

RESEARCH HIGHLIGHTS

Blockbuster film*Phys. Rev. E* 74, 051602 (2006)

Could you make a soap film twice the size of Tiananmen Square without it popping? In space it might be possible, say Rui Zheng and Thomas Witten of the University of Chicago, Illinois.

They suggest using, instead of soapy water, a commercial pump oil stabilized by surfactants, which would evaporate slowly enough to create a film that is 1 kilometre across by 1 micrometre thick and stable for at least a year. In Earth orbit, the film would absorb enough sunlight not to freeze. And with the right surfactant, the film shouldn't rupture spontaneously; meteoroids that punch holes, however, are more problematic.

Why go to all the bother? The film would be a great laboratory for investigating two-dimensional flow and turbulence, the authors argue — important in geophysics and astrophysics but hard to study in the lab.



N. JORGENSEN/REX FEATURES

CELL BIOLOGY**DNA is pulled into line***Genes Dev.* 20, 3269–3282 (2006)

When cells divide, they replicate their DNA so that each daughter cell can receive a full chromosomal complement. Eukaryotes use specialized motor proteins to pull chromosomes into each new cell, but how bacteria divvy up genetic material has remained a mystery.

Michael Fogel and Matthew Waldor of Tufts University in Boston, Massachusetts, now show how the cholera-causing bacterium, *Vibrio cholerae*, distributes its chromosomes when dividing. A protein called ParAI stretches across the cell, grabs a chromosome, and retracts, pulling the DNA along with it.

The finding provides the first example of pulling forces that distribute bacterial DNA, and suggests a possible evolutionary link between chromosome segregation in bacteria and in eukaryotes.

NANOTECHNOLOGY**Spare the rod***J. Am. Chem. Soc.* doi:10.1021/ja064535p (2006)

Nanorods of an organic material increase in length by an average of 15% when illuminated with ultraviolet light, report Christopher Bardeen and his colleagues at the University of California, Riverside. Surprisingly, randomly shaped crystals of the same material shatter under identical treatment.

Ultraviolet light induces pairs of molecules in the nanorods to react with each other, forming a product that takes up more molecular space than the starting material. The resulting crystal strain is more easily released at the surface of the rod than within it. The high surface-to-volume ratio of nanorods, the authors propose, allows strain to be released non-destructively by the extending rods.

This effect could eventually be harnessed to trigger nanoscale motion from light energy, the researchers suggest.

IMMUNOLOGY**Body invaders***J. Clin. Invest.* 116, 3258–3265

(2006)

In the disease type-1 diabetes, the body's immune system attacks many different proteins. Previous research had shown that the first victim is proinsulin. But until now, it was not clear whether immune responses spread from there to other proteins, or whether those others were attacked simultaneously.

Thomas Kay from St Vincent's Institute in Victoria, Australia, and his colleagues used genetically modified mice to study two proteins — proinsulin, and another called IGRP — that trigger inflammatory responses. Mice engineered to not respond to proinsulin also had no response to IGRP, and did not get sick. But those designed not to respond to

IGRP did recognize proinsulin and developed diabetes — suggesting that the response to proinsulin is 'upstream' of that to IGRP.

Proinsulin triggers the first and key response, the authors suggest, and thus should be the main target for diabetes treatments early in the course of the disease.

SUPRAMOLECULAR CHEMISTRY**Tying the knot***Angew. Chem. Int. Edn* doi:10.1002/

anie.200603521 (2006)

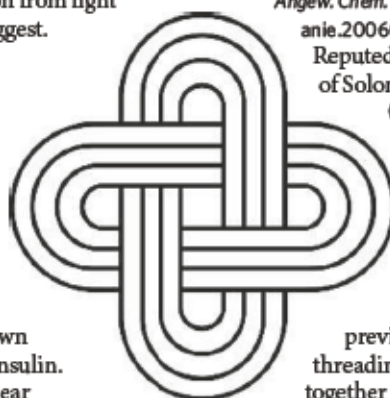
Reputed to encode all the wisdom of Solomon, King Solomon's Knot (pictured left) is found in sacred art from Africa to Celtic Britain. An interweaving of two chain links, it is not strictly speaking a knot, but a link.

Chemists have

previously figured out ways of threading two molecular hoops together to make Solomon's link, but the method now described by Fraser

Stoddart of the University of California, Los Angeles, and his co-workers has a new twist. They assemble the link from 12 distinct components, using metal ions to template the threading of the organic components of the two rings.

The Solomon link is just one of the possible interwoven products, but its formation can be amplified during crystallization, promoting this molecule over the others.



To flee or not to flee

If not exposed to predators regularly, animals seem to lose the physiological triggers that could save them from being eaten, a new study suggests.

For millions of years, until dogs and cats showed up a century and a half ago, marine iguanas (pictured right) lived in the Galapagos Islands without major land predators. In an experiment, researchers chased iguanas that still live in isolated areas; the animals, they found, did not generate the stress hormone corticosterone. But iguanas living near a town, and presumably more used to being attacked by predators, did produce the hormone.

The iguanas were fast learners, however. Even the naive animals quickly started producing the hormone after being chased a couple of times, says a team led by Thomas Rödl of Princeton University, New Jersey.

Same key, different lock

Enzymes are so good at chemical reactions that they are sometimes too good to be useful when making new compounds. The enzymes are too specific, and always carry out the same reaction. One way around this is to engineer or evolve the enzymes to do what's wanted of them. Another solution, highlighted by Stephen Withers at the University of British Columbia in Vancouver, Canada, and his group, is to instead tweak the substrate, or landing pad for the enzyme, through a temporary modification.



When joining sugars together, adding an extra molecular group to one of them allows enzymes called glycosyltransferases to link them up in different configurations. This relatively simple strategy can produce a range of strikingly different products. Synthetic chemists, take note.

A one-two punch for cancer

Adding a genetically modified bacterium to chemotherapy drugs fights cancer better than the drugs alone, new work suggests.

Getting cancer treatments to kill cancerous, but not healthy, cells is a major problem for drug delivery. Now, a team led by Shibin Zhou and Bert Vogelstein at the Johns Hopkins Kimmel Comprehensive Cancer Center in Baltimore, Maryland, says that adding the bacterium *Clostridium novyi-NT* can help.

The combination treatment caused both large and small tumours in mice to shrink. The bacterium may draw the drug more selectively to the cancer cells, helping it to better penetrate and destroy the tumour.

Drilling deep

The first-ever deep-drilling programme into California's infamous San Andreas Fault is starting to yield information about the mineralogy of the rocks there.

John Solum of the US Geological Survey in Menlo Park, California, and his colleagues have identified several distinct mineral assemblages, down to nearly the bottom of their 4,000-metre-deep hole.

In particular, the mineral serpentine — sometimes seen along the San Andreas at the surface — appeared in the layer where the fault was actively deforming the rocks. The presence of serpentine, some think, could help explain certain details of how the fault behaves.

Coring at the San Andreas Fault Observatory at Depth, a US government project, begins again next summer.

Liquid-controlled light

The passage of light through a layer of silicon can be controlled using liquid droplets less than 1 femtolitre in volume.

Francesca Intonti of the European Laboratory for Non-Linear Spectroscopy in Firenze, Italy, and her colleagues placed drops of water into selected pores in a silicon wafer. Light moving through the silicon was then constrained to the S-shaped path marked out by the drops.

The technique could provide a new method for constructing analogues of electronic semiconductor devices, such as optical switches.

JOURNAL CLUB

The molecular dance of a protein allows a chemist's secret wish to come true.

One fascinating aspect of molecular function is the way information propagates between parts of a molecule that can be many tens of angstroms apart.

Our understanding of how proteins do this, a process termed allostery, emerged from Max

Perutz's pioneering studies of oxygen-carrying haemoglobin. Three-dimensional images show that when a ligand binds to part of the molecule, a discrete set of structural changes take place at distinct sites. This, in turn, influences the ease with which subsequent ligands bind.

Nature has chosen this model in designing many allosteric proteins. However, as a practising nuclear magnetic resonance (NMR) spectroscopist with a strong interest in protein dynamics, I was secretly hoping she might design

proteins in which information is communicated through changes in the dynamics between distal sites, with little or no change in overall structure. Moreover, I was rooting for NMR to play a major role in characterizing such a system.

How exciting it was, therefore, to read that Charalampous Kalodimos and his co-workers recently found such a case by studying the motional properties of a protein in different ligated states (N. Popovych *et al.* *Nature Struct. Mol. Biol.* 13, 831; 2006). Using NMR spectroscopy, the team quantified

protein dynamics for a wide range of timescales. Remarkably, ligand binding at one site is linked to changes in motion far removed, over the complete set of timescales, while a corresponding propagation of structural changes does not occur.

The work of Popovych *et al.* provides a striking example of the importance of protein dynamics to information transfer. I eagerly await the discovery of more molecular dances and of how they, too, will relate to biological function.