A review of the clinical spectrum of BRAT1 disorders and case of developmental and epileptic encephalopathy surviving into adulthood

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Keywords: BRAT1 Epilepsy Lethal neonatal rigidity and multifocal seizure syndrome Epileptic encephalopathy Adult

1. Introduction

Sporadic developmental and epileptic encephalopathies (DEE) are most commonly attributable to monoallelic variants arising de novo but can be transmitted with recessive and x-linked variants. Pathogenic biallelic variants in BRAT1 are associated with a spectrum of disease including Lethal Neonatal Rigidity and Multifocal Seizure Syndrome (RMFSL, OMIM: 614498), Epilepsy of Infancy with Migrating Focal Seizures (EIMFS), and Neurodevelopmental Disorder with Cerebellar Atrophy and with or without Seizures (NEDCAS). RMFSL is characterized by early-onset multifocal seizures with microcephaly. Death occurs during infancy although a less severe course with later onset seizures and longer survival into childhood has been described. Here, we summarize published cases of BRAT1 disorders and present the case of a 20-year-old man with two heterozygous BRAT1 variants and a relatively later age of seizure onset with survival into adulthood. This case expands the spectrum of disease associated with BRAT1 variants and highlights the utility of genetic testing to identify the cause of developmental and epileptic encephalopathies where clinical heterogeneity within a spectrum of disease exists.

2. Case report

A 20-year-old man with non-consanguineous parents was born by caesarean section at 35 weeks due to severe intrauterine growth restriction. His mother and father did not have any history of epilepsy, neurological disorders, or any other medical problems.
| Case ID | BRAT1 Variant | Age at onset of seizures | Seizure type | Syndrome | MRI head findings | EEG findings | Survival  |
|---------|---------------|--------------------------|--------------|----------|------------------|-------------|-----------|
| Puffenberger, 2012 [2] | Homozygous c.638_639dupA, p. (Val214Glyfs*189) | Shortly after birth | Facial and arm jerks. | RMFSL | Normal or mild frontal hypoplasia | Bilateral medium–high voltage spikes over temporal and central regions, frequent multifocal seizures, background slowing, and no posterior rhythm. | < 4 months |
| Saunders, 2012 [3] | Homozygous c.453_454insATCTTCTC, p. (Leu152Ilefs*70) | Day 1 | NA | RMFSL | Normal | Focal epileptiform and sharp wave activity. | NK |
| Saiitsu, 2014 [21] | Compound heterozygous c.176 T > C, p.(Leu59Pro); c.962_963delTC, p.(Leu321Profs*81) | Day 1 | Myoclonic | RMFSL | Normal | Sharp waves and bilateral spikes predominantly over the right hemisphere. | 5 months |
| Straussberg, 2015 [4] | Homozygous c.1177delG, p. (Ala393Leufs*1) | Day 1 | Myoclonic | RMFSL | Normal | Bilateral epileptic activity with bilateral discharges. | 6 months |
| Van de Pol, 2015 [22] | NA | 1st month | Myoclonic | RMFSL | Normal | Severe abnormal background, multifocal sharp waves, and frequent multifocal epileptic seizure activity. | 3 months |
| Hanes, 2015 [5] | Compound heterozygous c.294dupA, p. (Leu99Thrfs*92); c.1825 > T, p.(Arg609Trp) | 5 months | Focal with impaired awareness | NEDCAS | Cerebellar atrophy | Slow awake background. No epileptiform discharges. | Alive at 4 years |
| Fernández-Jaén, 2016 [23] | Compound heterozygous c.1564G > A, p.(Glu522Lys); c.638dupA, p. (Val214Glyfs*189) | None | None | NEDCAS | Cerebellar atrophy | Normal | Alive at 4.5 years |
| Horn, 2016 [6] | Compound heterozygous c.638dupA, p. (Val214Glyfs*189); c.1134 + 1G > A | 5 months | Myoclonic | RMFSL | Thin corpus callosum, dilated lateral ventricles, delayed myelination. | Focus continuous spike discharges in the right more than in the left occipital region | 5 years and 9 months |
| Mundy, 2016 [7] | Compound heterozygous c.294dupA, p.(Leu99Thrfs*92); c.1925 > T, p.(Ala642Glu) | 3 months | NA | RMFSL | Thin corpus callosum, cerebellar hypoplasia. | Left- sided temporo-occipital epileptiform discharges and absence of a posterior dominant rhythm. | Alive at 6 years |
| Smith, 2016 [17] | Compound heterozygous c.1857G > A, p.(Trp619*); c.2125_2128delTTTG, p.(Phe709Thrfs*17) | 4 months | Myoclonic, focal with impaired awareness, focal to bilateral tonic-clonic. | RMFSL | Global cerebral and cerebellar atrophy | Bilateral multifocal epileptiform activity | 15 months |
| Srivastava, 2016 [8] | Compound heterozygous c.638dupA, p. (Val214Glyfs*289); c.803 + 1G > C | No seizures reported | No seizures reported | RMFSL | Normal | Bilateral multifocal epileptiform activity | Alive at 4 years and 4 months | Alive at 10 years |
| Celik, 2017 [24] | Homozygous c.2230_2237dupAACACTGC, p. (Ser747Thrfs*36) | 1st month | Tonic, focal status epilepticus | RMFSL | Mild cerebral atrophy | Generalized and focal bi–posterior quadrant slowing, and multifocal epileptiform activity. Background activity of ~ 6 Hz theta, bilateral frontotemporal sharp waves, and 8–10 Hz ictal rhythm during clinical seizures. | Alive at 15 months |
| Case ID | BRAT1 Variant | Age at onset of seizures | Seizure type | Syndrome | MRI head findings | EEG findings | Survival |
|---------|----------------|--------------------------|--------------|----------|------------------|-------------|----------|
| Hegde, 2017 [25] | Homozygous c.617 T > A, p.(Leu206)^* | Day 3 | Clonic | RMFSL | Cortical and cerebellar atrophy | Occasional generalized bursts of epileptiform activity with relatively well-preserved background activity, burst suppression pattern | 4 months |
| Skafi, 2018 [26] | Homozygous c.638,639dupA, p.(Val214Glyfs*189) | Day 1 | Myoclonic | RMFSL | Normal | Low-voltage background without epileptic discharges. | 3 months |
| Szymańska, 2018 [16] | Homozygous c.1313_1314delAG, p. (Gln438Argfs*51) | Day 1 | Myoclonic | RMFSL | Cerebral atrophy, thinning of corpus callosum | NA | 6 months |
| Van Ommeren, 2018 [27] | Homozygous c.1395G > C, p.(Thr465Thr) | Day 1 | Myoclonic | RMFSL | Widened subarachnoid space | Generalized and focal sharp and spike waves. | 12 months |
| Mahjoub, 2019 [28] | Homozygous c.1395G > C, p.(Thr465Thr) | No seizures reported | None | NEDCAS | Cerebellar atrophy | NA | 2 months |
| Colak, 2020 [29] | Homozygous c.1395G > C, p.(Thr465Thr) | No seizures reported | None | NEDCAS | Cerebellar atrophy | Generalized epileptiform activity, migrating focal epileptiform activity, background deceleration. | Alive at 7 years |
| Pourahmadiyan, 2020 [30] | Homozygous c.2041G > T, p.(E681^*) | NK | Focal seizures | RMFSL | NA | NA | 6 days |
| Scheffer, 2020 [10] | Compound heterozygous c.964C > T, p.(Gln322^*); c.2284C > T, p. (Gln762)^* | Day 1 | Focal motor migrating between hemispheres | EIMFS | Underopercularization of the Sylvian fissure | Multifocal epileptiform discharges, discontinuous background lctal: focal seizure migrating from left central region to right hemisphere. | 1 month |
| 2 | Day 1 | Focal clonic seizures with apnea and tachycardia migrating between hemispheres | EIMFS | Thinning of corpus callosum | Multifocal epileptiform discharges, discontinuous background intermittently lctal: migrating focal seizures; central, right occipital spread to left occipital region, left temporal spread to left hemisphere then right hemisphere, bi-occipital onset. | 10 months |
| 3 | Day 1 | Focal, multifocal motor seizures with clonic features, apnea, eye deviation epileptic spasms; tonic seizures | EIMFS | White matter volume loss | Multifocal epileptiform discharges lctal: migrating focal seizures; seizures arising from right central region, left occipital, right temporal, and left temporal region. Epileptic spasms and periodic spasms, hypsarrhythmia | 4 years and 3 months |
| 4 | Day 4 | Focal motor clonic seizures, migrating between hemispheres | EIMFS | Subdural hemorrhage | Multifocal epileptiform discharges, discontinuous background lctal: myoclonic seizures, clonic seizures, facial clonic movements, with migration from right posterior occipital region to left posterior region. | 2 months |
| 5 | Day 1 | Focal seizures migrating between hemispheres | EIMFS | Subdural hemorrhage | Multifocal epileptiform discharges, discontinuous background lctal: migrating focal seizures from one region to another, most frequent onset from right posterior quadrant; other onsets in left posterior region and left fronto-central region. | 14 months |
| Balasundaram, 2021 [31] | Biallelic deletion of at least exons 1–2 | Day 1 | Tonic-clonic, myoclonic | RMFSL | Corpus callosum thinning | Bilateral medium–high voltage spikes over temporal and central regions, frequent multifocal seizures, background slowing, and no posterior rhythm. | 2.5 months |
| Li, 2021 [32] | Homozygous c.233G > C, p.(Arg78Pro) | 1 month | Myoclonic | RMFSL | Corpus callosum thinning | Sharp wave discharges in the left forehead-parietal region than in the right forehead-parietal region. | 7 months |
| Stödberg, 2020 [18] | Compound heterozygous c.1771-1G > C, p.? | 1 month | NA | NA | Pachygyria | NA | Alive at |
| 3 | c.294dup; p.(Leu99Thrfs*92) | 1 month | NA | NA | Pachygyria | NA | Alive at |
| Nuovo, 2022 | Compound heterozygous c.638dup, p. | No | NA | NEDCAS | Mildly shrunken | NA | Alive at |
His birth weight was 1215 g and his head circumference was 27.2 cm, both below the third centile. He spent four weeks on the neonatal unit due to low birth weight and was noted to have abdominal distention, axial hypotonia, limb hypertonia and episodes of both bradycardia and apnea. At the age of one year, he still exhibited axial hypotonia and limb hypertonia in addition to horizontal and upbeat nystagmus, a sluggish pupillary reflex to light, and inability to fix his gaze on objects. He had persistent constipation requiring treatment with regular laxatives and enemas. A viral serology screen did not demonstrate evidence of congenital viral infection with CMV, parvovirus B19, Rubella, HSV, or congenital toxoplasmosis.

He had severe global developmental delay without regression. He did not smile until the age of 16 months, made babbling noises at the age of 21 months, and rolled at 22 months. He was never able to walk and required a wheelchair to mobilize. He had a wide nasal bridge, bilateral epicantilic folds, and dysmorphic ears.

At 8 months of age, he presented with an afebrile transient loss of consciousness lasting 5 min. He was reported to have been unresponsive, grinding his teeth with his eyes rolled backwards, and was grey and floppy without limb movements.

By the age of 3 years, his parents had witnessed at least six unresponsive, cyanotic, and hypotonic episodes with twitching of his limbs lasting for a few minutes each which were presumed to be bilateral tonic-clonic seizures of unknown onset. This was in addition to multiple daily brief episodes of apparent loss of awareness and occasional myoclonic jerks. At this point, epilepsy was diagnosed, and he commenced sodium valproate 200 mg twice a day. After two-years without tonic-clonic seizures, these relapsed, increasing in frequency to clusters of 3–4 seizures occurring approximately once a week. We have not been able to obtain medical records or a collateral history covering the period from childhood to until he represented at the age of 15 years, when he commenced topiramate but continued to have seizures weekly. This was thought to be contributed to, in part, by non-concordance with anti-convulsant medication when he stopped eating and drinking. A gastrostomy tube was subsequently inserted to support nutrition and medication administration. Despite this, seizure frequency worsened over the following five years, and at the age of 20, levetiracetam was added with an improvement in seizure frequency. He died aged 20 years due to aspiration pneumonia complicating unexplained gastroparesis with vomiting.

Blood tests in early childhood including full blood count, urea and electrolytes, and liver and thyroid profiles were normal, as were serum lactate and pyruvate. While the original data from electroencephalography (EEG) study at four years of age is no longer available, it was reported as showing symmetric background alpha, beta, and theta rhythms with occasional increased amplitude delta activity, which was notched and sharp. There were also frequent focal sharp wave and spike discharges, both simultaneously and independently over both hemispheres.

Table 1 (continued)

| Case | BRAT1 Variant | Syndrome | Seizure type | MRI head findings | EEG findings | Survival |
|------|--------------|----------|--------------|-------------------|-------------|---------|
| 1    | (Val214GlyfsTer189) | widened interfolial spaces | tonic-clonic staring, blepharospasm, myoclonic-epileptic absence | symmetric background alpha, beta, and theta rhythms with occasional increased amplitude delta activity, which was notched and sharp. There were also frequent focal sharp wave and spike discharges, both simultaneously and independently over both hemispheres | Alive at 10 years | 8 months |
| 2    | c.1395G > A, p.(Thr465 = ) | shrunken cerebellum | tonic-clonic starting, blepharospasm | diffuse sharp and wave complexes | Alive at 20 years | 18 years |
| 3    | c.294dupA, p.(Leu99Thrfs*92); c.1925C > A, p.(Ala642Glu) | cerebellar hypoplasia | tonic-clonic, staring myoclonic, suspected absence | symmetric background alpha, beta, and theta rhythms with occasional increased amplitude delta activity, which was notched and sharp. There were also frequent focal sharp wave and spike discharges, both simultaneously and independently over both hemispheres | Alive at 20 years | 14 years |

By the age of 3 years, his parents had witnessed at least six unresponsive, cyanotic, and hypotonic episodes with twitching of his limbs lasting for a few minutes each which were presumed to be bilateral tonic-clonic seizures of unknown onset. This was in addition to multiple daily brief episodes of apparent loss of awareness and occasional myoclonic jerks. At this point, epilepsy was diagnosed, and he commenced sodium valproate 200 mg twice a day. After two-years without tonic-clonic seizures, these relapsed, increasing in frequency to clusters of 3–4 seizures occurring approximately once a week. We have not been able to obtain medical records or a collateral history covering the period from childhood to until he represented at the age of 15 years, when he commenced topiramate but continued to have seizures weekly. This was thought to be contributed to, in part, by non-concordance with anti-convulsant medication when he stopped eating and drinking. A gastrostomy tube was subsequently inserted to support nutrition and medication administration. Despite this, seizure frequency worsened over the following five years, and at the age of 20, levetiracetam was added with an improvement in seizure frequency. He died aged 20 years due to aspiration pneumonia complicating unexplained gastroparesis with vomiting.

Blood tests in early childhood including full blood count, urea and electrolytes, and liver and thyroid profiles were normal, as were serum lactate and pyruvate. While the original data from electroencephalography (EEG) study at four years of age is no longer available, it was reported as showing symmetric background alpha, beta, and theta rhythms with occasional increased amplitude delta activity, which was notched and sharp. There were also frequent focal sharp wave and spike discharges, both simultaneously and independently over both hemispheres, consistent with multifocal epilepsy. The report does not mention capture of the events characterized by loss of awareness, that we classify as unknown onset behavior arrest seizures on purely clinical grounds. Similarly, the original magnetic resonance images acquired the same year are no longer available for review but were reported as showing cerebellar hypoplasia with a prominent vermis. EEG during his terminal illness in the intensive care unit without sedation was consistent with severe encephalopathy, demonstrating widespread low amplitude theta and delta rhythms without reactivity to auditory stimuli.

Genetic testing yielded a normal karyotype with no evidence of fragile X syndrome. Whole exome sequencing was performed during his terminal admission at 20 years of age, and reported as an in silico panel of selected genes (gene list provided as a supplement) by GEMINI (Cambridge University Hospitals NHS Foundation
Trust). This revealed two variants in \textit{BRAT1}, c.294dupA, p. (Leu99Thrfs*92) and c.1925C>A, p.(Ala642Glu). Both variants have been interpreted as pathogenic in previous publications and are present in gnomAD (p.Leu99Thrfs*92 on 42 occasions and p. Ala642Glu 8 times) below the maximum frequency that would refute their pathogenicity in a recessive disorder and neither has been reported in the homozygous state in gnomAD [7,13,14]. We interpret them according to American College of Medical Genetics and Genomics criteria as pathogenic and likely pathogenic respectively [15]. The lack of parental DNA and the distance between variants has meant that it has not been possible to confirm that these variants are in \textit{trans} (compound heterozygous).

3. Discussion

This case illustrates some important clinical lessons applicable to the genetic investigation of epilepsy in adults. First, recessive forms of DEE may be identified with panel testing in adults, even without consanguinity. However, compared to pediatric patients, adults may be less likely to have parental DNA available to confirm the suspected genetic mechanism through segregation of variants; this is also a challenge to the interpretation of monoallelic potentially de novo variants. In the very near future, advances in genetic sequencing technologies will help with the interpretation of variants in recessive disease genes where segregation cannot be demonstrated. In this case, the variants were 5 kilobases apart, raising the possibility that a single long-read may span both loci. Second, among the increasing number of young adults transitioning to adult services with DEE, some have disorders that were previously not recognized in adulthood due to the biases of genetic discovery and initial phenotypic studies, which focus on pediatric cohorts, often comprising children with particularly severe phenotypes. Consequently, some genetic causes may thus far have been associated only with the more severe extreme of their latent clinical spectrum. Finally, for adult patients, the threshold for testing should take account of the clinical heterogeneity of each gene, and the possible incompleteness of historical records of clinical phenomenology, development, EEG, and MRI. These may have been destroyed or incomplete, or not followed the patient from hospital to hospital. When assessing an adult with early onset epilepsy, the historical records and previous investigations are likely to predate the discovery of the particular cause of their epilepsy. Accordingly, the absence of the corresponding classic gene-phenotype association in these medical records should not be assumed to mean that the phenotype was never present — it may not have been actively sought.

The \textit{BRAT1} gene (7p22.3) encodes the BRCA1-associated ATM activator 1 protein. This protein is involved in the response to DNA damage and apoptosis and in the regulation of mitochondrial homeostasis [10]. Few detailed descriptions of people with disorders attributable to \textit{BRAT1} variants have been published, but these have revealed heterogeneous phenotypic constellations with a varying degree of clinical severity. If seizures are present, they are typically but not exclusively seen in early infancy. The median age of seizure onset in the reported cases of \textit{BRAT1}-related disease is the first day of life (Table 1).

At the most severe end of the clinical spectrum of disease is RMFLS, typified by severe drug-resistant seizures beginning in utero or within the first days of life, microcephaly, rigidity, swallowing difficulties, and poor psychomotor development. Infants with RMFLS experience apneic and bradycardic episodes culminating in cardiorespiratory arrest and death, typically within the first year of life [5].

A less severe clinical course is characterized by features similar to RMFLS but with survival past infancy, later onset epilepsy and continued development [5–8]. Our case matches this phenotype, with evidence of microcephaly, limb hypertonia, severe global developmental impairment and drug resistant multifocal epilepsy, and represents the longest reported survival in such an individual with this phenotype to date: the previous oldest patient with this phenotype was reported to still be alive at the age of six years and could be as old as 11 now [7]. The mildest reported constellation of \textit{BRAT1}-related disease is characterized by intellectual disability, ataxia, and cerebellar atrophy with or without microcephaly and epilepsy [8,12].

It appears that the severity of disease is linked to zygosity of the \textit{BRAT1} variants with most of the reported homozygous cases developing seizures immediately after birth or suspected to have had seizures in utero. The longest reported survival for a homozygous patient is 12 months [16]. The reported cases with compound heterozygous \textit{BRAT1} variants, however, show a relatively delayed onset of seizures, typically within the first few months of life but in a few cases, after one year of age. These patients generally survive longer, with several previously reported cases of survival into childhood and the present case showing that survival into adulthood is possible. However, two sets of siblings with compound heterozygous variants have shown significant phenotypic variability, indicating that genetic expressivity is variable and other genetic or non-genetic factors contribute [6,17]. Overall, it appears that \textit{BRAT1}-related epilepsies can include both focal-onset and generalized-onset seizures. Beyond the individual we present here with unknown onset for seizures associated with behavioral arrest, only one other has been reported to have seizures compatible with absences (case 3 in Srivastava, 2016 [7]). This person had late onset epilepsy given the context of \textit{BRAT1}-related disorder (3 years) and survival to at least 4 years. This may reflect the difficulty of detecting non-motor seizures in infancy or the increased propensity for absent seizures to occur as the brain develops towards mid-childhood (onset of absences before 4 years of age is rare across the epilepsies and should prompt consideration of genetic testing). In either case, the recognition of non-motor seizures in a person with \textit{BRAT1}-related epilepsy may be a marker of survival beyond infancy, and thus a milder form of these diseases.

A person with identical \textit{BRAT1} variants has previously been described by Mundy et al., who were able to confirm compound heterozygosity and emphasized survival beyond infancy [7]. Similarly, to the present case, they describe microcephaly, mild dysmorphic features, abnormal tone, and developmental delay in addition to drug-resistant seizures. However, the individual that they reported had developed seizures at a younger age (3 months vs 8 months). The three \textit{BRAT1} cases reported in Stödberg et al all had shared the c.294dupA, p.(Leu99Thrfs*92) genotype with our case; all three were reported to have pachygyria – not reported in the individual we report, for whom we were not able to review the original MRI or repeat the study [18]. The other variant we report here, c.1925C>A, p.(Ala642Glu), has been described in compound heterozygosity in someone with a neurodevelopmental syndrome with progressive neurodegeneration that fits neither the syndromes of EIMFS nor RMFLS, and who had focal impaired awareness seizures [7]. It is possible that the events with apparent loss of awareness of the person we report were focal impaired awareness seizures.

The prevalence of epilepsy due to \textit{BRAT1} variants in adults is currently unknown, although none was detected among 150 adult patients with neurodevelopmental disorders with epilepsy (using a gene panel that included \textit{BRAT1}) [19]. The national screening study of children with seizures in Scotland did not include \textit{BRAT1} on its panel [20].
4. Conclusion

Biallelic pathogenic BRAT1 variants cause a spectrum of clinical disease with and without seizures. With wider testing and a higher index of suspicion, we may see cases with longer survival in the context of a later age of seizure onset and compound heterozygosity. This case expands the clinical spectrum further to include people reaching adulthood with multifocal epilepsy, dysmorphic features, and developmental delay.

Declaration of Competing Interest

The authors declare the following financial interests/personal relationships which may be considered as potential competing interests: R.H.T. reports Honoraria from Arvelle, Bial, Eisai, GW Pharma, Sanofi, UCB Pharma, UNEEG and Zogenix. E.A. reports Honoraria from Arvelle, R.F., M.E., C.J.M., and D.L.-S. have nothing to declare.

Acknowledgements

D.L.-S. is a Wellcome Clinical PhD Fellow funded through the 4ward North Clinical PhD Academy. This research was funded in whole, or in part, by the Wellcome Trust [Grant number 203914/Z/16/2]. For the purpose of open access, the author has applied a CC BY public copyright license to any Author Accepted Manuscript version arising from this submission. This work was also supported by the Academy of Medical Sciences [Grant number SGL015/1029].

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.ebr.2022.100549.

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