscale crystal model. We report mature nacre has a correlation length of 5.5 μm, and translational order is lost by 16 layers. Correlation length is defined as the length where the pair-correlation envelope disappears (see Methods and SI Appendix, Fig. S11). Thus, although nacre maintains extraordinary translational symmetry across a dozen or so layers, it lacks true long-range order. In paracrystals, a deviation in layer thickness displaces subsequent layers and thus disorder propagates. Defects and tablet variation degrade translational symmetry with distance. The progressive broadening of each peak in the pair-correlation functions quantifies the long-range order and paracrystallinity (Fig. 2D). In a perfect paracrystal, the SD of peaks, $\sigma_n$, broadens by the square root of its peak number ($\sigma_n = \sigma_1 \sqrt{n}$) (4, 51).

Nacre, at the mesoscale, has both crystalline-like and paracrystalline disorder. As shown in Fig. 2D, nacre falls below the paracrystal curve (more ordered) and above the flat crystalline curve. The paracrystallinity, or broadening of peaks, is approximately equivalent in early, mid, and mature-stage nacre, suggesting the growth mechanisms may be equivalent. However, the sharper first-order peaks in mature nacre reflect improved initial conditions.

Nacre maintains discrete translational symmetry through self-correcting growth processes. The bumps and valleys in one nacre layer are attenuated in the subsequent layers. If one tablet is thicker than average, the next tends to be thinner. This negative correlation of thickness between adjacent nacre layers is quantified in the early, mid, and mature nacre in Fig. 2C. The negative correlation ($-0.66, R^2 = 0.46, P = 10^{-24}$) follows an autoregressive model that allows ordered periodic growth to reach steady state in ~200 layers, which demarcates maturity. In uncorrelated growth, thickness variation of each nacre tablet would devastate the periodic long-range order—much like Jorge Méndez Blake’s brick wall (1). Unlike a brick wall, multiple nacre layers grow simultaneously within continuous organic sheaths. Ultimately, nacre is able to self-correct for fluctuations in tablet thicknesses and occasional screw dislocations (Fig. 3 A and B and SI Appendix, Figs. S12 and S13).

**Topological Defects in Nacre.** Local disorder in pearls encompasses topological defects (i.e., dislocations) of the tablets. SEM images at the pearl surface and a corresponding schematic illustrate how screw dislocations originate in nacre (Fig. 3 A and B and SI Appendix, Figs. S12 and S13). High-resolution annular dark-field scanning transmission electron microscopy (ADF-STEM) images show a dislocation in cross-section (Fig. 3 C–L and SI Appendix, Fig. S13). Screw dislocations in nacre are known (52–54) and recently reported to couple as chiral pairs through a dissipative distortion field that helps accommodate the space filling requirements of nacre (34). In cross-sections, screw dislocations can be misidentified as edge dislocations (although both may be present). When viewed parallel to the slip plane, a mineral bridge at the dislocation marks the defect origin. Viewed normal to the slip plane the screw dislocation appears as an additional partial organic boundary. The defect density (54) of nacreous pearls herein is $5.3 \pm 0.4 \times 10^6$ m$^{-2}$ in plane view and $6.5 \pm 0.4 \times 10^6$ m$^{-2}$ in cross-section (Fig. 3A and SI Appendix, Fig. S14). Our quantification is measured from top view and cross-
sectional SEM by dividing the total number of defects by the total area (SI Appendix, Fig. S14).

Despite the prevalence of dislocations, nacre’s self-correcting growth processes preserve mesoscale translational symmetry via substantial thickness changes (∼28%) at the topological singularity. Fig. 3 F, G, K, and L show tablet thicknesses at and around screw dislocations viewed in cross-section parallel or perpendicular to the slip plane. The singularity causes tablets to become thinner to accommodate an additional layer without disrupting the long-range order of the mesoscale periodicity. Within just a couple layers, above or below the dislocation, tablet thicknesses at and around screw dislocations viewed in cross-section parallel or perpendicular to the slip plane. The singularity causes tablets to become thinner to accommodate an additional layer without disrupting the long-range order of the mesoscale periodicity. Within just a couple layers, above or below the dislocation, nacre visually returns to uniform, periodic growth.

**Nacre Growth Fluctuations Follow 1/f Behavior.** To understand the stochastic growth processes over longer lengths and time, nacre tablet thicknesses are analyzed across the entire pearl (2.5-mm diameter) at high resolution (∼3 nm) (Fig. 4A). This pearl contained 2,615 tablet layers deposited over 548 d, giving a mean growth rate of 4 to 5 tablets (1.4 to 1.7 μm) per day—in this regard, distances may also be considered in terms of time (Methods). Every tablet of nacre is measured from a mosaic of BSE-SEM images, and a profile of their thicknesses is plotted in chronological order in Fig. 4C. Initial thicknesses fluctuate substantially but reduce quickly within ∼10 layers (Fig. 4C, yellow box), as shown in the thickness map (Fig. 4D). Notably, sawtooth-shaped bursts in systematic thickness variation are visible (Fig. 4C, red box) and repeat aperiodically throughout the entire growth (Fig. 4E). The stochastic processes of nacre growth are revealed in a power spectral density (PSD [i.e., squared magnitude of the Fourier transform]) of the thickness profile (Fig. 4B). The spectral density shows two distinct regimes: power-law decay across a low- to mid-frequency regime corresponding to distances longer than ∼25 tablets and a high-frequency regime where the density slightly increases.

The spectral density of a pearl’s tablet thicknesses follows power-law decay across low to mid frequencies, colloquially called 1/f noise, suggesting nacre growth has correlations that extend over a wide range of time scales with cooperative effects linked to environmental changes. In stochastic Markov processes, the distribution of future events are determined by recent past events (55, 56), and even simple Markov processes can exhibit a spectral density following $f^{-2}$ where $1 < \alpha < 2$. This behavior characterizes a wide range of phenomena from fluctuations in river flow to heartbeats or economic markets (57, 58). The spectral density of nacre follows $f^{-1.5}$ across low to mid frequencies (≥25 tablets)—here, $f = 1 / (\text{number of tablets})$. In real-space, this corresponds to the sawtooth-shaped bursts in the thickness profile of Fig. 4C. Such aperiodic fluctuations can be associated with the $f^{-1.5}$ spectral density, and the behavior can be explained by the additive effect of one or several Markov processes representing the external effect of environmental and physiological factors. The coefficient $\alpha$ of power-law decay is estimated by truncating the PSD and applying a square fit for log–log coordinates (Methods). The observed power-law decay, $f^{-1.5}$, suggests that nacre’s growth is influenced by a set of stochastic processes with different characteristic time scales. This makes sense: we expect growth to depend on many external factors in the surrounding habitat of the mollusk shell (13, 14, 59). Multiple clocks in nature govern these living animals, such as the 30-d circannual cycle (60, 61), the day–night circadian cycle (62, 63), tidal changes (64), hibernation in the winter (65, 66), and aestivation in the summer (67, 68). This fluctuating marine environment and the natural time scales of the mollusk are interconnected influences of the nonequilibrium thermodynamical conditions (69–71) of nacre growth and likely explains the observed power-law behavior.

At high frequencies (<25 tablets), there is a flattening of the 1/f noise and a slight increase in the power spectrum due to
corrective growth processes, whereby the thickness of each tablet is antirelated with the thickness of the previous layer. The opposing fluctuations in subsequent layers (shown in Fig. 2C) amplifies high-frequency variations in a manner consistent with a negative autoregressive process (72–74) and causes the slightly increasing profile of the spectral density at high frequencies (Fig. 4B). Mechanistically, the source of this correlational correction is not clear—it may be explained by a partial viscosity of the CaCO3 or the precursor phase upon deposition under hydrostatic pressure. The corrective behavior likely emanates when nacre tablets form simultaneously within multiple periodically spaced continuous organic sheaths—allowing not-yet-calcified layers to influence the thicknesses of adjacent layers. Physiological responses may also play a role.

In nacre, we see a fundamental tradeoff between achieving mesoscale translational symmetry and minimizing tablet thickness variation. Corrective growth occurs to counterbalance disorder and the natural tendency toward paracrystallinity, thereby enabling surprising and remarkable mesoscale periodic ordering. However, this corrective growth process amplifies variation, wherein perturbations in one layer causes an opposite perturbation in subsequent layers.

**Discussion**

Pearls are iconic mesoscale periodic structures grown by mollusks with no external templates along the direction of growth. Nanoparticles assemble within an organic matrix with increasing density until a massive, bulk aragonite forms and later followed by nacre deposition. Tablets are then grown with self-correcting mechanisms that allow mesoscale translational symmetry to accommodate tablet variation and topological defects. Mollusks strike a balance between preserving translational symmetry and reducing thickness variation by creating a paracrystal with medium-range order (∼5.5-μm correlation length). This balance allows pearls to attenuate the initial disorder during early formation and maintain order throughout a changing external environment. Over longer-length scales (>25 tablets), nacre growth is mediated by external stochastic processes and exhibits a “1/f” behavior that belongs to a universal class of vast and disparate phenomena. Quantifying the mesoscale periodicity and paracrystallinity of nacreous pearls reveals underlying processes of formation and illustrates metrics for assessing assemblies of synthetic nanocrystalline materials that seek to mimic nacre’s hierarchical design.

**Methods**

**Specimen.** Specimens of nacreous non-bead-cultured Akoya “keshi” pearls produced by the *P. i. fucata* pearl-oyster were collected at the Broken Bay Pearl Farm (Pearls of Australia Pty Ltd) located on the east coast of Australia. The term keshi pearls is used today to describe beadless pearls that were uninfluenced by nucleation. Mollusks strike a balance between preserving translational symmetry to accommodate tablet variation and topological defects. Mollusks strike a balance between preserving translational symmetry and reducing thickness variation by creating a paracrystal with medium-range order (∼5.5-μm correlation length). This balance allows pearls to attenuate the initial disorder during early formation and maintain order throughout a changing external environment. Over longer-length scales (>25 tablets), nacre growth is mediated by external stochastic processes and exhibits a “1/f” behavior that belongs to a universal class of vast and disparate phenomena. Quantifying the mesoscale periodicity and paracrystallinity of nacreous pearls reveals underlying processes of formation and illustrates metrics for assessing assemblies of synthetic nanocrystalline materials that seek to mimic nacre’s hierarchical design.

**Wedge-Polishing Preparation.** Cross-sections were cut from the white pearls measuring 3 to 5 mm in diameter using a diamond wire saw. The cross-sections for SEM were prepared by mechanical wedge polishing (11, 24), which provides large-area, electron-transparent specimens with structural stability. The lapping process was performed using an Allied MultiPrep polishing system and a series of micrometer-sized diamonds embedded in polishing film (30 μm down to 0.1 μm). The lapping sequence was dictated by the “trinity of damage,” where a damaging layer is assumed to be less than three times as thick as the size of the grit. Therefore, with each subsequent lapping film, this damage layer was removed. To prevent CaCO3 etching during preparation, water-based lubricants were eliminated—instead, Allied alcohol-based “blue” lubricant was used.

**Electron Microscopy.** SEM images were recorded with a 2-contrast back-scattered electron detector of a JEOL 7500 system (10 kV) with an energy-dispersive X-ray spectroscopy detector. High-angle ADF-STEM was performed using a JEOL 31000RS microscope with Cs aberration-corrected STEM (300 keV, 15 mrad) and cold field emission gun. An ADF detector with 120- to 150-mm camera lengths and a detector angle of 59 – 74 (inner) to 354 – 443 mrad (outer) were used to produce Z-contrast images, where grey-scale intensity is sensitive to the atomic number in the specimen’s matrix. Column pressure in the STEM column at the specimen was ~1 × 10⁻⁷ torr.

**Low-dose methods, beam shuttering, and examination of regions exposed to the beam were used to separate electron beam irradiation from intrinsic phenomena (11, 24). For STEM measurements, with typical fields of view from ~500 nm to 10 μm, the electron dose was typically from ~0.4 to 90 e⁻ A⁻¹s⁻¹, and dose rates ranging from around 0.1 to 2.7 e⁻ A⁻¹ s⁻¹. The material was structurally preserved during imaging. However for the same imaging conditions at higher magnifications (e.g., 20-nm field of view), the radiation dose increases to ~9 × 10⁶ e⁻ A⁻¹ s⁻¹ and dose rate to ~3 × 10⁸ e⁻ A⁻¹ s⁻¹, which causes the material to show electron irradiation damage localized to the small field of view. Thus, larger fields of view are preferred to minimize dose and provide a large area of observation. Atomic-resolution STEM requires a small field of view with an on-axis region of interest.

**Raman Spectroscopy.** Confocal Raman spectroscopy (SI Appendix, Figs. S7 and S8) was performed on the cross-sections of pearls using a Tescan Raman imaging and scanning electron microscope equipped with a Witec confocal Raman microscope. Raman spectra in the spectral range (100 to 3,700 cm⁻¹) were recorded with a charge-coupled device camera using a solid-state 532-nm laser as the excitation source (spatial grating: 600 g/mm, laser power: 10 – 15 mW, optical lens: 100x). Each Raman spectrum was acquired typically for 0.5 ~ 1 s, and with ~100 of scans to minimize noise effects. The parameters for excitation power, acquisitions and accumulations are selected to prevent the sample from laser burn. Spectral reproducibility was confirmed by taking several spot analyses.

**Data Analysis.** The pair-correlation function describes the probability distributions of NN tablets in space about a known tablet position (50). The one-dimensional pair-correlation function is extracted from raw BSE-SEM images with high resolution (~3 nm). Mechanical scratches are removed using the image filtering (78). Interlamellar sheaths are identified by finding local minima in BSE intensity line profiles taken perpendicular to the nacre layers. The resulting interfacial positions are then used to calculate the thickness of nacre layers by measuring the distance between positions. The N⁰ tablets thickness is calculated as the difference between the N⁰ and N⁰-1 interfacial positions. This is done throughout all positions and layers across the entire pearl cross-section to generate a thickness map visualized as a color overlay atop an SEM image. A one-dimensional (1D) histogram of tablet thickness was then calculated and normalized by a factor of 1/√n to obtain a discrete version of the pair-correlation function, where n is the number of samples and bw is the histogram bin width. A sum of normalized gaussians is fitted to the histogram peaks, and the SDs are plotted versus peak order. Occasional, narrow two-tablet thicknesses appear in the thickness maps due to lack of contrast between layers, which may be due to topological defects and sometimes large asperities or mineral “Checa bridges” between tablets (20)—these minority regions were not included in generating the pair-correlation function.

**Correlation between the widths of adjacent tablets is calculated by measuring the deviation in tablet width from the local mean tablet width.** The local mean is calculated from the list of all sampled tablet widths using a rolling average with a width of three tablets. Thus, the deviation in thickness of the nth tablet is given as widthₜₙ – (widthₜₙ₋₁ + widthₜₙ₊₁)/2. Deviations in thickness of pairs of adjacent tablets are plotted against each other and the correlation coefficient is estimated by linear least squares fitting. Correlation length is defined as the length where the pair-correlation envelope disappears. It becomes negligible when the magnitude of the following peak is below 10% of the first peak (i.e., pₙ < 0.1). Translational symmetry is lost by 16 layers (i.e., 5.5 μm).

**Autocorrelation of tablet thicknesses can be interpreted by PSD, which is generated in the frequency domain by squaring a fast Fourier transform of thickness profile.** The tablet thickness is measured from BSE-SEM images using the peak-detection algorithm described above and is plotted in chronological
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