Tracking Zearalenone: Placental Transfer of a Fungal Toxin

Carrie Arnold

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For a developing fetus, the role of the placenta is simultaneously simple and complex. This ephemeral organ must allow nutrients and oxygen from the maternal bloodstream to reach the fetus but keep out pathogens and other harmful compounds. The placenta also secretes sex hormones, including estrogen and progesterone, to help direct fetal growth and development. Certain chemicals, including endocrine-disrupting chemicals (EDCs), can disrupt this delicate process. A recent article in *Environmental Health Perspectives* suggests that one such EDC, the fungal toxin zearalenone (ZEN), can readily cross the placenta and enter the fetal bloodstream.

The potential impact of disrupting the delicate balance of hormones during pregnancy is one of the major reasons that scientists are concerned about EDCs. “This work showed, in principle, that the transfer and metabolism of zearalenone across the placenta is possible,” says senior author Tina Buerki-Thurnherr, a toxicologist at the Swiss Federal Laboratories for Materials Science and Technology.

ZEN, produced by multiple *Fusarium* species, is found in a broad range of foods, including cereal- and legume-based products. It mimics natural estrogens, says Buerki-Thurnherr, but it is not easily absorbed by the human body, and it is undetectable in the plasma of the majority of the population. A 2019 study found ZEN in the plasma of only 6.5% of participants, with concentrations of 0.063–0.418 μg/L. This work is important in the context of the hypothesis that environmental exposures early in life can have an outsized impact on adult health. Importantly, however, the study did not assess potential health effects of ZEN exposure and is not intended as guidance for people to make dietary decisions.

*Fusarium culmorum*, shown here, is one of several fungal species that produce ZEN. This micrograph shows the fungus’s fibrous white hyphae and globular orange sporodochia grown on pink medium. The sporodochia, which form from the hyphae, may comprise thousands of spores each. Image: © Lesny Ludek/Shutterstock.
authors did not know was whether it could cross this barrier in humans, and if so, how quickly.

Buerki-Thurnherr had previous experience studying the placenta in human cell and tissue models. For this study, her team used a dually perfused *ex vivo* human placenta to model the transfer of ZEN and its nine key metabolites from mother to fetus. In this model, cannulae are inserted into blood vessels on both the maternal and fetal sides of the placenta. Investigators can then introduce compounds of interest on either side to observe how they transfer between mother and fetus. The process is labor-intensive and often fails due to placental leakage and other issues. “It’s definitely not high throughput,” Buerki-Thurnherr says.

The authors successfully perfused six placentas. The maternal circulation of three placentas received a control solution, and three were perfused with 1 μM (318 μg/L) ZEN for 6 hours. Both solutions were collected on the fetal side and analyzed using a new ultra-high-performance liquid chromatography–tandem mass spectrometry method developed by members of the team in a previous study. This method allowed the researchers to analyze up to 75 metabolites simultaneously instead of individually.

The new findings showed that two ZEN metabolites, including a highly estrogenically active form, were detected in fetal circulation within 15 minutes of being introduced on the maternal side. After 6 hours of ZEN perfusion, approximately 15% of the compound was present in the fetal circulation, 31% was present in the maternal circulation, and the rest was metabolized or present elsewhere in the placenta.

The results indicate that ZEN transfer was both fast and efficient, according to Loch-Caruso. She points out that the study is limited by its small sample size and the lack of demographic data about the mothers who donated their placentas. Still, she praised the work as “an important, early first study” in understanding how endocrine disruptors cross the placenta, with potential to impact fetal development.

**References**

1. Bukovsky A, Caudle MR, Cekanova M, Fernando RI, Wimalasena J, Foster JS, et al. 2003. Placental expression of estrogen receptor beta and its hormone binding variant—comparison with estrogen receptor alpha and a role for estrogen receptors in asymmetric division and differentiation of estrogen-dependent cells. Reprod Biol Endocrinol 1(1):36, PMID: 12740031, https://doi.org/10.1186/1747-7827-1-36.
2. European Food Safety Authority Panel on Contaminants in the Food Chain. 2011. Scientific opinion on the risks for public health related to the presence of zearalenone in food. EFSA J 9(6):2197, https://doi.org/10.2903/j.efsa.2011.2197.
3. Warth B, Freindl K, Manser P, Wick P, Marko D, Buerki-Thurnherr T. 2019. Transfer and metabolism of the xenoestrogen zearalenone in human perfused placenta. Environ Health Perspect 127(10):107004, PMID: 31596610, https://doi.org/10.1289/EHP4860.
4. Fan K, Xu J, Jiang K, Liu X, Meng J, Di Mavungu JD, et al. 2019. Determination of multiple mycotoxins in paired plasma and urine samples to assess human exposure in Nanjing, China. Environ Pollut 248:865–873, PMID: 30856502, https://doi.org/10.1016/j.envpol.2019.02.091.
5. Barker DJ. 1995. Fetal origins of coronary heart disease. BMJ 311(6998):171–174, PMID: 7613432, https://doi.org/10.1136/bmj.311.6998.171.
6. Wang Y, Zhao S. 2010. Placental blood circulation. In: *Vascular Biology of the Placenta*. San Rafael, CA: Morgan & Claypool Life Sciences.
7. Mitro SD, Johnson T, Zota AR. 2015. Cumulative chemical exposures during pregnancy and early development. Curr Environ Health Rep 2(4):367–378, PMID: 26341623, https://doi.org/10.1007/s40572-015-0064-x.
8. Bernhoff AH, Behrens GH, Ingebrigtsen K, Langseth W, Berndt S, Haugen TB, et al. 2001. Placental transfer of the estrogenic mycotoxin zearalenone in rats. Reprod Toxicol 15(5):545–550, PMID: 11780962, https://doi.org/10.1016/S0890-6238(01)00159-9.
9. Hutson JR, Garcia-Bournissen F, Davis A, Koren G. 2011. The human placental perfusion model: a systematic review and development of a model to predict *in vivo* transfer of therapeutic drugs. Clin Pharmacol Ther 90(1):67–76, PMID: 21562489, https://doi.org/10.1038/clpt.2011.66.
10. Freindl K, Braun D, Aichinger G, Sieri S, Fang M, Marko D, et al. 2019. A generic liquid chromatography–tandem mass spectrometry exposome method for the determination of xenoestrogens in biological matrices. Anal Chem 91(17):11334–11342, PMID: 31398002, https://doi.org/10.1021/acs.analchem.9b02446.