Data Related to Disease Similarity--A Case Study: PEACE Initiative in Pediatric Epilepsy

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In disease states affecting both adults and children, drugs are often approved for adult use before development in children is completed or even started.

Although antiepileptic drugs (AEDs) approved for use in adults can be prescribed off-label for children, this availability hampers pediatric drug development:

- Raises parents’ concerns about enrolling children with refractory epilepsy in a trial of a marketed drug with placebo.
- Creates an operational disincentive to undertake the challenges of conducting double-blind, randomized, controlled efficacy trials in children with seizures.
In the absence of pediatric-specific labeling, prescribers lack critical information (e.g., dosing, tolerability/safety, age-specific monitoring) that can facilitate the appropriate and safe use of AEDs in this vulnerable population.

Expediting pediatric access to new AEDs is compelling since epilepsy is the most common serious neurological disorder in children.

Almost none of the AEDs approved for the management of focal (partial-onset) seizures included children <12 years of age in the initial clinical development program and were therefore marketed for use in older adolescents and adults.
• Since focal seizures occur in both children and adults, efficacy data from adults can be successfully extrapolated to children if there is scientific consensus that disease progression and response to intervention are similar in adults and children.

• The focus of an argument for adult-to-pediatric efficacy data extrapolation in focal seizures is based on the similarity of seizure pathophysiology and the similarity of the clinical response to AEDs in terms of seizure control.
PREVALENCE OF GENERALIZED AND PARTIAL SEIZURES

Hauser, 1992

Generalized seizures are more common in the first five years of life, the incidence was similar for both between the ages of 6 and 24, and partial seizures were at least twice as common as generalized onset seizure in adults over 24 years.

Pediatric Patients <15 Years
- Other partial: 7%
- Tonic-clonic: 23%
- Other generalized: 11%
- Complex partial: 49%
- Unknown/multiple: 3%
- Other partial: 6%
- Myoclonic: 7%
- Simple partial: 11%

Adults 35-64 Years
- Other partial: 3%
- Tonic-clonic: 27%
- Other generalized: 11%
- Complex partial: 49%
- Simple partial: 13%
- Myoclonic: 2%
- Unknown/multiple: 3%
PEACE Rational

• Because anti-epileptic drugs are not evaluated as disease-modifying drugs (i.e. not anti-epileptogenic) the focus of an argument for adult-to-pediatric efficacy data extrapolation is based on the similarity of seizure pathophysiology.....
  – Key factors in E/I balance as function of age
  – Network maturation
  – Neurophysiological maturation
  – Seizure and EEG semiology

• and the similarity of the clinical response to AEDs in terms of seizure control.
90% of a child's brain development happens before age 5.
My Hobby: Extrapolating

As you can see, by late next month you'll have over four dozen husbands.

Hold on - shouldn't you be using more than two data points?

Number of Husbands

Ooops, you're right.
Developmental Aspects of Receptor Development

**Newborn**
- AMPA: Silent
- NMDA: Depol.
- GABA_A: Silent
- GABA_B: Delayed
- KCC2: K^+ / Cl^-

**Adult**
- AMPA: Depol.
- NMDA: Hyperpol.
- GABA_A: K^+ / Na^+
- GABA_B: K^+ / Na^+ + Ca^{2+}
- KCC2: K^+ / Cl^-
GABA is Excitatory in the Neonatal Brain

- Cl⁻ taken up by NKCC1
- Overwhelms Cl⁻ extrusion by KCC2
- High intracellular [Cl⁻]
- +ve Cl⁻ equilibrium pot.
- GABA_A activation results in Cl⁻ efflux & depolarization
- Depolarization causes glutamate release, further excitation via GLU-R
Bumetanide Potentiates the Effect of Phenobarbital

- Bumetanide selectively inhibits NKCC1
- Reduces intracellular [Cl\(^-\)]
- Reversal of Cl\(^-\) equilibrium potential
- GABA\(_A\) activation causes Cl\(^-\) influx & hyperpolarization
- Hyperpolarization prevents excitation, glutamate release
Neuronal Receptor Expression vs Age

- GABA (depolarizing)
- GABA (inhibitory)
- NMDA
- AMPA
- Kainate

% Adult Function

Rodent Human

P0 20 PCW
P5 30 PCW
P10 40 PCW
P15 1-2 years
P20 >3 years
P25
P30
Adult

Jensen and Silverstein, 2007
Theta Rhythm

Rats: 4-12Hz

Time-locked to lever pressing behavior (Sample & Match presses)

Humans: 4-8Hz

1 second
Single Unit Recording Method

Rate Coding

![Graphs illustrating rate coding with time and distance metrics.](image)
Beyond Rate Coding – Temporal Coding of AP’s by the Phase of Theta
Replay of Place Cells in Sleep

Matt Wilson

O’Neill, Trends in Neurosciences, 2010

Diekelmann and Born, Nature Neuroscience Reviews, 2010

modified from Ji and Wilson, Nature Neuroscience, 2007
Maturational Dynamics of Place Cells in Immature Rats

- Ontogeny of the place cell system in rats studied between P22 and P43, a time during which there was a rapid improvement in spatial behavior.
- Place cells with adult like firing fields were observed at the earliest ages, but were few in number.
- Firing rate and stability increased with age and the average spatial signal of all pyramidal cells improved.
Alpha Frequency

![Graph showing Alpha Frequency across different ages]

- **4 yo**
- **16 yo**
The Ontogeny of Partial Seizures in Infants and Young Children

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Summary: Purpose: To describe the clinical manifestations of partial seizures in the pediatric population as a function of age.

Methods: Using the database of the pediatric epilepsy monitoring unit (Children’s Hospital of New York), clinical and EEG characteristics of partial seizures were distributed by age groups 0–2, 2–6, and 6+ years for 123 patients who had at least one such seizure with a clear EEG correlate during their admission. χ² tests for trend were used to examine clinical and EEG features as a function of age.

Results: The frequency of aura, limb automatism, dystonic posturing, secondary generalization, and unresponsiveness increased with age, whereas asymmetric clonus and symmetric tonic posturing decreased with age. There were no clear changes in the types of EEG ictal patterns observed with age; however, partial seizures emanating from the anterior regions of the brain tended to increase with age, whereas those from the posterior regions tended to decrease with age.

Conclusions: Important differences exist in the clinical expression of seizures between young children and adults. These findings will contribute to a better understanding of ictal ontogeny that will promote more accurate classification of seizures and of the epilepsies in young patients. Such efforts can be used to identify young patients for focal epilepsy surgery and to select appropriate anticonvulsive medications. Key Words: Epilepsy—Seizures—Infants—Age-related features.
Seizure Characteristics that Change with Age

Nordli et al., 2001
## EEG Parameters by Age

<table>
<thead>
<tr>
<th>Location</th>
<th>0–2 (48)</th>
<th>2–6 (21)</th>
<th>6+ (54)</th>
<th>Total (123)</th>
<th>$\chi^2_{\text{trend}}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Frontal</td>
<td>8 (17)</td>
<td>5 (24)</td>
<td>14 (26)</td>
<td>27 (22)</td>
<td>1.3</td>
</tr>
<tr>
<td>Anterior temporal</td>
<td>4 (8)</td>
<td>3 (14)</td>
<td>8 (15)</td>
<td>15 (12)</td>
<td>1.0</td>
</tr>
<tr>
<td>Rolandic</td>
<td>8 (17)</td>
<td>2 (10)</td>
<td>3 (6)</td>
<td>13 (11)</td>
<td>3.3</td>
</tr>
<tr>
<td>Occipital</td>
<td>10 (21)</td>
<td>1 (5)</td>
<td>1 (2)</td>
<td>12 (10)</td>
<td>10.8&lt;sup&gt;a&lt;/sup&gt;</td>
</tr>
<tr>
<td>Temporoparietal</td>
<td>14 (29)</td>
<td>5 (24)</td>
<td>10 (19)</td>
<td>29 (24)</td>
<td>1.6</td>
</tr>
<tr>
<td>Hemispheric</td>
<td>3 (6)</td>
<td>3 (14)</td>
<td>8 (15)</td>
<td>14 (13)</td>
<td>1.8</td>
</tr>
<tr>
<td>Vertex</td>
<td>0</td>
<td>0</td>
<td>4 (7)</td>
<td>4 (3)</td>
<td>2.7</td>
</tr>
<tr>
<td>Feature</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Isolated spike</td>
<td>1 (2)</td>
<td>0</td>
<td>1 (2)</td>
<td>2 (2)</td>
<td>0</td>
</tr>
<tr>
<td>Rhythmic delta</td>
<td>14 (29)</td>
<td>4 (19)</td>
<td>12 (22)</td>
<td>30 (24)</td>
<td>0.6</td>
</tr>
<tr>
<td>Rhythmic theta/alpha</td>
<td>25 (52)</td>
<td>15 (71)</td>
<td>27 (50)</td>
<td>67 (55)</td>
<td>0.1</td>
</tr>
<tr>
<td>Run (sharps/spikes)</td>
<td>7 (15)</td>
<td>3 (14)</td>
<td>5 (9)</td>
<td>15 (12)</td>
<td>0.7</td>
</tr>
<tr>
<td>Attenuation</td>
<td>1 (2)</td>
<td>0</td>
<td>1 (2)</td>
<td>2 (2)</td>
<td>0</td>
</tr>
<tr>
<td>Low-voltage fast</td>
<td>5 (10)</td>
<td>0</td>
<td>9 (17)</td>
<td>14 (13)</td>
<td>1.0</td>
</tr>
<tr>
<td>Irregular slowing</td>
<td>1 (2)</td>
<td>0</td>
<td>0</td>
<td>1 (1)</td>
<td>0.2</td>
</tr>
</tbody>
</table>

Nordli et al., 2001
Efficacy of antiepileptic drugs in adults predicts efficacy in children
A systematic review

ABSTRACT

Objective: Due to the challenges inherent in performing clinical trials in children, a systematic review of published clinical trials was performed to determine whether the efficacy of antiepileptic drugs (AEDs) in adults can be used to predict the efficacy of AEDs in the pediatric population.

Methods: Medline/PubMed, EMBASE, and Cochrane library searches (1970–January 2010) were conducted for clinical trials of partial-onset seizures (POS) and primary generalized tonic-clonic seizures (PGTCS) in adults and in children <2 and 2–18 years. Independent epidemiologists used standardized search and study evaluation criteria to select eligible trials. Forest plots were used to investigate the relative strength of placebo-subtracted effect measures.

Results: Among 30 adjunctive therapy POS trials in adults and children (2–18 years) that met evaluation criteria, effect measures were consistent between adults and children for gabapentin, lamotrigine, levetiracetam, oxcarbazepine, and topiramate. Placebo-subtracted median percent seizure reduction between baseline and treatment periods (ranging from 7.0% to 58.6% in adults and from 10.5% to 31.2% in children) was significant for 40/46 and 6/6 of the treatment groups studied. The ≥50% responder rate (ranging from 2.0% to 43.0% in adults and from 3.0% to 26.0% in children) was significant for 37/43 and 5/8 treatment groups. In children <2 years, an insufficient number of trials were eligible for analysis.

Conclusions: This systematic review supports the extrapolation of efficacy results in adults to predict a similar adjunctive treatment response in 2- to 18-year-old children with POS.
Efficacy Comparison of Differences in Median % Seizure Reduction Between the Baseline and Treatment Periods by Drug for Children and Adults

Pellock et al, 2012
Placebo-subtracted median percent seizure reduction between baseline and treatment periods (ranging from 7.0% to 58.6% in adults and from 10.5% to 31.2% in children) was significant for 40/46 and 6/6 of the treatment groups studied. The ≥50% responder rate (ranging from 2.0% to 43.0% in adults and from 3.0% to 26.0% in children) was significant for 37/43 and 5/8 treatment groups.
Summary

• Based on anatomical and physiological data, from a focal seizure standpoint, physiological function of the brain of a 4 year old is similar to that of an adolescent.

• By age 4 years the EEG is quite similar to that of an adolescent and adult.

• The clinical semiology and EEG features of focal seizures in a 4 year old does not differ substantially from adolescents and adults.

• The response to AEDs in focal seizures do not differ in children older than 4 years of age than adults.