sive promoter and removes any residual target-gene activity. To turn the switch on, a drug is added that inhibits expression of the Lac inhibitor, thus allowing expression of the target gene and the Tet repressor, which is needed to shut off shRNA expression from the tet-responsive promoter.

Collins and Dean showed that this intricate system relying on crosstalk between the different modules does indeed tightly regulate gene expression in mammalian cells. They used diphtheria toxin to prove that in the off state there was no detectable gene expression; otherwise, the cells—sensitive to even small amounts of the toxin—would not have survived. The cells only died once the researchers flipped the switch on, resulting in the expression of the diphtheria toxin.

Other useful features of this genetic circuit are that it can be used with any gene of interest and expression can be fine-tuned as desired.

Collins anticipates applications of the system in functional genomics, as well as for the study of normal development or disease onset and progression. He says, "Starting from a very tight off state you can explore phenotypic responses to titrated levels of gene expression, to establish threshold responses."

And of course it will be useful in synthetic biology. A well-characterized toggle switch can be incorporated into much more complicated circuit designs, which are needed to recapitulate pathways in mammalian cells.

Nicole Rusk

RESEARCH PAPERS

Deans, T.L. et al. A tunable genetic switch based on RNAi and repressor proteins for regulating gene expression in mammalian cells. Cell 130, 363-372 (2007).

as well as the speed of incorporation, which provides insight into metabolic dynamics.

In a recent paper, Rabinowitz and his graduate student Jie Yuan took the KFP method a step further "to be able to figure out when you have branch points in metabolism, and which branch is being used," explains Rabinowitz. More specifically, they showed that KFP can distinguish between metabolites resulting from macromolecular decomposition versus those that are synthesized de novo (Yuan and Rabinowitz, 2007). Using the rapid isotope switching method with [15N]ammonia as the label source, they added a carbon starvation step by transferring the cells on the filter from medium containing glucose to medium lacking glucose (or vice versa). De novo-synthesized metabolites incorporate ¹⁵N, whereas those produced by macromolecular decomposition remain unlabeled. The researchers thus observed that under glucose starvation conditions cells switch from producing amino acids by synthesis to producing them by protein degradation.

Whether in NMR or mass spectrometry, systems biology or medical diagnostics, isotope labeling provides researchers with a powerful way to trace small-molecule metabolites.

Allison Doerr

RESEARCH PAPERS

Shanaiah, N. et al. Class selection of amino acid metabolites in body fluids using chemical derivatization and their enhanced ¹³C NMR. Proc. Natl. Acad. Sci. USA 104, 11540-11544 (2007).

Yuan, J. et al. Kinetic flux profiling of nitrogen assimilation in Escherichia coli. Nat. Chem. Biol. 2, 529-530 (2006).

Yuan, J. & Rabinowitz, J.D. Differentiating metabolites formed from de novo synthesis versus macromolecular decomposition. J. Am. Chem. Soc. 129, 9294-9295 (2007).

NEWS IN BRIEF

RNA INTERFERENCE

MicroRNA matchmaking

Computational microRNA target prediction programs usually yield a large number of targets for each microRNA, and deciding which to choose for validation is difficult. Grimson et al. describe five new features that determine the pairing of a microRNA to its target. These features are incorporated into their target-discovery algorithm, TargetScan, and will provide a more stringent selection for microRNA-target pairs. Grimson, A. et al. Mol. Cell 27, 91-105 (2007).

MICROSCOPY

Target-locking microscopy

Several microscope systems can target-lock an object and image it as it moves randomly in three dimensions. But these methods generally treat the object as a point, which makes the tracking of large complex objects such as cells difficult. Lu et al. describe target-locking acquisition with realtime confocal (TARC) microscopy that can follow a cell moving in three dimensions even as it changes shape, size and orientation. Lu, P.J. et al. Optics Express 15, 8702-8712 (2007).

GENOMICS

Mapping an abundance of SNPs

By resequencing the genomes of 15 different laboratory mouse strains, including 11 classical and 4 wild-derived ones, Frazer et al. identified more than 8 million unique singlenucleotide polymorphisms (SNPs) across the mouse genome with oligonucleotide arrays. They used the data to generate an ancestral haplotype map, available online (http://mouse. perlegen.com).

Frazer, K.A. et al. Nature, published online 29 July 2007.

PURIFICATION AND SEPARATION

Assaying sisterly cohesion

Sister chromatid cohesion is an essential process during cell division, requiring a multiprotein complex called cohesin. Ivanov and Nasmyth developed an in vitro system based on sucrose gradient centrifugation and gel electrophoresis, which allowed them to directly show that the cohesin ring in yeast affects sister chromatid cohesion by trapping the sisters within its ring. Ivanov, D. & Nasmyth, K. Mol. Cell 27, 300-310 (2007).

BIOSENSORS

Artificial tongues made from synthetic pores

There has been considerable interest in developing 'electronic tongues' for understanding the biology of taste and for developing sensing applications. Litvinchuk et al. report a universal stimulus-responsive pore to detect flavor molecules by using the concept of 'reactive amplifiers': molecules that react with analytes to enhance their pore-blocking ability and thus amplify the pore response.

Litvinchuk, S. et al. Nat. Mater. 6, 576-580 (2007).