DEVELOPMENTAL GENETICS

Micromasters

New work from Javier Palatnik and colleagues provides the first direct evidence that microRNAs (miRNAs) control aspects of plant development by guiding the cleavage of mRNAs.

Like small interfering RNAs (siRNAs), which are the effectors of RNA interference, miRNAs can bind to complementary mRNAs and silence the genes that encode them. Despite evidence that they can silence plant genes, previously there was no support for any specific biological function for miRNA silencing.

In an Arabidopsis developmental mutant (jaw-D) that has uneven leaf shape and curvature in adult plants, Palatnik and colleagues identified five TCP genes that were downregulated, which were closely related to a snapdragon gene (CIN) that is involved in leaf morphogenesis. In all five TCP genes, the authors found a motif that was partially complementary to a previously identified plant miRNA and a sequence at the JAW locus.

By hybridizing a probe that was based on the JAW sequence to a low molecular-mass RNA blot, the authors showed that it encoded a 20-nucleotide miRNA (miR-JAW). To try to understand the interaction between miR-JAW and the TCP genes, the authors engineered miRNAresistant forms of TCP4. Constructs with mutant TCP4 produced stable transcripts that were unaffected by the

co-injection of a JAW construct into a heterologous host, unlike wild-type TCP4 constructs, which led to an accumulation of cleavage products.

The same mutations introduced into TCP4 in situ resulted in a range of developmental defects, most of which led to growth arrest at the seedling stage. This clearly showed that miRNA-guided cleavage of TCP4 is essential for normal plant development.

Neatly completing the circle, the authors then showed that miRNAresistant TCP2 could rescue the jaw-D phenotype. So, the production of TCP2 transcripts that are resistant to miRNA-mediated degradation might correct aberrant phenotypes that are associated with lower expression levels of the gene.

As miRNA target sequences were found in TCP genes from plants with a range of leaf forms, including lotus and maize, it seems that these micromasters might have a ubiquitous role in controlling leaf development. Perhaps the time is now ripe for a wider search for such motifs to help identify other developmental pathways that could be influenced by miRNAs.

Nick Campbell

References and links

ORIGINAL RESEARCH PAPER Palatnik, J. et al. Control of leaf morphogenesis by microRNAs. Nature 20 August 2003 (10.1038/nature.01958) FURTHER READING McManus, M. T. &

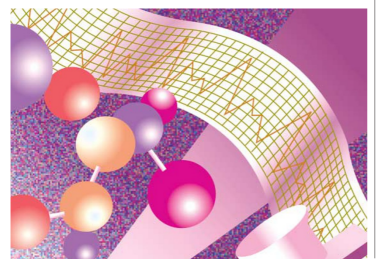
Sharp, P. A. Gene silencing in mammals by small interfering RNAs. Nature Rev. Genet. 3, 737-747 (2002)

WEB SITES

Detlef Weigel's laboratory: http://weigelworld.org

Arabidopsis small RNA project in James

Carrington's laboratory: http://cgrb.orst.edu/smallRNA/index.html



IN BRIEF

DEVELOPMENTAL BIOLOGY

Conserved role of nanos proteins in germ cell development.

Tsuda, M. et al. Science 301, 1239-1241 (2003)

In Drosophila, nanos encodes an RNA-binding protein that is required maternally for the migration of the primordial germ cells into the gonad. Now, two mouse nanos homologues, nanos2 and nanos3, have been cloned. The knockout phenotypes of these two genes — lack of male germ cells in nanos2 -/- mice and lack of germ cells in both sexes in nanos3-/- animals — indicate that despite apparent differences, the mechanism of germ-cell specification is conserved between vertebrates and invertebrates.

DEVELOPMENTAL BIOLOGY

GATA-3: an unexpected regulator of cell lineage determination in skin.

Kaufman, C. K. et al. Genes Dev. 17, 2108-2122 (2003)

The GATA-3 transcription factor is known to be required in lymphocyte differentiation. A microarray screen showed that GATA-3 is also expressed in the inner root sheath (IRS) cells — those that guide the developing hair shaft to the skin surface — early during hair morphogenesis. This evidence, together with the absence of the IRS in GATA-3-null mice and the aberrant structure of the mutant hair, points to a new function for GATA-3 in the differentiation of epidermal stem cells.

EVOLUTION

Parasite selection for immunogenetic optimality.

Wegner, K. M. et al. Science 301, 1343 (2003)

Theoretical models predict that the highly polymorphic genes of the vertebrate major histocompatibility (MHC) complex, which help to fight parasite infection, should exist at an intermediate level of diversity: too little diversity would impair pathogen recognition whereas too much would increase the likelihood of reacting against self-peptides. Wegner and colleagues have now tested the theory by infecting stickleback fish of varying levels of MHC diversity with several parasites. As predicted, fish with intermediate MHC diversity levels had the lowest infection rate.

FUNCTIONAL GENOMICS

The human transcriptome map reveals extremes in gene density, intron length, GC content, and repeat pattern for domains of highly and weakly expressed

Versteeg, R. et al. Genome Res. 13, 1998-2004 (2003)

The authors construct a new human transcriptome map to physically characterize clusters of highly expressed genes. They find that they are GC rich and contain many SINE but few LINE repeats. Genes that lie within these so-called ridges have shorter introns than genes that lie elsewhere. They also find evidence of clusters of weakly expressed genes, which they call antiridges.