# **RESEARCH HIGHLIGHTS**

stabilize the LOV-Cdc42 interaction with additional hydrogen bonds in the caged, 'dark' state, allowing the researchers to make a PA-Cdc42. This suggests that on the basis of a structural understanding of the steric block for the protein family, the interface can be optimized for each individual protein to create other caged GTPases.

Hahn hopes that the LOV domain has an ideal structure to become a general caging tool, but he notes that the tagging approach will need to be worked out for each individual protein family. "The GTPases have a similar enough structure that we could come up with a general solution," he says. "But if you now try to do something with a different shape, you are going to have to ask how you put the yo-yo on a string,... but I hope that there will be broadly applicable solutions for each family." His group is now working with other protein families to see how general the approach can be.

And as more such caged proteins are made, tools will be needed to keep track of them. In a companion paper (Machacek *et al.*, 2009), Hahn's group, along with Gaudenz Danuser's team at Scripps, used a computational multiplexing approach to delineate the spatiotemporal relationships between activities of the GTPases Rac1, RhoA and Cdc42 during cell protrusion.

As for PA-Rac1, because the protein controls cell motility, according to Hahn others are using it to make cells move around in animals. So more work is yet to come on many fronts.

Irene Kaganman

#### **RESEARCH PAPERS**

Machacek, M. *et al.* Coordination of Rho GTPase activities during cell protrusion. *Nature* **461**, 99–103 (2009).

Wu, Y.I. *et al.* A genetically encoded photoactivatable Rac controls the motility of living cells. *Nature* **461**, 104–108 (2009).

minutes—with a diverse array of protein-coding sequences, although the efficiency of folding varied depending on the protein in question. As a further demonstration of the utility of their method, the researchers monitored diffusion and interaction behavior of individual proteins directly in the cell-free translation mixture using fluorescence correlation spectroscopy.

Based on these initial experiments, Alexandrov is encouraged that his method offers a strong combination of efficiency and the capacity to ratchet up eukaryotic protein production to the liter scale at a fraction of the cost of existing commercial systems—a serious boon to scientists looking to generate crystals or produce antibodies. In contrast, there are also important advantages in 'going small', and the capacity for highly multiplexed protein production using the speciesindependent translational sequence leader, either with L. tarentolae extract or any other extract system, could enable powerful, high-throughput interaction analysis studies or even new approaches for tackling proteomic research. "You can find an organism, make an extract and back-translate its genome," says Alexandrov. "Craig Venter estimates a total pool of 20 million genes on the planet, and most of them are in microorganisms on which we have no handle-and so this could be a way of guickly generating expressed proteomes." **Michael Eisenstein** 

#### **RESEARCH PAPERS**

Mureev, S. *et al.* Species-independent translational leaders facilitate cell-free expression. *Nat. Biotechnol.* **27**, 747–752 (2009).

# **NEWS IN BRIEF**

## STEM CELLS

### Mice from iPSCs

Two groups show that mouse induced pluripotent stem cells (iPSCs) can generate fertile adult mice in tetraploid complementation assays, in which all tissues are derived from the pluripotent cells. Failure to achieve this so far has raised questions about the pluripotency of iPSCs, but these results show that full pluripotency via reprogramming is possible. The mice will be useful models to study the function of cells and tissues that are entirely iPSC-derived.

Boland, M.J. *et al. Nature* **461**, 91–94 (2009).

Zhao, X.Y. et al. Nature **461**, 86–90 (2009).

## PROTEOMICS

## Mammalian tethered catalysis

The tethered catalysis method, originally developed in yeast, is used to identify post-translational modification–specific binding proteins (PTMBPs). A substrate peptide is tethered to an enzyme that posttranslationally modifies the peptide, which then serves as bait for PTMBPs. The PTMBPs can be identified by affinity purification and mass spectrometry. Spektor and Rice now report an expansion of the tethered catalysis approach for application in mammalian cells. Spektor, T.M. & Rice, J.C. *Proc. Natl. Acad. Sci. USA* **106**, 14808–14813 (2009).

#### CHEMICAL TOOLS

## Cleavable linkers for affinity chromatography

Biotin-streptavidin-based affinity chromatography is widely used for identifying protein-protein and protein-ligand interactions. However, releasing the biotinylated target protein from the streptavidin support is a harsh and inefficient process. Park *et al.* now report an acylhydrazone-based linker that can be incorporated into a biotin tag, and which is cleaved under mildly acidic conditions. This allows for a clean release of the target protein without requiring denaturation, facilitating downstream analysis. Park, K.D. *et al. Chem. Biol.* **16**, 763–772 (2009).

### GENOMICS

## Predicting copy-number variation

Though duplicated regions of the human genome are the culprits of various conditions ranging from color-blindness to lupus, these regions have been very difficult to study. Alkan *et al.* describe an algorithm called mrFAST, which can predict absolute copy-number variation in duplicated genome segments from next-generation sequencing data. By examining three human genomes, Alkan *et al.* estimated that 73–87 genes vary in copy number between individuals.

Alkan, C. et al. Nat. Genet. advance online publication (30 August 2009).

#### DRUG DISCOVERY

#### Mapping drug space

Drugs that target pathogenic bacteria or cancer cells can potentially also interfere with human enzyme function. Thus, an understanding of how drugs can affect metabolism is very important. Adams *et al.* present an online resource of interactive maps of potential drug action in metabolism, which compares the chemical structural similarities of drugs and human metabolites. The resource contains information on 246 drug classes and 385 organisms; such maps could be used to design more effective drugs. Adams, J.C. *et al. PLoS Comput. Biol.* **5**, e1000474.