Case Report

Intravenous ganaxolone in pediatric super-refractory status epilepticus: A single hospital experience

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Abstract

Synaptic GABA A receptor (GABA A R) internalization contributes to the drug resistant nature of super-refractory status epilepticus (SRSE). Ganaxolone is a 3β-methylated synthetic analog of the endogenous neuroactive steroid, allopregnanolone, that has positive allosteric modulatory activity on synaptic and extrasynaptic GABA A receptors. Ganaxolone is currently in clinical trials to treat rare pediatric seizure disorders and established and refractory SE.

Two pediatric patients with SRSE (age 17 and age 7) were treated under emergency investigational new drug (E-IND) applications with intravenous (IV) ganaxolone administered as an initial bolus and a maintenance infusion for up to 4.5 days with intermittent IV boluses as-needed followed by taper on day 5 and transitioned to chronic treatment using ganaxolone suspension.

Adjunctive ganaxolone was effective in terminating SRSE in both patients, safely permitting IV anesthetics to be weaned. Seizure control has been maintained after transitioning to enteric ganaxolone. Further investigation of ganaxolone as a safe and effective treatment for SRSE is warranted.

1. Introduction

Super-refractory status epilepticus (SRSE) is a life-threatening neurological emergency and carries a risk for major morbidity and mortality [1–3]. Because of the lack of high-quality comparative clinical studies of SRSE treatment strategies, therapeutic decisions mainly rely on case series or expert opinion [1,4]. In addition to seizure control, the treatment objectives in SRSE include minimization of neuronal injury as well as other organ injury often observed during prolonged treatments with intravenous (IV) anesthesia [5]. Prolonged seizures as well as prolonged GABA stimulation are known to cause internalization of synaptic GABA A receptors (GABA A R). This may play a role in the development of pharmaco-resistance [5,6]. By contrast, extrasynaptic GABA A R have been found to be preserved in SE and identified as a potential target for therapeutic intervention for drug resistant SE [6].

The investigational drug ganaxolone (GNX) is a neuroactive steroid and a positive allosteric modulator of GABA A R that targets a unique binding site distinct from benzodiazepines or barbiturates [7–10]. Ganaxolone is a synthetic derivative of an endogenous neuroactive steroid, allopregnanolone (a metabolite of progesterone), and both exhibit very similar pharmacological properties [8]. GNX acts on both synaptic and extrasynaptic GABA A R to maximize inhibitory signaling as well as maintaining activity when synaptic receptors are downregulated. Intravenous GNX is currently in clinical trials to treat established and refractory SE.

Here we summarize a single institutional experience in treating two pediatric patients with SRSE with IV ganaxolone under emergency investigational new drug (E-IND) applications.

2. Case studies

2.1. Patient #1

A 17-year-old female with a remote history of sporadic seizures in early childhood was transferred to our pediatric hospital for inpatient rehabilitation after 7-month hospitalization for recurrent...
refractory SE (as defined by ILAE criteria) [11] at an outside facility. Over the course of 7 months at the initial tertiary hospital, she had 6 separate episodes of SE and required intubation 4 times with medically induced coma for seizure suppression. She was discharged to our inpatient rehabilitation unit on 5 maintenance anti-seizure medications (ASMs): cannabidiol, perampanel, phenobarbital, lacosamide, and lorazepam; along with pyridoxine, a ketogenic diet, anakinra, and menstrual suppression. A thorough infectious, metabolic, genetic, vascular, and autoimmune evaluation was nondiagnostic (Table 1a).

While in the inpatient rehabilitation unit, the patient developed a fever of 41.7 °C, and a respiratory viral panel was positive for parainfluenza. Recurrence of SE required transfer to the intensive care unit for midazolam and pentobarbital infusions. In addition to the prior antiseizure medications and therapies, enteric felbamate was introduced at this time. Video-electroencephalogram (vEEG) monitoring throughout the course of status epilepticus demonstrated multifocal seizure onset. She was seizure-free for 2 days after infusions were weaned, until seizures returned, prompting placement of a vagus nerve stimulator and a prolonged 2-week course of IV pentobarbital titrated to EEG burst suppression, leading to the prior antiseizure medications and therapies, enteric felbamate being discontinued; by day 3, clinical and electrographic seizures stopped (Fig. 2). The patient tolerated the protocol well. As pentobarbital was discontinued and GNX treatment initiated, she became more alert and interactive. On day 5 she was transitioned from IV to enteric GNX (maximal adult dose of 1800 mg/day divided three times a day) without recurrent seizures. During the introduction of ganaxolone she was maintained on the same anti-seizure regimen and ketogenic diet though was weaned off phenobarbital.

2.2. Patient #1 disposition and update

One month after GNX initiation, the patient was transferred to inpatient rehabilitation service and was discharged home one month later. While in the hospital she was weaned off the ketogenic diet and anakinra, and as an outpatient she has been weaned off felbamate with minimal breakthrough clinical seizures (primarily in the setting of intercurrent illness or decreasing ASMs). More than eighteen months later, the patient’s seizure control has been maintained as she continues to be on four ASMs (perampanel, brivaracetam, lorazepam and valproic acid—the latter two at lower doses) and GNX. To note, the patient remains with significant morbidity, with gastrostomy tube and tracheostomy dependence, requiring assistance with all activities of daily living.

2.3. Patient #2

A 7-year-old, previously healthy female presented with abdominal pain, encephalopathy and fever and developed SRSE concordant with febrile infection-related epilepsy syndrome (FIRES). The patient was admitted to PICU and placed on vEEG which demonstrated multifocal independent electrographic seizures arising from posterior head regions. Upon admission, MRI demonstrated FLAIR changes in bilateral temporal lobes consistent with posterior head regions. After two failed attempts to wean IV anesthetics, she was treated with IV immunomodulatory therapies with IV methylprednisolone and IVIG were initiated concomitantly in the ICU. After two failed attempts to wean IV anesthetics, she was treated with IV-to-enteric GNX approved under E-IND.

Table 1a

| Infectious | Autoimmune/inflammatory | Metabolic |
|------------|-------------------------|----------|
| B cell panel: negative | CSF 4/2/20: 1 cell, 57 protein, 71 glucose | Serum carbohydrate deficient transferrin: normal and inconsistent with a CDG |
| HIV and hepatitis: negative | Autoimmune encephalopathy panel: negative | Serum biotinidase level, 13.8 U/L (within normal limits) |
| COVID Ag testing <3: negative | Oligoclonal band panel: negative | Homocysteine: low < 3.5 |
| NMO: negative | Urine purine and pyrimidine panel: normal | Urine sulfocysteine: normal |
| | | Urine mucopolysaccharides screen: elevated heparan sulfate, 0.33 |
| | | Serum carbohydrate deficient transferrin: inconsistent with CDG |
| | | Acylcarnitine profile: normal |
| Imaging | Free and total L-carnitine: plasma level normal | Free and total L-carnitine: plasma level normal |
| Normal unenhanced and enhanced MRI of the brain (<5) Negative MRA of the head | Urine organic acids: normal | Serum organic acids: normal |
| Epilepsy panel with 5 VUS: PCDH19, POLG, DOCK7, NRXN1, and PLCB1 | Plasma amino acids: normal | Plasma amino acids: normal |
| c.39-5T>G DOCK7 gene: father; X mother | CSF metabolic studies: negative | CSF metabolic studies: negative |
| c.2299A>G p.Met767Val NRXN1 gene: father; X mother | | |
| c.199A>C p.Ser67Arg PLCB1 gene: father; X mother | | |
| c.545C>G p.Gly182Arg PLCD19 gene: mother; X father | | |
| c.3131T>C p.Val1044Ala POLG gene: father; X mother | | |
| All are responsible for autosomal recessive disorders, and a second change was not found | | |
| c.545C>G p.Gly182Arg PLCD19 gene: mother; X father | | |
| Mother asymptomatic; uncertain significance | | |
| Whole exome sequencing with mitochondrial genome (Trio): negative | | |

Ag, antigen; CDG, congenital disorders of glycosylation; COVID, coronavirus disease; CSF, cerebrospinal fluid; HLH, hemophagocytic lymphohistiocytosis; MRA, magnetic resonance angiography; MRI, magnetic resonance imaging; NMO, neuromyelitis optica; US, ultrasound.

* GeneDx, Inc., Gaithersburg, MD.
As anesthesia was weaned on a third attempt and seizures returned while on the same antiseizure medication regimen, GNX was initiated on hospital day 17 using an IV bolus, followed by infusion over 4.5 days (with boluses as needed for breakthrough seizures) followed by a taper on day 5 and transition to enteric maintenance GNX. On day 1, pentobarbital was discontinued; by day 3, clinical and electrographic seizures stopped.

2.4. Patient #2 disposition and update

At discharge, the patient remained on several ASMs (levetiracetam, lacosamide, perampanel, phenobarbital, lorazepam) ganaxolone (63 mg/kg/day divided three times a day), anakinra, and tetrabenazine. Note, tetrabenazine was introduced for her transient hyperkinetic movement disorder which became more apparent as she became more active and awake. MRI throughout the inpatient course demonstrated FLAIR changes in the basal ganglia explaining these symptoms. By 8 months from discharge, she was weaned off perampanel and tetrabenazine as the movement disorder resolved. More than one year after initial presentation, she continues on levetiracetam, lacosamide, phenobarbital, lorazepam, anakinra (the latter both at lower doses) and ganaxolone in addition to iron and pregabalin with rare breakthrough seizures in the setting of intercurrent illness or attempts at medication.
weans. Cognition remains age-appropriate without any physical limitations. She also attends school with modifications.

3. Discussion

3.1. Pathophysiology

The International League Against Epilepsy (ILAE) has defined SE as a condition of prolonged seizure activity resulting from either the failure of mechanisms responsible for seizure termination (mainly through GABA<sub>A</sub> receptors) or the initiation of mechanisms which lead to enhanced excitation (mainly through NMDA/AMPA/KA receptors) [11]. Refractory SE is defined as SE resistant to treatment with first-line ASM (benzodiazepines) and one second-line IV ASM such as fosphenytoin/phenytoin, valproic acid or levetiracetam [12,13]. SRSE is defined as SE that fails to respond or recurs after a trial of a third-line agent (IV anesthesia). Overall, one-third of patients with SRSE die and another one-third recover with chronic neurologic deficits [14,15].

3.2. Neurosteroids: targeting extrasynaptic GABA<sub>A</sub> receptors

Neurosteroids increase tonic inhibition via their allosteric modulation of both synaptic and extrasynaptic GABA<sub>A</sub>R [16]. While synaptic GABA<sub>A</sub>R (a target for benzodiazepines and barbiturates) are progressively internalized with prolonged seizures leading to pharmacoresistance [6], the function of extrasynaptic GABA<sub>A</sub>R is preserved [16].

A recent randomized controlled trial in adults and children failed to demonstrate the efficacy of IV allopregnanolone (brexanolone) compared to placebo in the resolution of SRSE, when it
was added to the standard care [17]. However, novel dosing regimens with ganaxolone described here offers a new opportunity to these highly refractory and medically complex patients. Ganaxolone is a proprietary IV solution solubilized by Captisol® (betadex sullobutyl ether sodium). The dosing targets concentrations of ganaxolone that are expected to demonstrate anticonvulsant properties [18]. In preclinical models, GNX has exhibited broad-spectrum antiseizure properties, including when benzodiazepine

| Infectious                          | Autoimmune/inflammatory                                                                 | Metabolic                                                                 |
|------------------------------------|-----------------------------------------------------------------------------------------|---------------------------------------------------------------------------|
| CSF and Blood cultures: negative   | CSF 12/30/20: 4 WBC, 49 protein, 71 glucose                                              | Vit. B12, TSH/FT4, folate, copper, ceruloplasmin: Normal                   |
| HSV PCR CSF: negative              | Autoimmune encephalopathy panel: negative (serum and CSF)                               | Acylcarnitine profile: normal                                             |
| CSF and Serum Arbovirus            | Pelvic US: negative for teratoma                                                        | Urine organic acids: normal                                               |
| CSF EBV, VZV PCR: negative         | CSF oligoclonal bands: negative                                                        | Plasma amino acids: normal                                                |
| CSF RPR and VDRL (CSF) negative    | NMO antibodies: negative                                                               | Heavy metals: normal                                                     |
| Bartonella serology and PCR:       | ANA, dsDNA, ACE                                                                        | Porphyria: negative                                                      |
| Negative                           | Lysozyme                                                                                | Very long chain fatty acids: normal                                       |
| Mycoplasma: Negative               | CSF cytokines showed no elevation and serum cytokines showed elevation of IL-6.         |                                                                           |
| COVID Ag and Ab testing negative   |                                                                                         |                                                                           |

| Imaging                            |                                                                                         | Genetic                                                                 |
|------------------------------------|-----------------------------------------------------------------------------------------|---------------------------------------------------------------------------|
| MRI Brain: Abnormal (*)            |                                                                                         | Epilepsy panel: negative                                                 |
| Negative MRA/MRV of the head and neck |                                                                                      | Whole exome sequencing (TRIO): negative                                  |

* MRI brain demonstrated FLAIR changes in bilateral temporal lobes consistent with prolonged seizures.

Table 1b
Diagnostic studies and evaluations for Patient # 2.

a. One day prior to Ganaxolone Initiation, Patient # 2. Double Banana Bipolar Montage

b. Day 8 after of Ganaxolone Initiation, Patient # 2. Double Banana Bipolar Montage

Double Banana Bipolar Montage, LFF 1 Hz, HFF 70 Hz, Sensitivity 10 uV/mm, Timebase 30 mm/sec

a. This EEG demonstrates rhythmic BIPL EDS (periodic lateralized epileptiform discharges) in the bipo
terior regions which subsequently evolve to electrographic seizure upon attempts to wean ketamine infusion prior
to initiation of GNX.

b. This interictal EEG demonstrates resolutions of seizures after 8 days of protocol once transitioned to enteric
ganaxolone and ketamine and midazolam infusions discontinued, with improvement in interictal background, less
epileptiform discharges, and reactivity.

Fig. 3. EEG prior to and after ganaxolone initiation for Patient # 2. One day prior to Ganaxolone Initiation, Patient # 2. Anterior-posterior longitudinal bipolar montage (a.k.a. “Double Banana”) Day 8 after Ganaxolone initiation, Patient # 2. Double Banana Bipolar Montage.
resistance developed [8, 10]. In an open-label, Phase 2 study with IV ganaxolone in refractory SE, ganaxolone achieved rapid and durable seizure control in patients with RSE, and showed acceptable safety and tolerability [9]. This study provided the rationale for specific IV dosing paradigm for SRSE [19,20] as well as for further investigation of ganaxolone in an ongoing, phase 3 trial in refractory SE (NCT04391569). Specifically, infusion parameters were optimized to allow for rapid loading of ganaxolone (plasma concentration ~900 ng/ml) designed to abort SE (median time to SE cessation following ganaxolone initiation was 5 min), followed by maintenance doses aimed at sustaining seizure control (no patient required escalation to third-line IV anesthetics during the 24-h period following ganaxolone initiation) (for details see [19]). Of note, target serum ganaxolone levels achieved by the current IV dosing paradigm were 5–10 times higher as compared to serum brexanolone levels in the STATUS trial [9, 21]. Ganaxolone’s properties and mechanism of action along with the availability of both IV and oral formulations allows reaching adult and pediatric patients in acute and chronic care settings [22].

3.3. Patient-specific discussion

Although both patients in our series developed SRSE, their prior histories and courses were quite different.

Our first patient had a history of isolated early childhood seizures but was off daily antiseizure medication for more than one decade when she initially developed SRSE as a teenager. Although her course was similar to those who develop NORSE (New-onset refractory status epilepticus), she did not meet the traditional criterion or definition of de novo onset of seizures. In contrary our second patient had no history of seizures, and her presentation was most consistent with FIRES (a subset of NORSE) with the clinical presentation characterized by de novo onset of RSE or SRSE without a clearly identifiable acute or active structural, toxic, or metabolic cause [23, 24]. Regardless of cause or etiology, mortality estimates in pediatric RSE range from 14 to 44% [3].

In SRSE, traditional antiseizure management has poor success rates, especially during the acute phase [5, 23, 24]. The majority of patients require anesthetics during the acute phase and prolonged burst-suppression coma is often necessary. At least one-third of patients require multiple anesthetics drugs to achieve seizure control. The challenge is that seizures may recur upon attempts to wean off the continuous infusions. Exact pathophysiology remains unclear, though emerging evidence suggests a role for a postinfectious or inflammatory cytokine-mediated mechanism [25]. Though treatment protocols vary among institutions, most often utilize other available nonpharmacologic therapies: immunomodulatory therapy, ketogenic diet, therapeutic hypothermia, electroconvulsive therapy, vagus nerve stimulator, each of which has varying efficacy, though without substantial change in the overall morbidity or mortality [5, 25]. The treatments for SRSE are guided by preclinical studies and retrospective case reviews as there are no published randomized control studies to guide clinicians.

Over the recent years both the ketogenic diet and novel immunomodulatory therapies are often utilized in unknown or presumed autoimmune causes of SRSE [5]. The evidence to use these agents to treat SRSE is based on preclinical and observational studies. Modulation of proinflammatory cytokines with anakinra (IL-1 receptor antagonist) or tocilizumab (IL-6 receptor antagonist) is being increasingly utilized in treatment of RSE, SRSE and FIRES [5, 26, 27]. Ketogenic diet has also gained significant attention in recent years for the treatment of SE as an adjunctive therapy due to its anti-inflammatory and anti-seizure properties and has been effective in FIRES [28, 29]. To note, our first patient relapsed back into SRSE while on therapeutic doses of anakinra and the ketogenic diet. Our second patient was initiated on the ketogenic diet briefly but did not ever achieve ketosis beyond trace ketones in the acute ICU stay and thus was weaned off at parents’ request once seizures became under control.

Although the exact cause of SE in each of our patients remains unknown despite comprehensive diagnostic evaluations ruling out acute and remote symptomatic causes, such is often the case in pediatric SRSE. Time to achieve seizure cessation for each patient varied again reflecting the heterogenous nature and course of patients with SRSE: Patient 1 was on a single continuous IV infusion and achieved seizure control after 5 days of Ganaxolone initiation. However, Patient 2 was on two continuous IV infusions and took longer to achieve seizure control as both infusions had to be weaned (8 days). Each patient had an extremely complicated course with multiple concurrent medication changes that could be confounding. Despite that, ganaxolone’s mechanism of action and preclinical data was suggestive of its potential role in treatment of SRSE in our patients.

4. Conclusions

Despite recent advances in novel therapies for the treatment of pediatric SRSE beyond IV anesthetics, morbidity and mortality especially in the acute stages remains high. The patient’s experience in this series reflects the heterogenous nature of various treatment approaches for SRSE which often employs a trial-and-error manner until treatment response is achieved [5]. Ganaxolone appeared to contribute to termination of SRSE in two pediatric patients at our institution, permitting IV anesthetics to be weaned and ultimately discharge from the hospital. Long-term seizure control has been maintained in both patients after transitioning to adjunctive oral ganaxolone excepting rare breakthrough seizures.

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Marinus Pharmaceuticals, Inc. provided IV and oral ganaxolone. No monetary support was provided.

Ethical statement

Each patients E-IND trial was approved by the FDA and was conducted according to the guidelines of the Declaration of Helsinki and approved by the Institutional Review Board of Atrium Health System.

Declaration of Competing Interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: Rani K. Singh – Reports research funding from the Pediatric Epilepsy Research Foundation, has served on the advisory board for Zogenix, Prasco, and AKPharma, and is a member of the executive board of the Western Region of the Epilepsy Foundation of North Carolina. Joseph Huihui, Heather van Heusen, Henrikas Vaitkevicius, Maciej Gasior are all employees of Marinus Pharmaceuticals, Inc.

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