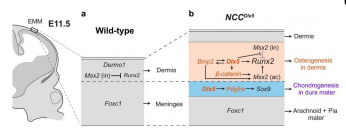


Get your head in the game: One gene's role in cranial development

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Proposed regulatory molecular cascades in the initial layer of the calvarium (early migrating mesenchyme: EMM) at E11.5 in the wild-type mouse (a) and the mouse in which DIx5 is overexpressed in NCC (NCCDIx5: b). ac, activator; in, inhibitor. Credit: Department of Molecular Craniofacial Embryology,TMDU

Mammalian embryonic development is an extremely complex and precise process. Specific molecular events act as cues that tell cells in the embryo where to move and what type to mature into. The expression levels of different genes in these cells can change at certain points of development, helping produce the signals that further the progression. Now, researchers at Tokyo Medical and Dental University (TMDU) have used experiments with mice to show how a particular gene helps direct generation of tissues in the head.

In an article published in *Scientific Reports*, a group of researchers from TMDU determined that expression of a gene called Distal-less homeobox 5 (DIx5) assists certain cells in the mouse head mature into <u>cartilage cells</u>, while others become <u>bone cells</u>. This is critical for proper cranial formation.

Neural crest cells (NCCs) have been a recent focus of developmental biology research because they can transition into many cell types including neurons, and those in the head region additionally differentiate into <u>bone</u> and cartilage. NCCs form an initial layer that constructs dermis and the meninges in early stages of development, then another group of NCCs present at the supraorbital region becomes bone forming <u>cells</u> and the cell domain expands apically as the overlying layer to form the calvarium to protect the brain. From these previous observations, the TMDU group became interested in how experimentally increasing DIx5 expression would affect NCC differentiation.

"We worked with a mouse model to further understand how over-expressed DIx5 affects cranial development," says lead author of the study Tri Vu Hoang. "We used a method where we could force higher expression levels of DIx5 in NCCs in a group of <u>mice</u>, then compared their head development with mice not genetically engineered in this way."

The researchers first confirmed that DIx5 was being expressed in NCCs in the engineered moue as they expected before examining cranial bone and cartilage formation in the two groups of mice. They observed enhanced layers of both bone and cartilage in the mice with higher NCC DIx5 expression compared with the controls.

"We saw an interesting response with DIx5 overexpression in the NCCs," describes Sachiko Iseki, senior author. "NCCs intrinsically forming soft tissues of the initial layer transform into bone and cartilage in the vertex of the DIx5-engineered mice."

The group also examined expression of several bone and cartilage growth-promoting genes in the regions where new cartilage and bone were formed. They observed upregulation of multiple major genes for bone and cartilage formation including ?-catenin and Pdgfr?, bone genes were expressed in the region close to the skin and cartilage genes near the brain in the Dlx5-engineered mice that was not seen in the control animals. The observation shows that the



initial soft tissue layer transform into the two different layers around the time of bone and cartilage formation.

"Our results shed light on early development of the head layer of NCCs in the mouse vertex; there two cell layers exhibiting differentiating potential for bone and <u>cartilage</u>." says Vu Hoang. "Overexpression of DIx5 in NCCs is an artificial situation in <u>embryonic development</u>, but it showed us an interesting phenomenon to discuss uncovered developmental mechanisms in the vertex of the head."

This study provides fascinating data that will help move the developmental biology field forward. The researchers have successfully helped reveal the complicated mechanisms involved with cranial development.

More information: Tri H. Vu et al, Dlx5-augmentation in neural crest cells reveals early development and differentiation potential of mouse apical head mesenchyme, *Scientific Reports* (2021). DOI: 10.1038/s41598-021-81434-x

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