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The Eppendorf Award for Young European Investigators was established in 1995 to recognize outstanding work in biomedical science. It also provides the opportunity for young European researchers to showcase their work and communicate their research to a scientific audience. *Nature* is pleased to partner with Eppendorf to promote the award and celebrate the winner's work in print and online. Thea Cunningham from *Nature* talks to the 2014 winner Madeline Lancaster about her work — which showed that complex neuronal tissues resembling early states of fetal human brain can be created *in vitro* from pluripotent stem cells — and how it felt to win the award. To listen to the full interview, visit: nature.com/nature/awards/eppendorf.

About the Award

Madeline Lancaster is the nineteenth recipient of the Eppendorf Award for Young European Investigators, which recognizes talented young individuals working in the field of biomedical research in Europe. The Eppendorf Award is presented in partnership with *Nature*. The winner is selected by an independent jury of scientists under the chairmanship of Reinhard Jahn, Director at the Max Planck Institute for Biophysical Chemistry, Göttingen, Germany. *Nature* and Eppendorf do not influence the selection. For more information see: eppendorf.com/award.

Thea Cunningham: Congratulations on winning the award. How did it feel when you found out?

Madeline Lancaster: It was a big surprise. It's a wonderful opportunity to have my work spotlighted and a real honour to be chosen this year.

TC: Give listeners an insight into what your research involves.

ML: I'm working on a new model system that we've developed for studying human brain development in a dish. We've been able to generate brain tissue from human embryonic stem cells or we've reprogrammed induced pluripotent stem cells to develop into brain tissues that we can use to model brain development and neurological disease.

TC: What led you into this research?

ML: I've been interested for as long as I can remember in the human brain: how it's special and how it develops. We've started

to reach a point in neurobiology where we need some kind of model system where we can look at human-specific processes of brain development. That is what got me interested in going *in vitro* to try to develop some kind of system where we could start to do that.

TC: You've turned pluripotent stem cells into miniature brains. How did you do that?

ML: We give them a few signals to try to make them become neural first. Then we give them the right environment to allow them to develop on their own using their own endogenous signals, to develop into the different brain regions that they would normally develop into *in vivo*.

TC: What did you see as the cells began to grow?

ML: Initially you end up with a white clump of cells. We didn't know what we were looking at. One organoid in particular showed a dark area of pigment [an organoid is a structure that

resembles an organ]. It was a round, circular shape that looked just like eye tissue. When I cut it open I realised that is exactly what it was. It was retinal pigmented epithelium of the developing retina. That was really striking. Once we started staining for different regions not only did we see retina but we saw dorsal cerebral cortex, which is the most extensive part of the human brain. We also saw regions like the hippocampus, which is important for learning and memory, and the choroid plexus, which gives rise to cerebrospinal fluid. We could even see tissues of the ventral forebrain and the midbrain-hindbrain boundary, which is a very important boundary in early brain development.

TC: In a normal brain, there is a lot of interplay between these regions. Did you notice that in your tissue?

ML: One of the most striking aspects is that when you have an organoid that has different regions within the same brain tissue, these

regions can actually interact with each other. It is well known that there are neurons produced in the ventral forebrain that migrate into the dorsal forebrain. When we have organoids that contain both ventral and dorsal regions, that's exactly what we can see. So even over millimetres of distance, these cells automatically know where to go. They know to go from the ventral region to the dorsal region and they don't require any exogenous factors to do that.

TC: Did these regions contain neurons?

ML: Yes. In general, the organoids have a large number of neurons. We have looked at the neurons of the dorsal cerebral cortex, which seem to be functional. They can fire; we see spontaneous calcium surges; they can put out axons; and they can even perform axon guidance together in a way that looks a lot like axon bundling that you would see *in vivo* [axon guidance is the name given to the mechanism in which neurons send axons to a specific target cell]. They even know where to send their axons without any exogenous factors.

TC: So this miniature brain by Oh la Eppendorf A

APPLY FOR THE 2015 EPPENDORF AWARD FOR YOUNG EUROPEAN INVESTIGATORS

We invite biological and biomedical researchers not older than 35 years and working in Europe to apply for the 2015 Eppendorf Award. The deadline for entries is **January 15, 2015**. The prize ceremony on June 25, 2015 will take place at the EMBL Advanced Training Centre (ATC) in Heidelberg, Germany. To find out more visit eppendorf.com/award.