

How to SHINE

A GUIDE TO NAVIGATING LIFE WITH
SHINE SYNDROME/DLG4 SYNAPTOPATHY



Love Fearlessly! Shine Fearlessly!



How to SHINE

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All underlined words in this guide are clickable and linked to corresponding content for your convenience.



Welcome, we are here for you.

We would like to extend you a warm welcome to the SHINE Syndrome/DLG4 Synaptopathy community. Getting a diagnosis can be an overwhelming experience. You may feel a whole range of emotions, from confusion to grief or even a sense of relief. Wherever you are on your journey, please know that we are here for you.



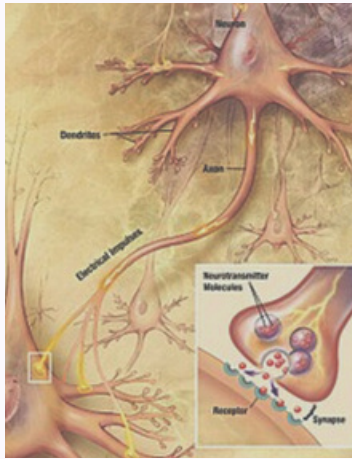
important tip

Please feel free to share this guide with health practitioners, teachers, family and friends so that they can learn more about SHINE Syndrome and how to best support you, your child and the SHINE community.

This guide includes information to help you learn more about SHINE Syndrome, otherwise known as DLG4-related Synaptopathy and to connect you with our community. We hope that you use it as a resource that you can refer back to as needed. Please take your time reading and feel free to skip around as you see fit. Given that SHINE is a rare neurological condition, there is a great probability that your support team has little or no knowledge about SHINE. Please share this guide with them. We are here to give you the support, information and resources you need to be a more confident advocate for you and your SHINE loved one.



What is SHINE Syndrome?



SHINE Syndrome, also known as DLG4 Synaptopathy, is an ultra rare neurodevelopmental disorder. SHINE is an acronym for its most common symptoms.

Sleep Disturbances

Hypotonia

Intellectual Disabilities

Neurological Disorders

Epilepsy



SHINE Symptoms

SHINE Syndrome is characterized predominantly by global developmental delay, intellectual disability, autism spectrum disorder, attention deficit hyperactivity disorder, hypotonia and epilepsy. Many individuals with SHINE Syndrome also present with sleep disturbances, skeletal abnormalities, and/or structural brain abnormalities (seen on an MRI). Families and patients also report symptoms such as: Sensory Processing Disorder, Dyspraxia, Apraxia and Speech Disorders. *SHINE patients do not always have all these symptoms. Symptoms can be experienced as mild to severe.*

Over 50% of people affected by SHINE develop epilepsy. Of those diagnosed with epilepsy, many families have reported that their loved one has been diagnosed with ESES (or CSWS) which is a rare and severe form of epilepsy that stands for Electrical Status Epilepticus of Sleep. For more information about ESES, click [here](#).

Epilepsy can be challenging to detect and diagnose especially when they are brief, non-convulsive or atypical, or generally happen during sleep (like ESES). *If epilepsy or seizures are suspected, please ask for a referral to a neurologist (or epileptologist).* An EEG combined with clinical evaluation is generally needed for a diagnosis.

Please work closely with your medical team for personalized testing and treatment for all neurological symptoms.



The Rarity of SHINE Syndrome

We currently are aware of over 75 patients in the world but we know there are more undiagnosed patients out there. People are yet to be diagnosed for a few reasons:

- Families receive a diagnosis like autism, intellectual disability, or epilepsy and discontinue their research or are not offered genetic testing by their medical team
- Cost and access to genetic testing can be a barrier to getting a genetic cause for the patient's symptoms
- Prior to 2019, DLG4 variants were considered VUS (variants of unknown significance). This changed in 2019 to confirm that DLG4 variants may be pathogenic (disease causing). This means that those tested prior to 2019 may not have had tests that include DLG4 or they may not know that their variant changed from VUS to pathogenic.



*alone we are rare,
together we are strong®*



SHINE Genetics

The human body is made of trillions of cells. Each cell contains 23 pairs of chromosomes (46 total). Each chromosome contains thousands of genes. Most genes also come in pairs and we get one copy from each parent. The role of genes is to produce proteins. Proteins are used to regulate the body's tissues and organs. A gene can stop working or no longer work properly when a variant/mutation occurs. A variant is a mistake that happens, similar to a typo, when the DNA is copied from cell to cell or due to environmental factors.

SHINE Syndrome is inherited in an autosomal dominant fashion. This means that only a single copy of the disease-associated mutation is enough to cause the disease. Most individuals with SHINE syndrome are found to have de novo (new) variants in DLG4, meaning the variant occurs for the first time in them and is not inherited from a parent. There are families, however, where there has been sibling reoccurrence of SHINE syndrome. This is hypothesized to be due to a phenomenon known as mosaicism (the variant is present in small amount of the parent's cells, but all of the child's cells).

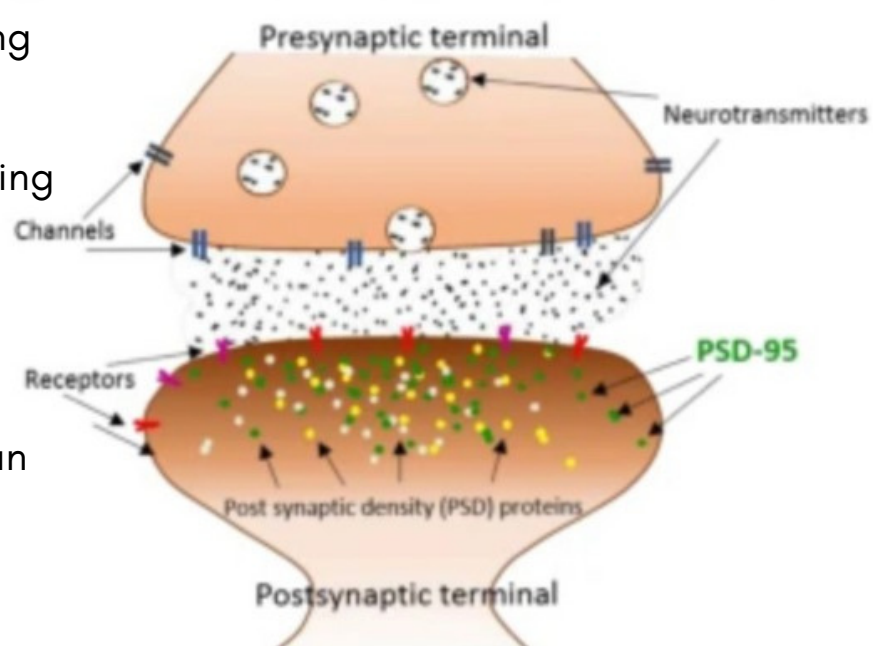
To learn more about this please visit our informational series on YouTube hosted by one of our medical advisors, Alexandre White-Brown.



SHINE and PSD-95

SHINE Syndrome is caused by variants/mutations in the DLG4 gene. The DLG4 gene is located on the 17th chromosome and is an important gene that encodes the protein PSD-95 (Postsynaptic Density Protein 95). *PSD-95 plays a major role in brain development and function through its implications in synaptic strength and plasticity.*

These mechanisms, along with PSD-95's role in organizing and interacting with other proteins, represent a gene with many capabilities of which, when altered, can induce susceptibility to SHINE Syndrome.



To learn more about this please visit our informational series on YouTube hosted by one of our medical advisors, Alexandre White-Brown.



We are grateful to one of SHINE's medical advisors, Dr. Zeynep Tümer, who created this informative piece for the SHINE community.

VARIANTS IN DLG4 and THEIR PREDICTED EFFECTS ON PSD-95

Prepared by Professor Zeynep Tümer, MD, PhD, DMSc

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Rigshospitalet, Denmark

A METAPHORIC EXPLANATION OF DIFFERENT DLG4 VARIANTS (SEE FIGURE 1)

A stretch of three nucleotides (combinations of A, C, G, or T) code for a specific amino acid, and these three nucleotides are called codons. Proteins are composed of amino acid chains. Some codons are called STOP codons, and these signal that the protein chain should stop.

In the next illustration we made a sentence (protein) composed of words (codons) with three letters (nucleotides). The start of the protein is shown with THE and the stop codon with END.

Apart from the normal sequence, we illustrate three common types of variants (mutations) detected in DLG4.

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DIFFERENT DLG4 VARIANTS AND THEIR PREDICTED EFFECT ON THE PROTEIN

According to their predicted effect on PSD-95, we can divide the disease-causing variants in DLG4 into two groups:

A. Protein truncating variants: Frameshift and nonsense variants (see figure 2)

These variants are predicted to result in two possibilities.

Possibility 1: Truncated protein which has lost its normal function

Possibility 2: The protein is not produced at all due to a mechanism called nonsense mediated mRNA decay.

The result is that the affected person has only one normal gene-copy producing a normal protein, while no protein is produced from the other copy of the gene. In short, the protein has lost its function.

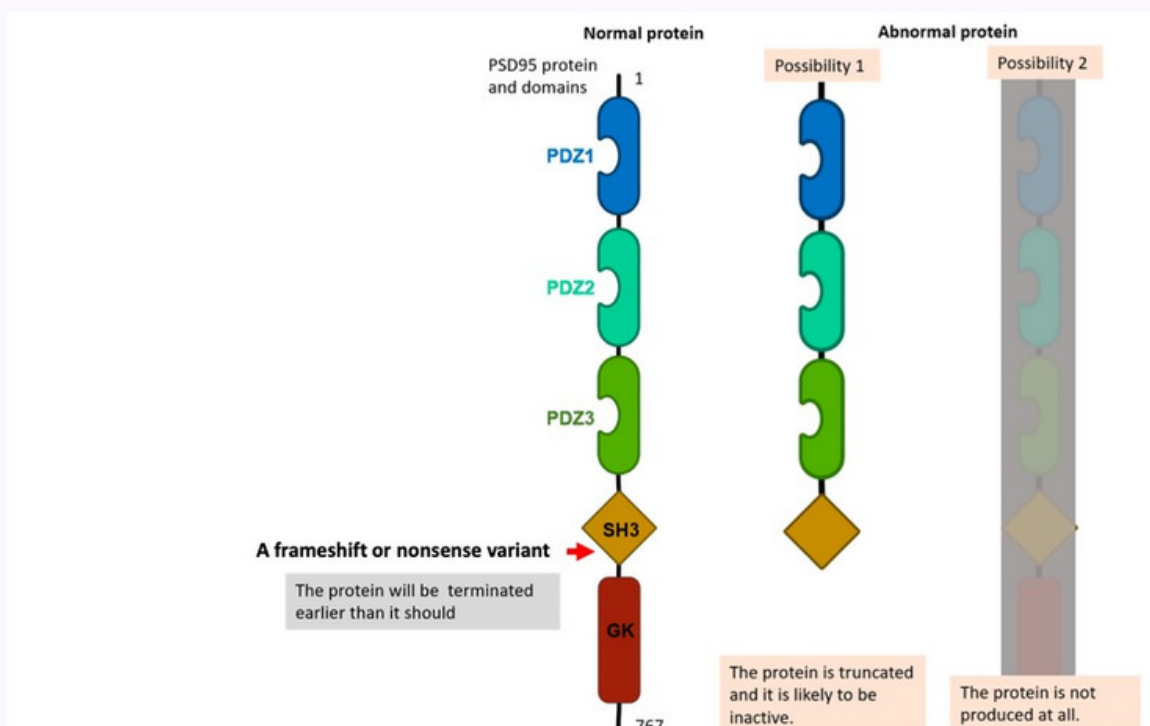
B. Missense variants (see figure 3)

These variants change a single amino acid and they do not truncate the protein. The effect of these variants are not known, but we predict that they result also in the loss of function of the protein. With other words, the protein is there but it does not function as it should.

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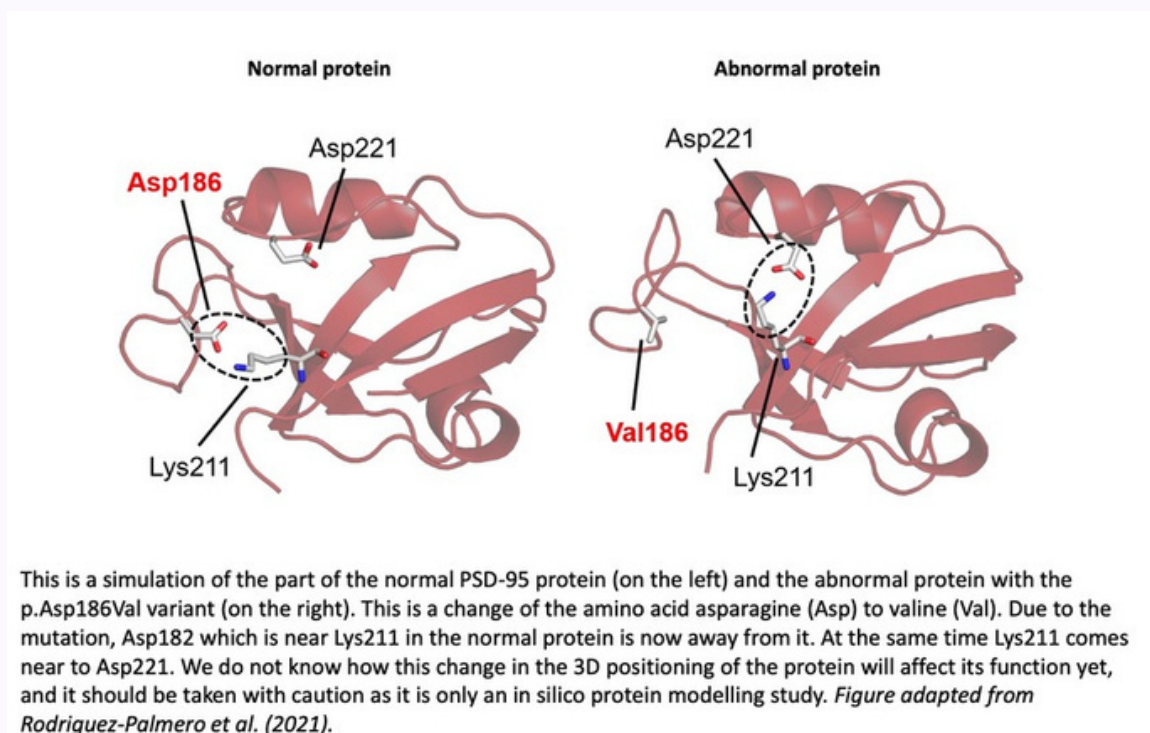
Figure 2 - Protein Truncating Variant



PSD-95 protein has some functional domains: PDZ1, PDZ2, PDZ3, SH3, and GK.

See, Rodriguez-Palmero et al. (2021) or Levy et al. (2022) for more details. *Figure adapted from Levy et al. (2022)*

Figure 3 - Missense Variant (Example p.Asp186Val in the PDZ2 domain)



SHINE Syndrome Foundation

Our foundation is founded and run entirely by parents of children living with SHINE Syndrome. Our board members are listed [here](#). We would like to thank all the other individuals that have helped us get where we are today!

Timeline

In just a few short years, our organization has come a very long way. Explore our timeline to see what we've done recently to expand awareness and support for SHINE Syndrome.



PRE-2019 VUS

RESEARCH HAS NOT YET IDENTIFIED THE EFFECTS OF VARIANTS WITHING THE GENE. DLG4 VARIANTS CLASSIFIED AS VUS OR VARIANCE OF UNKNOWN SIGNIFICANCE



2019 PATHOGENIC STATUS

VARIANTS IN DLG4 WERE CONFIRMED TO BE DISEASE CAUSING OR PATHOGENIC



2019 PARENTS UNITE

PATIENTS INFORMED OF THE PATHOGENIC STATUS OF THEIR VARIANTS. PARENTS FORMED CONNECTIONS AND A FACEBOOK GROUP. WITHIN A YEAR, THE COMMUNITY GREW TO MORE THAN 15 FAMILIES.



2021 DISEASE NAMED

WITH THIS **PUBLICATION**, DLG4 SYNAPTOPATHY WAS NAMED. SOON AFTER, A FEW INVESTED MOTHERS OF SHINE PATIENTS CREATED THE NAME SHINE SYNDROME FOR THE COMMON CHARACTERISTICS OF THE DISEASE.



2021 FOUNDATION FORMED

SHINE SYNDROME FOUNDATION WAS OFFICIALLY FORMED IN DECEMBER, 2021 BY A DEDICATED GROUP OF PARENTS OF SHINE PATIENTS



SHINE Research and Awareness

The SHINE Syndrome Foundation has been extremely fortunate to gain the support of a dedicated group of researchers and clinicians around the world. Several of those individuals have also agreed to participate on our Medical and Science Advisory Board in hopes of helping guide us towards research funding opportunities and raising awareness for SHINE Syndrome throughout the medical community.

To view published research articles on DLG4, learn more about current research and our medical advisory board please visit [here](#).

If you would like to get involved in research and awareness for the SHINE community, please see opportunities below*. Please know that we recognize that all of this can be overwhelming. We encourage you to participate in whatever capacity is comfortable for you and your family.

SHINE Awareness Opportunities

- Please consider sharing photos [here](#) for our social media bank to help spread awareness of SHINE Syndrome. Read more about our community [here](#).
- We'd also love to share your personal SHINE story on our community page. Please email us at contact@shinesyndrome.org if you are interested.

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SHINE Research Opportunities

Simons Searchlight is a research registry for DLG4 Synaptopathy and other rare genetic neurodevelopmental disorders. They collect data and blood samples and share the information with leading researchers around the world to improve the lives of people living with rare neurodevelopmental disorders. Participation is open worldwide to people who speak English, Dutch, French, and Spanish, and more languages are coming soon. People of any age with a DLG4 Synaptopathy diagnosis and their family members can sign up [here](#).

Dr. Zeynep Tümer (one of our medical advisors) is leading a research initiative to study DLG4 and PSD-95. They have submitted a large grant application through the Chan Zuckerberg Initiative to expand this research even further. You can participate by sharing your variant with the team. A detailed explanation on how to share this information can be viewed [here](#). Alternatively, it can be completed [here](#).

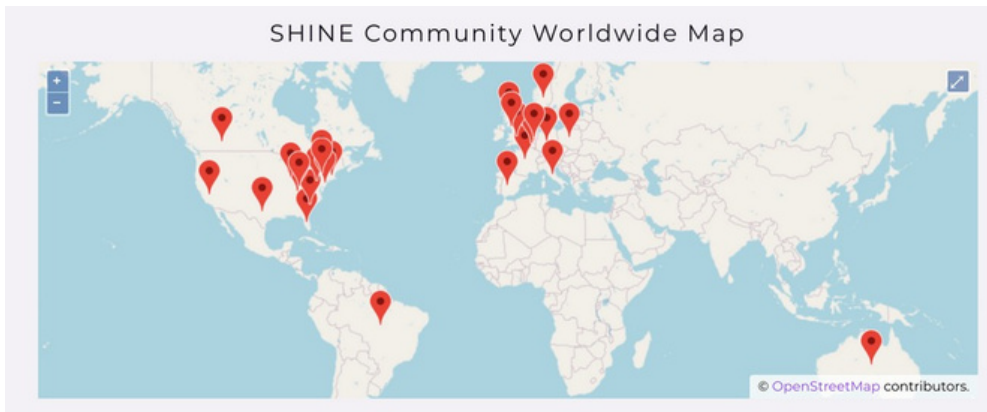
Please check back here shortly for the CoRDS (Coordination of Rare Diseases at Sanford) Registry.

Please check back here for information upcoming epilepsy research initiative.



Join Our Community

In just a short period of time, (as of Summer 2022) our community has grown to include over 75 known patients around the world. We'd love to get to know you and be a part of your support system. For community information and patient stories please visit [here](#).



Please join our private Facebook support group [here](#). The group primarily functions as a support group for families. We warmly welcome, support and appreciate our DLG4 adult patients and caregiver members. Select DLG4 researchers are also welcome and participate only to help answer questions related to research. While respecting group rules, you can ask questions and participate with no judgment and cheer for each other as progress is made.

Please also join our public Facebook foundation [group](#) where we share SHINE Syndrome Foundation news and all things pertinent to DLG4 Synaptopathy.

We are here for you! If you prefer private support or Facebook is not your thing, please reach out to Courtney, our Director of Patient & Family Engagement at croche@shinesyndrome.org.



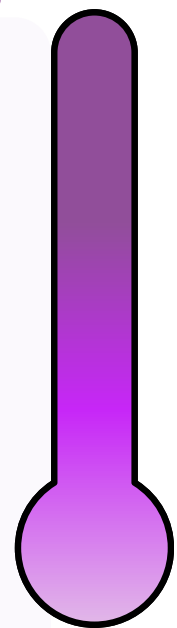
SHINE Syndrome Foundation: How to Get Involved

At SHINE Syndrome Foundation, we strive to build awareness for our growing community as well as to help provide funding for researchers around the world who are searching for answers and working to unravel the secrets of SHINE Syndrome. Through sharing our stories on our website, increasing visibility on social media, and participating in fundraising efforts, we can connect with newly diagnosed patients, provide a community for the families living with SHINE, and assist in finding strategies and treatments for those in need.

Though we understand that not everyone will be able to contribute financially, we appreciate anyone willing to share the word about SHINE Syndrome on social media or among family and friends. For more information on fundraising, please visit [here](#).

SHINE Syndrome Foundation set a goal of reaching \$50,000 in July 2022 for our sunSHINE campaign. We beat our goal and were able to raise over \$73,500. We are currently at \$90,000 total raised funds (as of September 2022). Thank you to all who have participated in our fundraising. Proceeds for the sunSHINE campaign will be used to fund a postdoctoral or PhD fellow focused on repurposing drugs for treatment of SHINE Syndrome.

\$90,000



Resources

"A hero is an ordinary individual who finds the strength to persevere and endure in spite of overwhelming obstacles."

Epilepsy Management

Epilepsy Foundation: epilepsy.com

Danny Did Foundation: dannydid.org

Child Neurology Foundation: childneurologyfoundation.org

Autism

ASAN autisticadvocacy.org

Autism Society autismsociety.org

Sensory Processing Disorder

Star Institute: spdstar.org

Apraxia

Apraxia Kids: apraxia-kids.org

Rare Diseases:

Global Genes: globalgenes.org

NORD: rarediseases.org



Stay Up to Date on SHINE Syndrome

Visit our [website](#) to stay current on SHINE news.

To make sure that you don't miss anything important, please [sign up](#) for our newsletter.

Read our latest newsletter [here](#)



Save the date!

JULY							
Week	I	II	III	IV	V	VI	VII
27			1	2	3	4	5
28	6	7	8	9	10	11	12
29	13	14	15	16	17	18	19
30	20	21	22	23	24	25	26
31	27	28	29	30	31		

**July 17 is
SHINE Syndrome
Awareness Day**



How to SHINE

CHECKLIST

Join Facebook Support and Foundation Pages

Sign Up for Research with Simon's Searchlight

Share Variant for Research with Dr. Tümer

Sign up for SHINE Newsletter

Share 'How to Shine' Guide with Support Team, Family and Friends

Help Fundraise for SHINE





Love Fearlessly! Shine Fearlessly!

