ALTERATIONS IN NEURAL FATTY ACID METABOLISM CAUSED BY VITAMIN E DEFICIENCY

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Outline

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Significance

- Oxidative damage in the brain is believed to play a role in degenerative diseases such as Alzheimer's and dementia.
- Patients with Alzheimer's have low levels of vitamin E in their cerebrospinal fluid (Kontush et al., Acad Science 2004).
- Alpha tocopherol supplementation of 2000 IU has been shown to slow the progression of Alzheimer's and dementia, but the mechanism is unknown. (Sano M, Ernesto C, Thomas RG, et al, N Engle J Med. 1997.)
Vitamin E
(Alpha tocopherol)

Exists in 8 isomers:

- Alpha, beta, gamma, and delta tocopherol
- Alpha, beta, gamma, and delta tocotrienol
- Alpha tocopherol is preferentially retained due to the action of the tocopherol transfer protein in the liver (DRI, 2000)
Vitamin E
(Alpha tocopherol)

- Lipid soluble vitamin
- Principal role is to protect polyunsaturated fatty acids from peroxidation
- Deficiency can cause peripheral neuropathy
Vitamin E
(Alpha tocopherol)

- Donates a hydrogen from the free hydroxyl group on the aromatic ring
DHA (Docosahexaenoic acid) (22:6 ω-3)

- Important long chain fatty acid created by elongating omega 3 fatty acids or consumed as fish oil
- Comprises ~50% of brain fatty acids (Bazar, et al. AnnuRev Nutr 2011)
- Wide variety of cellular functions:
  - Gene regulation, membrane fluidity, precursor for some signaling molecules
DHA is very susceptible to peroxidation due to the high amount of unsaturation in the aliphatic tail.
Hypothesis

We hypothesize that vitamin E protects DHA from lipid peroxidation.
Prediction

Zebrafish fed a vitamin E deficient diet:

1) Will have increased levels of lipid peroxidation

2) Will have an increase in mRNA expression of enzymes responsible for polyunsaturated fatty acid synthesis to replace any DHA lost to lipid peroxidation

3) However, due to the deficiency of vitamin E in the diet, the increase in mRNA expression will be insufficient to replace DHA lost in the brain.
Model Organism
Zebrafish

- Currently established model in the Traber/Tanguay laboratories
- Similar fatty acid metabolism pathways as humans (Lebold et al J Nutr, accepted)
- Easily manipulated diet with defined ingredients
Method Overview

- Defined diet without Vit E (E-, n=30)
- Defined diet with Vit E (E+, n=30)

- Brains
- Eyes
- Livers

- Protein expression changes
- mRNA expression changes
- Fatty Acid concentrations
Proteins involved in fatty acid synthesis

1. Sterol regulatory binding factor 1 (SREBP1)
2. Fatty acid desaturase (fads2)
3. Elongase (elovl2)
4. Elongase (elovl4)
Sterol regulatory binding factor 1 (SREBP1)

- Regulates genes related to lipid metabolism and fatty acid synthesis
Fatty acid desaturase (*fads2*)

- Removes two hydrogens from a fatty acid in order to create a double bond
- Required for the synthesis of DHA from Omega-3 fatty acids
- mRNA measured in the liver and brain
Elongase (*elovl2*)

- Catalyzes the synthesis of polyunsaturated very long chain fatty acids
- Elongates fatty acids by adding 2 carbons
- mRNA measured in the liver and brain
Elongase (*elovl4*)

- Catalyzes the synthesis of polyunsaturated very long chain fatty acids
- Elongates fatty acids by adding 2 carbons
- Specifically expressed in neural tissues
- mRNA measured in the eye and brain
Tripathy S, Torres-Gonzalez M, Jump DB. J Lipid Res. 2010 Sep;51(9):2642-54
Methods

- Adult zebrafish fed a vitamin E sufficient or deficient diet were euthanized and brains, livers, and eyes removed for analysis
Methods

- Fatty acid concentrations were determined using high pressure liquid chromatography coupled to a single-quad mass spectrometer.
- Vitamin E concentrations were determined using high pressure liquid chromatography with electrochemical detection.
Methods

- mRNA expression, evaluated using quantitative real-time PCR, were:
  - fatty acid desaturase (fasd2)
  - elongase (elovl2, elovl4)
- Protein expression of SREBP1 was determined using western blotting
Results
Diet impact on vitamin E levels

Diet

Tissue

Diet x Tissue

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<th>Diet x Tissue</th>
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<tbody>
<tr>
<td>P&lt;0.0001</td>
<td>P=0.2772</td>
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Columns not sharing the same letter are significantly different.
Vitamin E deficiency did not affect fatty acid concentrations in the liver.
Vitamin E deficiency did not affect fatty acid concentrations in the liver.

- **Alpha Linolenic Acid**
- **DHA**
- **EPA**
Vitamin E deficiency did not affect Elovl2 or FADs2 mRNA expression in the liver.
Fatty Acid Concentration in the Brain

- No data available due to equipment failure.
Vitamin E deficiency did not affect mRNA expression in the brain

**Elovl4**

**Elovl2**

**Fads2**
Vitamin E deficiency increases eye EPA and linoleic acid concentrations

**Linoleic Acid**

- E-: 20 ng/mg tissue
- E+: 10 ng/mg tissue
  - p = 0.002

**EPA**

- E-: 0.4 ng/mg tissue
- E+: 0.2 ng/mg tissue
  - p = 0.03
Vitamin E deficiency tended to decrease DHA concentration in the eye.

**Alpha Linolenic Acid**

**Arachidonic Acid**

**DHA**

*P* = 0.09
Correlation between eye DHA and $\alpha$-tocopherol

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<tr>
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Vitamin E deficiency did not affect eye Elovl4 mRNA expression
Summary

- With regard to neural tissues, vitamin E deficiency in eyes (brain unstudied):
  - tended to decrease DHA levels
  - vitamin E and DHA concentrations were positively correlated
  - Other PUFA concentrations increased

- With regard to the liver, vitamin E deficiency:
  - did not change liver fatty acid concentrations
Vitamin E deficiency did not alter mRNA expression of FADs2, Elovl2, and Elovl4 in any tissues studied (brain, eye, liver)
Conclusion

- Vitamin E in the eye was not depleted sufficiently to allow DHA oxidation
  - With lower vitamin E concentrations less DHA was observed
- Vitamin E deficiency did not alter mRNA expression of FADs2, Elovl2, and Elovl4 in any tissues studied (brain, eye, liver)
- Vitamin E deficiency did not change liver fatty acid concentrations
- More samples are being analyzed
Limitations

- SREBP expression: Data not available due to western blotting optimization problems
- Data not available for DHA in the brain due to equipment failure
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