NEUROGENESIS

Give it a REST!

The REST protein helps to prevent the premature activation of genes that are only expressed in mature neurons, and is now found to protect the genome of neural progenitor cells.

STEVEN M POLLARD AND MARIA ANGELES MARQUES-TORREJON

To construct the mammalian brain, the right neurons must be produced in the right place at the right time. For example, the outer layer of the brain, the cortex, consists of six layers of neurons and is built up one layer at a time. This is achieved by controlling how cells called apical progenitors become basal progenitors, which then specialize into new neurons (Gotz et al., 2002; Figure 1). If the apical progenitors become basal progenitors too soon, several developmental brain abnormalities can result, the brain may be too small (a condition known as microcephaly), or the layering of the cortex may be disrupted. Now, in eLife, Gail Mandel of the Oregon Health and Science University and co-workers – including Tamilla Nechiporuk as first author – report that a protein called REST has an unexpected role in protecting the genome of these progenitor cells (Nechiporuk et al., 2016).

The identity of a cell is determined to a large extent by which of its genes are transcribed. Therefore, when a new neuron first develops from a progenitor cell, a process of ‘transcriptional resetting’ must occur so that the genes that need to be expressed in mature neurons can be activated. The REST protein, first identified in 1995, is expressed in all cells except for mature neurons, and so researchers immediately suspected that it was involved in repressing neuronal genes (Schoenherr and Anderson, 1995; Chen et al., 1998; Bruce et al., 2004).

Support for this idea came from experiments that showed that REST binds to and represses genetic elements that are associated with many genes that are specific to mature neurons. As part of this repression, REST recruits a series of repressor complexes that alter how the gene is packed into a structure called chromatin. However, a previous study involving knockout mice who could not produce the REST protein failed to identify any significant abnormalities in the developing or adult brain (Gao et al., 2011; Aoki et al., 2012; Yang et al, 2012). The role of REST has therefore remained uncertain.

Nechiporuk et al. – who are based in the US and Germany – have now used a technique called conditional genetic ablation to explore the role of REST in neural progenitors. This revealed an unexpected requirement for REST in protecting the genome of the apical progenitors. Loss of REST induces DNA damage during the S phase of the cell cycle: this is the phase during which DNA is replicated. A consequence of this damage is the acquisition of chromosomal abnormalities in the apical progenitors.
This triggers cells to commit suicide – following the orders of a protein called p53 – and the result is microcephaly. Nechiporuk et al. also show that the combined loss of REST and p53 results in the formation of a highly aggressive brain tumour called a glioblastoma. REST therefore performs a dual role during brain development: it protects the progenitor cells from genetic catastrophe, and it silences neuronal gene expression until the time is right.

These findings came as a surprise, given the results of the previous REST knockout studies. However, in a series of elegant experiments Nechiporuk et al. showed that these earlier mouse models did not fully delete the REST coding sequences: the knockout mice still produced a C-terminal peptide that was able to recruit the repressor complexes that helped to silence certain genes. This is a valuable lesson for all researchers – knocking out a gene does not always result in a complete loss of function.

Why does the premature removal of the repressor complexes recruited by REST inflict widespread genomic damage? One possibility proposed by Nechiporuk et al. is that the associated loss of chromatin repression might lead to a subset of neuronal genes being incorrectly transcribed. Thus, REST seems to provide a ‘license’ for progenitors to transform into their final neuronal form by guarding the genome and preventing the premature transcription of genes specific to mature neurons. These new findings address a question that has received little attention to date – how are genome maintenance and transcriptional control coordinated as new neurons develop from progenitor cells?

As with all interesting discoveries, many new questions arise: how is REST protecting neuronal gene integrity during S-phase? How are the processes of cell cycle exit, chromatin repression, and the DNA replication timing coordinated during the birth of new neurons? New insights into how REST orchestrates gene regulation during the construction of the nervous system will clearly enhance our understanding of diseases such as microcephaly and brain cancer. It seems that for neural
progenitors, a little REST is what it takes to ensure you reach your full potential!

Steven M Pollard is at the MRC Centre for Regenerative Medicine, University of Edinburgh, Edinburgh, United Kingdom

Steven.pollard@ed.ac.uk

Maria Angeles Marques-Torrejon is at the MRC Centre for Regenerative Medicine, University of Edinburgh, Edinburgh, United Kingdom

Competing interests: The authors declare that no competing interests exist.

Published 08 January 2016

References

Aoki H, Hara A, Era T, Kunisada T, Yamada Y. 2012. Genetic ablation of rest leads to in vitro-specific derepression of neuronal genes during neurogenesis. Development 139:667–677. doi: 10.1242/dev.072272

Bruce AW, Donaldson U, Wood IC, Yerbury SA, Sadowski MI, Chapman M, Gottgens B, Buckley NJ. 2004. Genome-wide analysis of repressor element 1 silencing transcription factor/neuron-restrictive silencing factor (REST/NRSF) target genes. Proceedings of the National Academy of Sciences USA 101:10458–10463. doi: 10.1073/pnas.0401827101

Chen Z-F, Paquette AJ, Anderson DJ. 1998. NRSF/REST is required in vivo for repression of multiple neuronal target genes during embryogenesis. Nature Genetics 20:136–142. doi: 10.1038/2431

Gao Z, Ure K, Ding P, Nasaht M, Yuan L, Ma J, Hammer RE, Hsieh J. 2011. The master negative regulator REST/NRSF controls adult neurogenesis by restraining the neurogenic program in quiescent stem cells. Journal of Neuroscience 31:9772–9786. doi: 10.1523/JNEUROSCI.1604-11.2011

Götz M, Huttner WB. 2005. The cell biology of neurogenesis. Nature Reviews Molecular Cell Biology 6:777–788. doi: 10.1038/nrm1739

Götz M, Hartfuss E, Malatesta P. 2002. Radial glial cells as neuronal precursors: a new perspective on the correlation of morphology and lineage restriction in the developing cerebral cortex of mice. Brain Research Bulletin 57:777–788. doi: 10.1016/S0361-9230(01)00777-8

Nechiporuk T, McGann J, Mullendorff K, Hsieh J, Wurst W, Floss T, Mandel G. 2016. The REST remodeling complex protects genomic integrity during embryonic neurogenesis. eLife 5:e09584. doi: 10.7554/eLife.09584

Schoenherr C, Anderson D. 1995. The neuron-restrictive silencer factor (NRSF): a coordinate repressor of multiple neuron-specific genes. Science 267:1360–1363. doi: 10.1126/science.7871435

Yang YJ, Baltus AE, Mathew RS, Murphy EA, Evrony GD, Gonzalez DM, Wang EP, Marshall-Walker CA, Barry BJ, Murn J, Tatarakis A, Mahajan MA, Samuels HH, Shi Y, Golden JA, Mahajnah M, Shenav R, Walsh CA. 2012. Microcephaly gene links trithorax and REST/NRSF to control neural stem cell proliferation and differentiation. Cell 151:1097–1112. doi: 10.1016/j.cell.2012.10.043