New Treatments in Fragile X Syndrome

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Early in childhood may be the best time to demonstrate efficacy

- ASD studies have demonstrated this with the Early Start Denver Model (ESDM Dawson et al 2010, 2012)
- Neurochemical abnormalities can be modified before the brain infrastructure is permanently changed so ongoing development can be normalized
SEROTONIN SYNTHESIS CAPACITY:
Autistic vs. Non-autistic Children

Chugani et al., 1999
Syndromic and non syndromic forms of ASD have reduced tryptophan metabolism

- Boccoto et al 2013 Mol Aut: Studied 137 lymphoblastoid cell lines of patients with neurodevelopmental disorders with (80) and without ASD and 78 controls (2.5 to 35yo)
  - Metabolic profiling demonstrated deficits in tryptophan metabolism in ASD with reduced NADH, not seen in ID or schizophrenia
  - Found abnormal gene expression of enzymes in tryptophan metabolism pathway
Tryptophan metabolic pathway

The figure illustrates the main intracellular pathways involving tryptophan. Genes with reduced expression in our microarray dataset are in blue, genes with increased expression are in red. Genes with statistically significant reduction of expression are underlined. Reactions generating NADH are indicated in the top section of the figure. FAD, flavin adenine dinucleotide.
Sertraline is an optimal SSRI for FXS and ASD

- SSRIs stimulate neurogenesis in the mouse and in humans (Malberg et al 2004; Warner-Schmidt et al 2006)
- SSRIs stimulate BDNF levels (Bianchi et al 2010; Taler et al 2013)
- Sertraline in FXS has a calming effect to the anxiety and may directly stimulate speech or improve language as a secondary effect from less anxiety
- Sertraline is the only SSRI to stimulate dopamine levels in the striatum and nucleus accumbens (Kitaichi et al 2010)
Sertraline Treatment in Early Childhood in FXS

A retrospective study of 45 children followed 12 to 50 months and 11 treated with sertraline: significant differences in expressive and receptive language in TX vs non treated (p=0.0001 and p=0.0071 respectively)

Winarni et al 2012 Autism Treatment and Research
Randomized, double blind, controlled trial of sertraline in 2 to 6yo with FXS lasting 6 months

Baseline and 6 month Follow-up:
1* CGI-I, MSEL Expressive Language

2* MSEL: Receptive, fine motor, visual reception, Composite T score, Sensory Processing Measure-(SPM) Preschool version, Visual Analogue Scale

Laura Greiss Hess PhD, OTR
Kerrie Lemons Chitwood PhD, CCC
initiated study

Sarah Fitzpatrick
NFXF summer student fellow
finalized the study and data
Clinical Global Impression-Improvement (CGI-I)

No significant difference between groups (2.28 [1.06] vs. 2.59 [0.84], \( P = 0.244 \))
Mullen Scales of Early Learning (MSEL)

- Cognitive T-Score Sum: $P=0.047$
- Visual Reception Age Equivalent Score: $P=0.031$
- Fine Motor Age Equivalent Score: $P=0.005$
- Visual Reception T-Score: $P=0.038$
- ELC Standard Score: $p=0.008$
- Fine Motor Raw Score: $P=0.007$
- Combined Age Equivalent Score: $P=0.007$
- Fine Motor T-Score: $P=0.007$
- Receptive Language T-Score: $P=0.007$
- Receptive Language Age Equivalent Score: $P=0.007$
- Receptive Language Raw Score: $P=0.007$
- Expressive Language Age Equivalent Score: $P=0.007$
- Sertraline - Placebo Change
Conclusions

• MSEL visual perception and fine motor subtests, the composite T score, and the combined mean of all the subtests scores were improved on sertraline vs placebo so cognitive and motor benefits and improvement in social participation on SPM-P. Those with ASD with most significant benefit and improved expressive language score. It was safe without signif AEs

• A significantly beneficial treatment by age 2 yo reinforces the need for early diagnosis and even newborn screening

• Families wanted to continue on sertraline after the study and long term follow-up is needed.

• Currently enrolling 2 to 6yo with ASD for a controlled trial of low dose sertraline
AFQ056 + PILI in 3-6yo FXS through NeuroNEXT

- Effects of AFQ056 on Language Learning in Young Children with FXS with lead in PILI for 6 months
- Change paradigm/create model for development of mechanism targeted pharmacotherapy in NDDs – effects of drug on plasticity
- Address many Quandries: incorporate young age (3-6y), longer trial, objective measures, learning intervention, biomarkers for target engagement
- Randomize to AFQ056 or placebo - lead-in period when adjust to best dose
- Extension for 8 months on AFQ056 for all participants

Study Team  PI: Elizabeth B-Kravis; Co-PIs: Randi Hagerman, Len Abbeduto; Co-Is: Walter Kaufmann, Craig Erickson, David Hessl, Flora Tassone
Controlled trial of lovastatin in 10 to 18 yo combined with Parent Implemented Language Intervention (PILI)

- **Lovastatin** is an inhibitor of the rate-limiting enzyme in cholesterol biosynthesis and an FDA-approved treatment for hyperlipidemia (Acosta et al 2011).
- Lovastatin down-regulates the RAS-ERK1/2 pathway and lowers the excessive protein synthesis in FXS.
- In FX KO mice lovastatin rescues seizures and lowers excess protein production in KO mouse (Osterweil et al 2013).
- Lovastatin was beneficial in open label trial in FXS.
- Lovastatin has anti-inflammatory effects.
lovastatin inhibits Ras farnesylation by targeting the upstream mevalonate pathway and thereby downregulates ERK1/2 activation
PILI has demonstrated efficacy in FXS (n=10 in Rx and n=9 in controls over 20 weeks) time in shared story telling jointly engaged.

Data from McDuffie and Abbeduto 2016
Maternal Use of Intervention Strategies were significantly improved with PILI

Data from McDuffie et al 2016
Every one gets PILI intervention

- PILI is twice a week with S and L therapist and with behavior interventionist on skype in your home
- Controlled trial of lovastatin increasing from 10 mg gradually up to 40 mg if tolerated
- Total trial lasting 18 weeks ages 10 to 18yo
- Contact Erika Bickel (esbickel@ucdavis.edu) at the MIND institute or rjhagerman@ucdavis.edu
GABA_A receptor expression is down-regulated in FXS

- **GABA_A** expression is down-regulated in the KO mouse (D’Hulst et al 2007; Kooy et al 2005)

- **GABA_A agonists**: Ganaxolone
  - Investigational medication with efficacy in infantile spasms and other types of epilepsy: A controlled double blind cross-over trial (each arm 7 weeks) in children with FXS (6-18y) funded by DOD is finished. Marinus supplied the ganaxolone; Ligsay et al 2016
  - Targeting improvement in anxiety and behavior
  - Frank Kooy has studied 11 patients in Belgium and we studied 48 patients at the MIND: total 59 in study
Primary Outcome Measures with no significant efficacy: CGI-I

\[ p = 0.448 \]
Post-hoc: High Anxiety (PARS) Significant efficacy

**VAS-Anxiety**

- GNX: Baseline 0, End 4
- Placebo: Baseline 0, End 3

**ADAMS-General Anxiety**

- GNX: Baseline 10, End 6
- Placebo: Baseline 10, End 10

$p=0.023$ and $p=0.035$ respectively.
Adverse events and plans

- Ganaxolone was generally safe, although some experienced sedation which was the main adverse event.
- Data shows those with high anxiety i.e. PARS above 13 demonstrated the best efficacy.
- Additional studies are warranted for this subgroup of patients with FXS.
Additional $\text{GABA}_{\alpha}$ agonists in FXS

- Alphaxolone improved anxiety and seizures in KO mouse (Huelens et al 2012)
- Gaboxadrol normalized neuronal hyperactivity in amygdala and PPI in KO mouse (Olmos-Serrano 2010;2011)
- Ovid announced plans to carry out controlled trial of gaboxadrol in Angelman syndrome and in FXS
- Could allopregnanolone be helpful in FXS?
Yoga and Mindfulness Meditation improves GABA inhibition
Cannabinoid Structures: Cannabidiol (CBD) is not psychotropic.

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<thead>
<tr>
<th>Cannabinoid</th>
<th>Structure</th>
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<tr>
<td>Δ⁹-Tetrahydrocannabinol</td>
<td><img src="image1.png" alt="Structure" /></td>
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<tr>
<td>Cannabidiol</td>
<td><img src="image2.png" alt="Structure" /></td>
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Cannabidiol (CBD)

- The lack of FMRP in FXS leads to a reduction of endocannabinoid system and lowering of endogenous ligands of CB1 and CB2 receptors, 2 arachidonoylclyclycerol (2AG) and anandamide (AEA)
- CBD increases 2AG and AEA and indirectly a GABA agonist effect
- 2 companies interested in CBD trials in FXS: GW pharmaceuticals and Zynerba
- Anecdotal evidence of improvements with CBD in individual cases and maybe helpful for carriers too, so parents have mentioned
Metformin a type 2 diabetes med

- Known to help overeating and obesity
- Can prevent cognitive deficits in diabetics
- Helpful in several patients with Prader-Willi Phenotype of FXS (obesity and hyperphagia), present in less than 10% of FXS
- Drosophila FX model: insulin-like peptide 2 (dilp2) in insulin-producing cells elevated insulin signaling via PI3K/Akt/mTOR pathway (Monyak et al 2016)
- Defect in circadian rhythm and short and long-term memory improves in FX fly with metformin
Defects in FX fly improved with metformin

(Monyak, et al., 2016)
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<tr>
<th>Age/Sex</th>
<th>Diagnosis</th>
<th>Concerns</th>
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<tbody>
<tr>
<td>18yo Male</td>
<td>FXS ID ASD, ADHD Overweight</td>
<td>Aggression Picking/scratching Hyperactivity Perseveration Hyperphagia</td>
</tr>
<tr>
<td>13yo Male</td>
<td>FXS:PWP ASD, ID OSA, Type II DM HgbA1c7.7 Glucose 174</td>
<td>Anxiety Perseveration Hand flapping Poor eye contact Hyperphagia</td>
</tr>
<tr>
<td>19yo Male</td>
<td>FXS:PWP ASD, ADHD, ID Obese</td>
<td>Behavioral outbursts Speaking in short phrases but not sentences Hyperphagia</td>
</tr>
<tr>
<td>60yo Female</td>
<td>FXS, ID HTN Obese</td>
<td>Memory Agitation and anxiety Hyperphagia</td>
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Neuren Phase 2 placebo controlled trial with IGF1 analogue

- Safe in males 14-40 yo with FXS over 42 days of treatment carried out at 5 US centers. Highest dose was most effective
Eye Tracking in FXS As Outcome

TD:

FXS:

Calm  Happy  Fear

Based on data from Farzin, Rivera, & Hessl (2009)
Eye Tracking as Outcome in AFQ056

Sansone et al 2016 Gaitlinberg
AFQ056 was well tolerated although more side effects in the 100 mg bid dose

Gradual improvement in symptoms with the ABC-\(C_{\text{FX}}\) but not more remarkable than placebo in the controlled trial. No learning paradigm or cognitive outcome measure

AFQ056 may be helpful in young children and this will be assessed in a study of 3 to 6 yo with FXS in combination with intensive language training (NeuroNext multicenter trial funded by NIH)
Targeted Treatments must be combined with innovative educational programs

- If synaptic connections are improved with targeted treatment we must enhance these connections with educational interventions
- Combine treatment trials with educational interventions, digital programs such as CogMed, Headsprout for reading, AT devices, iPAD apps
Collaborators

UC Davis School of Medicine

Dept. Biochem & Molec. Medicine
Paul Hagerman     Flora Tassone
Anna Ludwig
Greg Mayeur      Chris Raske

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Elizabeth Berry-Kravis  Deb Hall  Christopher Goetz

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Danuta Loesch      Richard Huggins

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