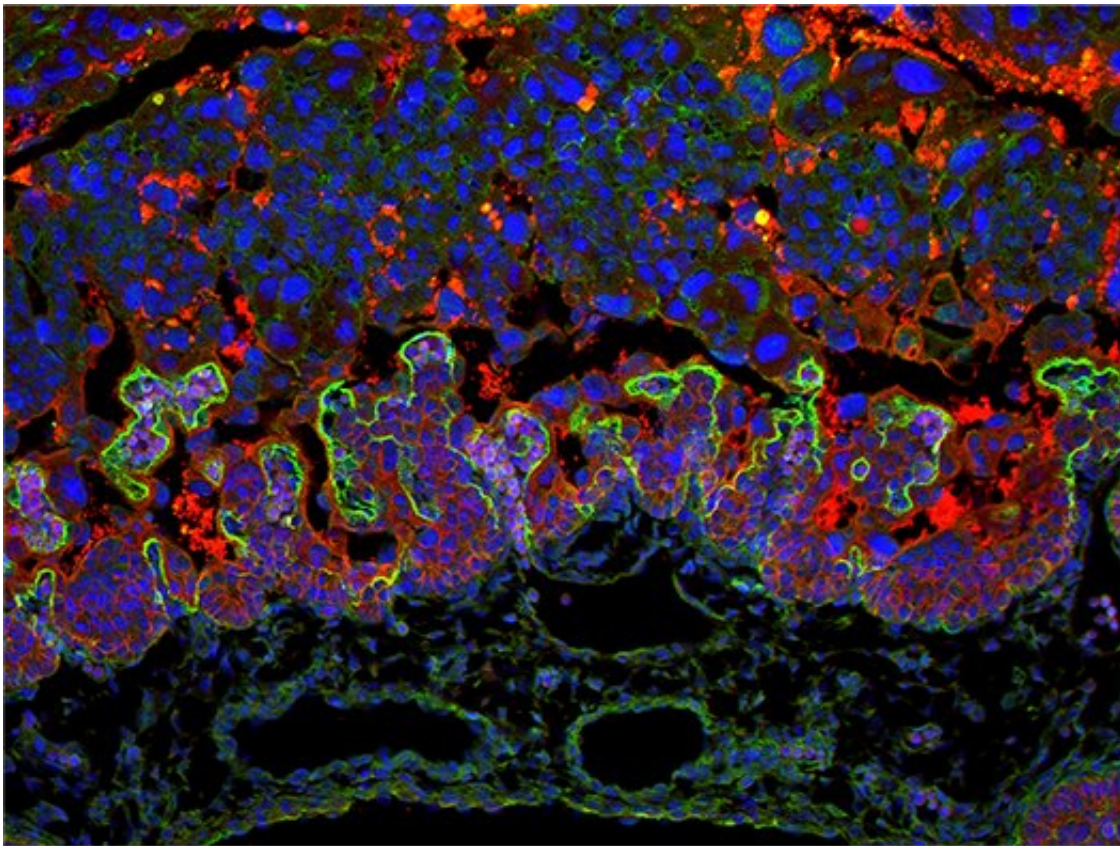


Alomst 70 percent of 103 genes linked to prenatal death affect the placenta

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Cells in a healthy placenta (from mice), the exchange surface between mother and baby is shown in green. Credit: Perez-Garcia, Fineberg et al.

The role of the placenta in healthy fetal development is being seriously under-appreciated according to a new paper published today (14th March). The study was part of the Wellcome Trust-funded "Deciphering

the Mechanisms of Developmental Disorders (DMDD)" consortium. Dr Myriam Hemberger at the Babraham Institute, Cambridge led the research, working with colleagues at the Wellcome Sanger Institute, Cambridge, the Francis Crick Institute, London, the University of Oxford and the Medical University of Vienna, Austria. Writing in the journal *Nature*, the team studied 103 genetic mutations in mice that cause embryos to die before birth. The results showed that the majority, almost 70%, cause defects in the placenta.

Each of the 103 gene mutations causes the loss of a particular factor. Many of these had not been previously linked to [placenta](#) development, and hence the study highlights the unexpected number of genes that affect development of the placenta. By studying a select group of three genes in further detail, the team went on to show that the death of the embryo could be directly linked to defects in the placenta in one out of these three cases. This may mean that a significant number of genetic defects that lead to prenatal death may be due to abnormalities of the placenta, not just the embryo.

Although this research uses mice, the findings are likely to be highly relevant to complications during human pregnancy and the study highlights the need for more work to be done on investigating development of the placenta during human pregnancies.

The placenta is vital for normal pregnancy progression and [embryo development](#) in most animals that give birth to live young, including humans. It provides a unique and highly specialised interface between the embryo and the mother, ensuring an adequate provision of nutrients and oxygen to the embryo. The placenta is also involved in waste disposal from the embryo and produces important hormones that help sustain pregnancy and promote fetal growth. Although previous research has highlighted the pivotal role of the placenta for a healthy pregnancy, its potential contribution to pregnancy complications and birth defects

continues to be overlooked.

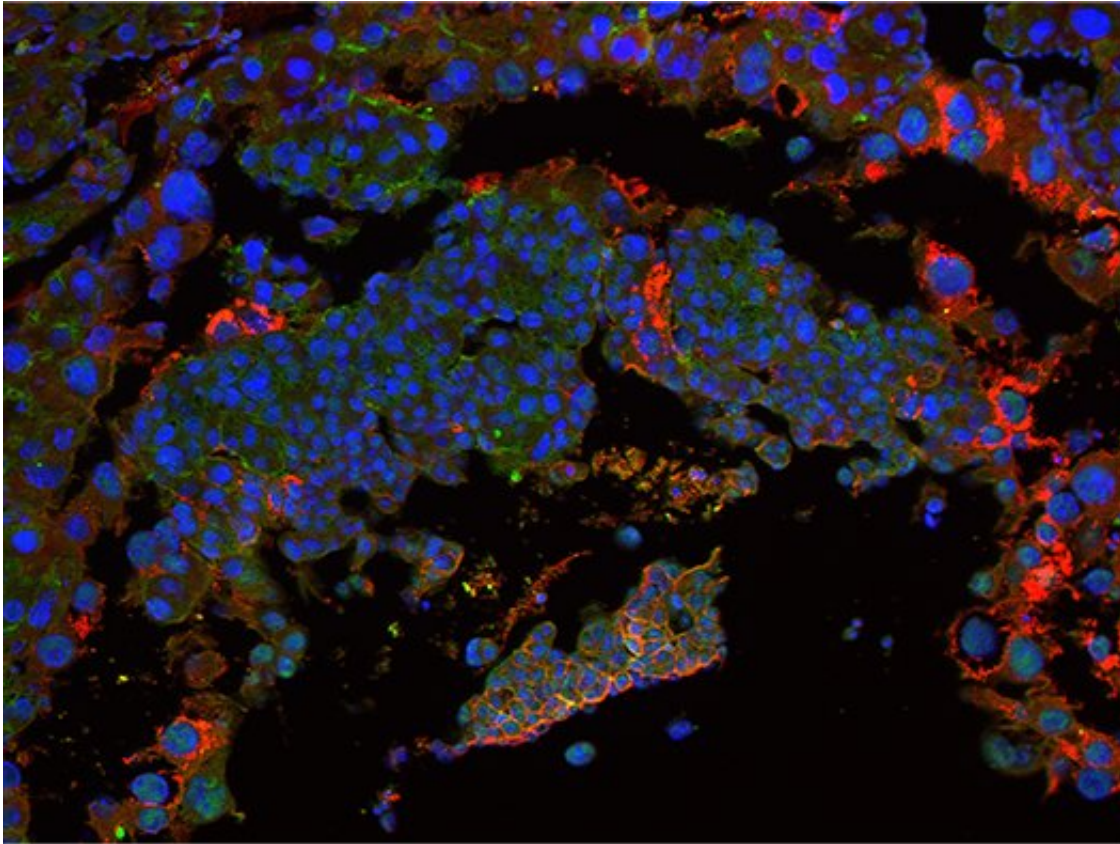
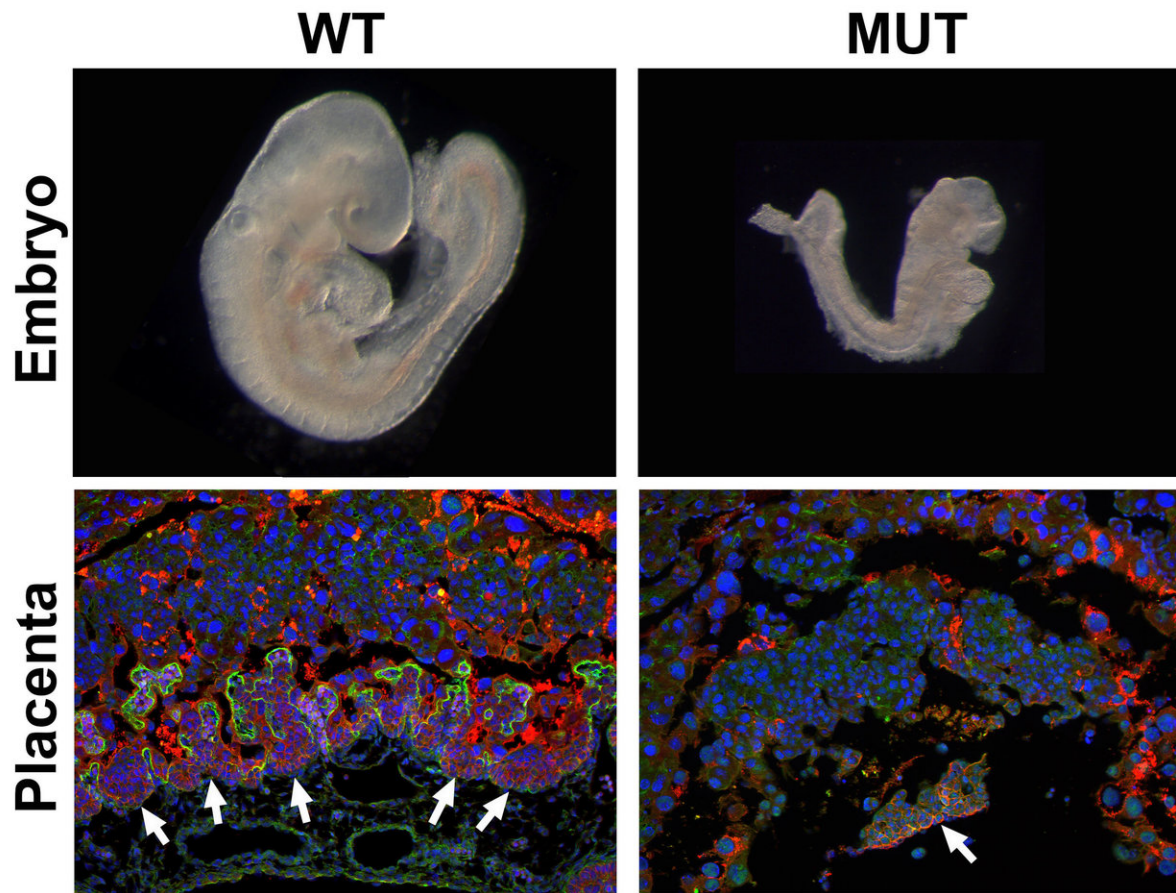


Image illustrating defects in the placenta (in mice) caused by loss of the Nubpl gene. The exchange surface between mother and baby is missing. Credit: Perez-Garcia, Fineberg et al.

Scientists call mutations that cause death in the womb embryonic lethal. Mouse lethal genes are enriched for human disease genes and the affected [embryos](#) often show morphological abnormalities, i.e. changes to their shape and structure. Around one-third of all gene mutants studied in mouse are lethal or subviable (i.e. mutant offspring are less likely to survive than non-mutant pups).

The DMDD consortium investigates the detailed morphological abnormalities in the embryo caused by such gene mutations. Importantly, the DMDD programme includes analysis of placentas from these mutants. Through this uniquely comprehensive assessment, the team was able to show a significant co-occurrence of placental defects alongside abnormalities in the brain, heart and vascular system.

The co-first authors on the paper are Dr Vicente Perez-Garcia and Mrs Elena Fineberg. Dr Perez-Garcia, said: "Analysis of embryonic lethal mutants has largely focused on the embryo and not the placenta, despite its critical role in development. Of the mutations we've studied, far more than expected showed defects in the placenta and this is particularly true for mutations that cause death during the early stages of pregnancy. Intriguingly, our analysis also indicates that issues in the placenta often occur alongside specific defects in the embryo itself."



Mouse embryos part way through development (top) showing a normal embryo and an embryo with a mutation in the *Nubpl* gene, which affects development and leads to prenatal death. Lower images show cells in the corresponding placentas. The green staining marks the exchange surface where nutrients are transferred between the mother and embryo; note that this layer is missing in the *Nubpl* mutant placenta. Red staining shows the cells underlying the exchange surface, these are also mostly missing in the mutant placenta. Credit: Perez-Garcia, Fineberg et al.

The senior researcher on the study was Dr Myriam Hemberger, Group Leader in Epigenetics at the Babraham Institute. Dr Hemberger, said: "Our data highlight the hugely under-appreciated importance of

placental defects in contributing to abnormal embryo development and suggest key molecular nodes governing placentation. The importance of a healthy placenta has often been overlooked in these studies and it is important that we start doing more to understand its contribution to developmental abnormalities."

The placenta-focussed aspect of the DMDD project was carried out at the Babraham Institute where all mutant placentas were analysed and characterised for their morphological defects. The Institute researchers also carried out work using mouse placental stem cells to understand further the functional roles of particular genes in the development of this organ. The mutant mouse lines were produced and bred at the Wellcome Sanger Institute. The detailed morphological analysis of embryos was performed by Dr Tim Mohun at the Francis Crick Institute together with colleagues at the Division of Anatomy, Medical University of Vienna, Austria.

More information: Vicente Perez-Garcia et al, Placentation defects are highly prevalent in embryonic lethal mouse mutants, *Nature* (2018). [DOI: 10.1038/nature26002](https://doi.org/10.1038/nature26002)

Provided by Babraham Institute

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