Succinic Semialdehyde Dehydrogenase Deficiency: Review of the Natural History Study

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Abstract

Objective: The SSADHD Natural History Study was initiated in 2019 to define the natural course and identify biomarkers correlating with severity. Methods: The study is conducted by 4 institutions: BCH (US clinical), WSU (bioanalytical core), USF (biostatistical core), and Heidelberg (iNTD), with support from the family advocacy group (SSADH Association). Recruitment goals were to study 20 patients on-site at BCH, 10 with iNTD, and 25 as a standard-of-care cohort. Results: At this half-way point of this longitudinal study, 28 subjects have been recruited (57% female, mean 9 years, range 18 months–40 years). Epilepsy is present in half and increases in incidence and severity, as do psychiatric symptoms, in adolescence and adulthood. The average Full Scale IQ (FSIQ) was 53 (Verbal score of 56, Non Verbal score of 49), and half scored as having ASD. Although there was no correlation between gene variant and phenotypic severity, there were extreme cases of lowest functioning in one individual and highest in another that may have genotype-phenotype correlation. The most common EEG finding was mild background slowing with rare epileptiform activity, whereas high-density EEG and magnetoencephalography showed reduction in the gamma frequency band consistent with GABAergic dysfunction. MR spectroscopy showed elevations in the GABA/NAA ratio in all regions studied with no crossover between subjects and controls. Conclusions: The SSADH Natural History Study is providing a unique opportunity to study the complex pathophysiology longitudinally and derive electrophysiologic, neuroimaging, and laboratory data for correlation and to serve as biomarkers for clinical trials and prognostic assessments in this ultra-rare inherited disorder of GABA metabolism.

Keywords
epilepsy, genetics, inborn errors of metabolism, intellectual disability, metabolism, neuroimaging

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Succinic semialdehyde dehydrogenase (SSADH) deficiency is an autosomal recessive inherited disorder of degradation of gamma-aminobutyric acid (GABA), the major inhibitory neurotransmitter of the brain. GABA is initially metabolized by GABA transaminase, which utilizes pyridoxal-5-phosphate (P5P) as a cofactor and leads to formation of succinic semialdehyde, an unstable and toxic intermediate converted into succinic acid via SSADH (Figure 1). This should lead to entry of succinate into the tricarboxylic acid, leading to the generation of ATP as well as formation of alpha-ketoglutarate, which further leads to synthesis of glutamate. There is a 1:1 conversion of glutamate, the major excitatory neurotransmitter of the brain, to GABA, the latter mediated by the enzyme glutamic acid decarboxylase (GAD), which also uses P5P as a cofactor. In the absence of SSADH, the succinic semialdehyde is instead converted to gamma-hydroxybutyrate (GHB), also called 4-hydroxybutyric

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acid, which is detected in physiologic fluids as well as other dicarboxylic acids, including 4,5-dihexanoic acid.

The disorder has protean manifestations, with most cases characterized by developmental delay and hypotonia identified within the first 2 years of life, communication disorder which persists as severe impairment in expressive language, intellectual deficiency, and in many cases a variety of neurologic and psychiatric manifestations including epilepsy, ataxia, movement disorders, obsessive-compulsive symptoms, anxiety, sleep disruptions, and hallucinations.1-3 Although the course is typically nonprogressive, there are acute presentations, including infantile encephalopathies,4 as well as evidence of deterioration, especially in adulthood.5

The disease was first discovered following reports of elevations in GHB6 leading to confirmation of the enzymatic deficiency.7 Mammalian SSADH was purified in 1992,8 and the ALDH5A1 gene was cloned in 19959 (Figure 2). Although the clinical phenotype was then being described,1,3,7,10 a series of experiments emanated from development of a murine model using genetic degradation technology.11 The affected mice manifest failure to thrive and evolution of absence to vibrissal twitching seizures with lethality by 3 weeks of age. Preclinical therapeutic trials included taurine (a high constituent of murine breast milk and used due to advent of seizures on weaning of the suckling mice), the GABA-transaminase inhibitor vigabatrin, GHB receptor antagonist NCH-382, GABA(B) receptor antagonist CGP35348 and later named SGS742,12 and the ketogenic diet13 although with concern regarding applicability to patients.14 These early studies led to later clinical trials15-17 (Figure 3). In addition, investigations elucidated widespread neurometabolic alterations18 and the molecular underpinnings of the disorder, from disease-associated gene variants19 to the crystal structure of the enzyme.20 Subsequent animal studies showed dysfunction of the mTOR pathway with potential rescue using rapalog therapy21 and early work showing feasibility and promise of enzyme replacement therapy.22

Meanwhile, neurophysiologic studies and immunohistochemistry disclosed abnormalities of both GABA (A) and (B) receptor function in the murine model,23,24 followed by evidence for chronic overuse downregulation of these receptors in patients based on flumazenil-ligand positron emission tomography25 and transcranial magnetic stimulation studies focused on GABAergic innervation.26

The SSADH deficiency (SSADHD) natural history study is an NICHD (National Institutes of Health [NIH]) sponsored study with a single US site for clinical activity (Boston Children’s Hospital), biorepository and bioanalytical core (Washington State University), and statistical core (Florida

Figure 1. The metabolic pathway of gamma-aminobutyric acid (GABA).

1. Glutaminase
2. Glutamic acid decarboxylase
3. Homocarnosinase/carnosinase
4. GABA transaminase
5. Succinic semialdehyde dehydrogenase
6. Tricarboxylic acid cycle
State University). There is international recruitment through the iNTD (international Neurotransmitter Disease Association) and involvement by the family advocacy organization, the SADH Association. Patients are evaluated on alternate years on site, and receive annual laboratory testing. Patients unable to attend one of the clinical centers are enrolled into a “standard of care” cohort, with questionnaire-based evaluations and procurement of laboratory samples (Figure 4).
As an observational study, the objectives on inception were to justify the incorporation of SSADHD in newborn screening panels by defining the clinical, neurophysiological, and biochemical spectrum and natural course of the disease. Secondary objectives were to identify biomarkers that correlate and predict the clinical presentation and severity, develop a blood spot-based GABA assay adaptable to high-throughput newborn screening platforms, and establish an SSADHD international registry and biobank for future investigations of pathogenesis and therapy. The primary outcomes are a clinical severity score, neuropsychological results, neurophysiological data from transcranial magnetic stimulation, electroencephalography (EEG), and in selected cases high-density EEG and magnetoencephalography. In addition, MRI and MR spectroscopy with sequences and quantification dedicated to quantification of GABA are administered. The primary goals of the study are to define age-dependent and interindividual variation of study outcomes and to correlate neurophysiological and biochemical outcomes with clinical outcomes. A secondary goal is to determine the power of neurophysiological and biochemical outcomes measured at baseline to predict clinical outcomes at the end of the study.

Methods
The design of the study is prospective, mixed with longitudinal and cross-sectional assessments over a period of 5 years. The enrollment target was 55 patients total: 20 at BCH, 10 at iNTD, and 25 who were unable to receive on-site diagnostic testing but are included as a standard-of-care cohort for whom historical and laboratory data could be obtained. The BCH patients are recruited from an ongoing registry that includes 143 patients (52% female) as of February 2020, with a median age of 9 years (range 8 weeks–63 years). Eligible patients are any individual with confirmed SSA DH deficiency, via persistent 4-OH-butyric aciduria and pathologic ALDH5A1 variants, of any age. Patients in all cohorts receive electronic surveys every 6 months.

The BCH cohort receives comprehensive evaluation every 2 years, including brain imaging (MRI, MRS), EEG, transcranial magnetic stimulation, and clinical and neuropsychological testing in addition
to laboratory specimens including blood, urine, hair, saliva, and stool samples, and in some cases skin biopsy. The laboratory specimens are entered into the biorepository, and measurements include blood GABA, GHB, glutamate, and allopregnanolone in addition to blood spots for GABA. Ancillary measurements and procedures include blood oxidative stress markers (glutathione [GSH], 4-hydroxyxonenal [4-HNE], 4-hydroxyxonenal glutathione [4-HNE-GSH]), metabolites (D-2-hydroxyglutarate [D-2-HG], succinic semialdehyde [SSA], 4,5-dihydroxyhexanoic acid [DHHA], homocarnosine, urine metabolites (GHB, D-2-HG, SSA), hair GABA and GHB, stool GABA and microbiota, and gene expression with samples banked for RNA-Seq next-generation sequencing. In addition, a subset of patients was able to receive high-density EEG and magnetoencephalography testing at BCH for spontaneous recordings as well as evoked somatosensory potentials.

Neuropsychological assessments include a clinical parent/subject interview and observations of current behaviors, age-appropriate cognitive and adaptive measures (Mullen Scales of Early Learning; Differential Abilities Scale, Second Edition; Wechsler Abbreviated Scales of Intelligence; Vineland Adaptive Behavior Scales, 2nd Edition), as well as academic and language measures (Receptive Language and Expressive Language [REEL-3]), behavioral measures (Achenbach Adult Behavior Checklist and Child Behavior Checklist), autism spectrum disorders and related symptom measures (Autism Diagnostic Observation Scale; Behavior and Sensory Interests Questionnaire), and motor capabilities using the Movement Assessment Battery for Children–2nd Edition or MABC-II.

Based on the constellation of neurologic and psychological impairments in this patient population, an overall Clinical Severity Score was developed to be employed and tested for reliability. This is based on a composite score ranging from 5 (profound impairment) to 25 (no impairment), calculated using scores from 5 clinically significant subdomains (cognition, communication, motor skills, psychiatric presentation), and epilepsy) (Table 1).

Given the recognized variable clinical presentation depending on age, the analysis of clinical characteristics was stratified as above and below 12 years of age. The standard of care cohort was excluded from the analysis because of the nature of the limited visit conducted off-site. Qualitative variables are expressed in percentages, and relationship contrasts were analyzed using the chi-square (or Fisher exact test). Statistical analyses were conducted with SPSS, version 24.0 (IBM Corp, Armonk, NY).

Results

At the time of the July 2020 Conference, 28 SSADH deficiency patients have been recruited into the natural history study, 12 males (43%) and 16 females (57%), with a median age of 9 years and age range of 18 months–40 years. Controls have been recruited for comparison to our neuroimaging and neurophysiologic procedures, specifically 13 (6 males/46%, 7 females/54%, median age 16 years with age range 6–35 years). The BCH on-site cohort is currently composed of 25 subjects, 16 in the pediatric group (<12 years) and 9 in the adolescent/adult group (12+ years). Their clinical characteristics are shown in Table 2. The SOC cohort has 3 subjects, 2 in the pediatric group (<12 years) and 1 in the adolescent/adult group (12+ years). Epilepsy is a comorbid diagnosis in half of the patients and increases in incidence during adolescence and adulthood (P < .001). The clinical severity of epilepsy also correlated with age based on seizure frequency, showing a stronger association in the older cohort (P = .002). There was also an increased incidence of obsessive-compulsive disorder in the older cohort (P = .003) as well as severity in psychiatric impairment based on the criteria used in the Clinical Severity Scale (P = .09).

### Table 1. Clinical Severity Score (25 = No Impairment; 5 = Profound Impairment).

<table>
<thead>
<tr>
<th>Cognitive</th>
<th>1</th>
<th>IQ &lt; 60</th>
<th>2</th>
<th>IQ 60-69</th>
<th>3</th>
<th>IQ 70-79</th>
<th>4</th>
<th>IQ 80-89</th>
<th>5</th>
<th>IQ 90+</th>
</tr>
</thead>
<tbody>
<tr>
<td>Communication</td>
<td>1</td>
<td>Nonverbal</td>
<td>2</td>
<td>Ambulatory with assistance</td>
<td>3</td>
<td>Ambulatory with ataxia</td>
<td>4</td>
<td>Incoordination</td>
<td>5</td>
<td>Normal</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>1</td>
<td>Profound OCD, interferences with ADLs</td>
<td>2</td>
<td>Active convulsive seizures, no status epilepticus</td>
<td>3</td>
<td>Intermittent seizures, including convulsive</td>
<td>4</td>
<td>Intermittent seizures, only nonconvulsive (absence)</td>
<td>5</td>
<td>No Seizures</td>
</tr>
</tbody>
</table>

### Table 2. Clinical Characteristics of BCH SSADHD Subjects Enrolled in Natural History Study.

<table>
<thead>
<tr>
<th></th>
<th>BCH Pediatric Cohort (&lt;12 y), n (%)</th>
<th>BCH Adolescent/Adult Cohort (&gt;12 y), n (%)</th>
<th>P Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intellectual</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>disability</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Fine motor delay</td>
<td>15 (94)</td>
<td>9 (100)</td>
<td>&gt;.99</td>
</tr>
<tr>
<td>Gross motor delay</td>
<td>16 (100)</td>
<td>9 (100)</td>
<td>&gt;.99</td>
</tr>
<tr>
<td>Speech delay</td>
<td>16 (100)</td>
<td>9 (100)</td>
<td>&gt;.99</td>
</tr>
<tr>
<td>Seizures</td>
<td>1 (6)</td>
<td>7 (78)</td>
<td>&lt;.001*</td>
</tr>
<tr>
<td>Sleep disturbance</td>
<td>8 (50)</td>
<td>7 (78)</td>
<td>.229</td>
</tr>
<tr>
<td>Hypotonia</td>
<td>16 (100)</td>
<td>9 (100)</td>
<td>&gt;.99</td>
</tr>
<tr>
<td>OCD/Anxiety</td>
<td>6 (38)</td>
<td>9 (100)</td>
<td>.003*</td>
</tr>
<tr>
<td>Ataxia</td>
<td>14 (88)</td>
<td>5 (55)</td>
<td>.143</td>
</tr>
<tr>
<td>ADHD</td>
<td>13 (81)</td>
<td>4 (44)</td>
<td>.087</td>
</tr>
</tbody>
</table>
Of 25 subjects studied with EEG, 16 (64%) had diffuse background slowing (mild in all cases), and only occasional epileptiform features manifest as generalized spike-and-wave in only 1 (Figure 5), focal spikes in 2 subjects (ages 5 and 9, and without history of seizures), and sharp waves in 2.

Neuropsychological evaluations have been completed on 21 subjects. The average full scale IQ measurement was 53 (IQR 49-61, range 30 [floor value] to 87), including verbal of 56 (IQR 46-65, range 30 [floor value] to 95) and nonverbal 49 (IQR 47-62, range 30 [floor value] to 84). Of 20 patients evaluated with autism scales (ADOS/AOSI), 10 (50%) scored as having autism spectrum disorder. On a test of movement ability, 20/21 (95%) scored <1st percentile and 1/21 (5%) >1st percentile. The median Clinical Severity Scale score for the 28 total subjects was 15 (IQR 12.75-17.25).

Although there was a correlation between age and epilepsy and psychiatric dysfunction, there was no overall correlation between gene variant and phenotypic severity, although 2 notable cases were observed at the extremes of severity. A single patient had the maximum severity level (5 of 25), who presented with neonatal hypotonia, infantile spasms and hypsarrhythmia on EEG, status epilepticus at age 14 months, neurologic regression, and daily seizures on 3 antiseizure medicines. The genotype was compound heterozygous c.1294A>C (p.Met432Leu) and c.610-2A>G.

A single patient had a normal IQ measurement and also had the least severity on the global scale (20 of 25). The genotype was a novel variant c.1321G>A (p.Gly441Arg) and a known variant present in more than 5% of the patients: c.612G>A (p.Trp204Ter). This patient had a clinical presentation at 16 months, expressive language deficit, and hypotonia with no seizures and only mild slowing on EEG without epileptiform features.

Additional neurophysiologic testing was done in a subset of subjects using somatosensory evoked responses during both high-density EEG (64 channels) and magnetoencephalography, showing suppression of both the gamma and beta activity bands from tactile stimulation of both upper and lower extremities of patients compared to controls (Figure 6).

Structural neuroimaging demonstrated T2 hyperintensity in the globus pallidi and dentate nuclei, with additional quantification of these and other signal abnormalities and volumetry as discussed in a separate article in this issue. MR spectroscopy measured GABA in voxels corresponding to the posterior cingulate gyrus, basal ganglia, and occipital cortex with consistent elevation in the GABA/NAA ratio in all subjects, having no overlap with controls.

**Discussion**

SSADH deficiency has protean manifestations, both clinically and metabolically, and prior investigations have led to landmark discoveries including development of a genetically modified murine model, identification of pathogenic \( \text{ALDH5A1} \) gene variants, and determination of GABA (A) and (B) receptor dysfunction in humans confirming animal findings. Preclinical and clinical trials have not led to definite therapy for this complex pathophysiology involving alterations in multiple neurotransmitter systems as well as evidence for disrupted autophagy and oxidative metabolism. In order to study the natural history of the disorder and develop biomarkers to better understand its pathophysiology, prognostic determinants, and potential for newborn screening, a natural history study was undertaken by an international collaborative group based at 4 institutions: BCH as the primary clinical site, WSU as the biorepository core, FSU for statistical core, and the iNTD for international patients, with support and outreach from the SSADH Association, the US family advocacy group. Patients were recruited from a registry maintained at BCH and with help from the SSADH Association for the BCH on-site.
comprehensive evaluations, as well as standard-of-care patients unable to travel to this site. The iNTD participating sites, based at the Children’s Hospital Heidelberg, is about to initiate recruitment.

At this half-way point into the natural history study and at the time of this International SSADH Deficiency conference, recruitment goals have been exceeded at BCH with 25 active subjects, along with recruitment of controls, addition of more sophisticated neurophysiologic studies (high-density EEG and magnetoencephalography) in a subset of patients, and enrollees into the standard-of-care cohort. The group’s clinical characteristics are similar to what has been reported from our registry.28 There is a significant relationship between the presence of epilepsy and its severity, and obsessive-compulsive behaviors and psychiatric dysfunction, in the adolescent and adult cohorts compared to the pediatric one (<12 years of age). Although there was no overall correlation between gene variant and severity, there were notable extremes of phenotypic severity in association with variants that may have prognostic value. That is, the missense variant c.1294A>C (p.Met432Leu) in the most severely affected patient was also published in a 23-month-old boy with severe neurologic deficits and a nearly identical progressive course to our patient.29 We are unaware of other cases with this variant. In addition, we had a single patient with a normal full-scale IQ (87) who concomitantly had the highest functioning on the other measures, including the

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**Figure 6.** Time-frequency analysis of SEPs for bipolar electroencephalograph (EEG) channel Cz-C4. (A) TFA for healthy controls (n=10); (B) TFA for patients with SSADHD (n=17). (C) Statistical differences between the 2 groups. Note the significant suppression of signal power in gamma band from ~40 ms till 50 ms and in beta band from ~40 ms till 150 ms. Analysis was performed using nonparametric 2-tailed t test [gamma band: P = .001, corrected for multiple comparisons across normal controls and patients with SSADH over a time span of 70 ms (10-80 ms), t = -4.6074; beta band: P = .001, t = -3.6018 over a time span of 130 ms (10-140 ms)]. TFA, time-frequency analysis.
Movement Assessment Battery and Clinical Severity Score who had a novel missense variant (c.1321G>A, p.Gly441Arg) that we suspect correlates with a milder phenotype.

The SSADHD Natural History Study has provided us with a unique, unprecedented opportunity to study this complex, ultrarare neurometabolic disorder of GABA degradation longitudinally and to collect a biorepository for a wide range of studies as well as correlation with our neurophysiologic and neuroimaging studies. Use of high-density EEG and magnetoencephalography-derived spontaneous and evoked activity demonstrate a decrease in the gamma frequency band in SSADH subjects compared with healthy controls, consistent with impaired GABAergic activity and presenting another biomarker that can be used to study neurophysiologic dysfunction. We propose additional studies on this population, including addition of polysomnography and new imaging sequences that allow for metabolic profiling, to further elucidate the complex mechanisms and potential for intervention in this disorder.

Authors’ Note
Contributing Authors for the SSADH Deficiency Investigators Consortium (SDIC), in addition to the authors of this study: Phillip L. Pearl, Jean-Baptiste Roullet, K. Michael Gibson, Christos Papadelis, Thomas Opladen, Alexander Rotenberg, Kiran Maski, Melissa Tsuboyama, Simon Warfield, Onur Afacan, Edward Yang, Carolyn Hoffman, Kathrin Jeltsch, Jeffrey Krischer, M. Angeles Garcia Cazorla, and Erland Arning.

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Author Contributions
PP contributed to the conception and design of this work; acquisition, analysis, and interpretation of data; and in drafting and revising the manuscript for content.

MD, CP, and EH contributed to the acquisition, analysis, and interpretation of data and participated in drafting and revising the manuscript for content.

TO, JBR, and KMG contributed to the conception and design of this work and in drafting and revising the manuscript for content.

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Ethical Approval
This study was approved by the Boston Children’s Hospital Institutional Review Board (P-00029917). Written informed consent was obtained for all subject participants.

References


