growth front, the silica-coated carbonate nanoparticles self assemble and aggregate free of the symmetry restrictions normally imposed by the carbonate’s crystal lattice. Perhaps not surprisingly, though, the large-scale form of the biomorph depends on the overall pH of the solution, which changes over a few hours due to CO₂ dissolution, as well as on local pH fluctuations.

Using video microscopy, the team identifies specific stages of growth. Initially, the carbonate crystal grows as a fragmenting, dendritic stalk whose head resembles a cauliflower, as in figure 1. Then, once the pH drops enough for cyclic coprecipitation to be established (within a day, typically), the biomorph grows through pulses of nucleation events to form the carbonate nanoparticles. At that stage, for reasons unexplained by the team’s mechanism, two-dimensional sheets emerge from the 3D cauliflower. Each sheet, consisting of partially aligned nanoparticles, grows radially outward, often to hundreds of square microns, until it curls up at points along its edge like a scroll. Those curls carve out smoothly curved borders that can become shaped like cardiods or interact to spawn filaments like those in figure 2.

Werner Kunz and Matthias Kellermeier (both at the University of Regensburg in Germany) speculate that interfacial tension between the liquid and 2D surface may account for the curling. Michael Marder (University of Texas at Austin) postulates another reason: “Once 2D sheets form, it’s actually hard for them to grow in a way that won’t curl or buckle for exactly the same reason that leaves tend to do so—spontaneous symmetry breaking. Simply inserting material or stretching the sheet changes its internal metric (the equilibrium lengths between things), which induces nonzero Gaussian curvature.” (See the article by Marder, Robert Deegan, and Eran Sharon in PHYSICS TODAY, February 2007, page 33.)

Garcia-Ruiz’s team presents a phenomenological model that describes a variety of emergent biomorph shapes using just four parameters: the radial velocity of the advancing sheet; the azimuthal velocity, or speed of the curl’s leading edge; the handedness of the curl; and its relative height. The myriad helical filaments in figure 2, captured by polarized optical microscopy, exemplify one case. Each is extruded from the cusp formed when any two curls moving along the edge of the wide (and out-of-focus) parent sheet approach each other with the same handedness and with nearly equal azimuthal velocities. Together, those conditions lock the curled segments of the sheet into a coupled growth process that causes a filament to twist as it lengthens, as sketched in the side-view inset.

Although mysteries still abound, Garcia-Ruiz argues that the research could provide insight on how biominerals like sea shells, bone, and teeth form. Traditionally, they’re thought to develop from a protein, surfactant, or other biomolecule that, like a scaffold, imprints its structure on the subsequent growth of inorganic minerals. But armed with a plausible mechanism for dynamic self-organization in biomorphs, one can now ask whether that static view of biomineralization is really correct. Whatever the answer, says Kunz, researchers are sure to start searching for novel materials that form out of similar chemically coupled pairs.

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References

Room-temperature magnetic refrigeration. To reject heat or to generate “cold,” refrigerators typically rely on the vapor-compression cycle of fluids such as the chlorofluorocarbon Freon. Now emerging is a new class of solid-state materials known as magnetocalorics; under an applied magnetic field or with changing temperature, they experience a large and sharp magnetic entropy change with the first-order paramagnetic–ferromagnetic phase transition. That entropy change, known as the magnetocaloric effect, makes the material an effective heat sink. In fact, such materials have already been exploited for cryogenic cooling in research settings. Recently discovered magnetocaloric metal alloys containing either gadolinium or arsenic have shown large MCE values at ambient temperatures; however, the high cost of the former and the undesirable toxicity of the latter present challenges to their commercial deployment.

Inspired by scientists at the Beijing University of Technology who replaced arsenic with the relatively benign germanium, an international collaboration has now measured the germanium-containing material’s MCE under varying temperatures and magnetic fields. The team also conducted neutron-scattering studies at NIST to examine the material’s crystal structure. Diffraction measurements revealed that the ferromagnetic structure, when compared with the paramagnetic phase, was noticeably contracted, confirming that the phase transition is driven by structural changes. The team also discovered that in the 250- to 270-K range, a 5-tesla magnetic field induced an MCE value in the germanium compound that is considerably higher than the arsenic-containing magnetocalorics.

The researchers say the MCE can be pushed even higher with further improvements in sample purity. (D. Liu et al., Phys. Rev. B 79, 014435, 2009.) —JNAM

Manipulating magnet-coated bubbles. Gas bubbles in a liquid driven by acoustic waves can be used in a variety of contexts—for example, as contrast agents for ultrasound imaging, as a delivery system for therapeutic drugs, as catalysts for sonochemical reactions, and as scrubbers of surfaces. The bubbles’ compressibility, which allows their volume to oscillate in response to the varying sound pressure, accounts for the wide applicability. Unfortunately, spatial gradients in the pressure field can make it nearly impossible to control the bubbles’ oscillations and position simultaneously. Researchers at Nanyang Technological University in Singapore have now found a simple recipe to dress the bubbles in a 1-μm-thick coat of magnetite nanoparticles. The coat, which self assembles in solution, stabilizes the bubbles (typically 40–350 μm in diameter) without sacrificing their compressibility; they remain intact for more than 6 months in a light-tight drawer and their position in solution can be controlled with a simple household magnet. The figure here illustrates an example: The oscillation of a single 130-μm-wide bubble, subject to sound at a frequency close to its resonant frequency, sets up eddies (signified by arrows) in the surrounding fluid. After the acoustic field is turned off, a permanent magnet nudges the bubble to the right. The authors expect magnetic bubbles to serve as remotely controlled microfluidic mixers and pumps, and, more generally, as tools to test fundamental fluid mechanics concepts. (X. Zhao, P. A. Quinto-Su, C.-D. Ohl, Phys. Rev. Lett. 102, 024501, 2009.) —RMW

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