Drosophila functional screening of de novo variants in autism uncovers damaging variants and facilitates discovery of rare neurodevelopmental diseases

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SUMMARY

Individuals with autism spectrum disorder (ASD) exhibit an increased burden of de novo mutations (DNMs) in a broadening range of genes. While these studies have implicated hundreds of genes in ASD pathogenesis, which DNMs cause functional consequences in vivo remains unclear. We functionally test the effects of ASD missense DNMs using Drosophila through “humanization” rescue and overexpression-based strategies. We examine 79 ASD variants in 74 genes identified in the Simons Simplex Collection and find 38% of them to cause functional alterations. Moreover, we identify GLRA2 as the cause of a spectrum of neurodevelopmental phenotypes beyond ASD in 13 previously undiagnosed subjects. Functional characterization of variants in ASD candidate genes points to conserved neurobiological mechanisms and facilitates gene discovery for rare neurodevelopmental diseases.

Graphical Abstract

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AUTHOR CONTRIBUTIONS

M.F.W. and S.Y. conceived and designed the project. P.C.M., J.A., S.L.D., and J.M.H. designed and conducted most fly experiments and analyzed the data. V.H.B. and Y.-H.C. performed cloning and mutagenesis. H.K.G. performed cloning and coordination. S.J. performed immunostaining and confocal microscopy. X.L. performed structural analysis and generated reagents. N.L., D.B., Y.-H.C., P.L., B.H., H.P., P.B., M.-C.H., C.M.L., H.-T.C., H.C., N.A.H., O.K., and S.N.M. contributed to reagent generation and some fly experiments. R.M., A.G., E.M., C.S., R.G., A.V., E.E., C.N.M., T.S.B., M.F.v.D., M.W., M.v.S., G.L., I.S., N.C., C.A.B., J.A.M., P.B.A., B.K., T.C., L. Perrin, M.B., T.R., S.L., F.T.-M.-T., J.D., E.S.-A., S.T., E.R., V.S., K.P., R.K., L. Pavinato, and A.B. reported and described GLRA2 subjects. L.Z.R., R.J.G., and J.A.R. aided in subject matchmaking and collection and organization of patient data. P.C.M., J.A., S.L.D., J.M.H., M.F.W., and S.Y. wrote and revised the manuscript.

SUPPLEMENTAL INFORMATION

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DECLARATION OF INTERESTS

The Department of Molecular and Human Genetics at Baylor College of Medicine receives revenue from clinical genetic testing completed at Baylor Genetics Laboratories.
Marcogliese et al. generate >300 Drosophila mutants and use complementary rescue-based and overexpression approaches to study the function of de novo missense variants found in autism. They find that 38% of missense changes have functional consequences and identify variants in GLRA2 that cause a variable neurological disorder.

INTRODUCTION

Autism spectrum disorder (ASD) is a complex neurodevelopmental condition with impairments in social interaction, communication, and restricted interests or repetitive behaviors (APA, 2013). Individuals affected by ASD, particularly in severe cases, exhibit a higher burden of de novo mutations (DNMs) in an expanding list of genes (Coe et al., 2019; Fischbach and Lord, 2010; Iossifov et al., 2014). The genetic burden of DNMs in ASD subjects has been estimated to account for ~30% of disease causation (Iossifov et al., 2014; Rubeis et al., 2012; Sanders et al., 2012; Satterstrom et al., 2020; Takata et al., 2018; Yuen et al., 2017). While these studies have implicated hundreds of genes in ASD pathogenesis, which of these genes and variants causally contribute to this disease remains unknown. Missense DNMs in particular present a unique challenge because most genes lack established functional assays. Drosophila melanogaster is a genetically tractable system that is widely used to study human diseases (Bellen et al., 2019; Link and Bellen, 2020; Marcogliese and Wangler, 2001). In addition to studying disease mechanisms by
establishing preclinical models, flies can be used as a “living test tube” to study functional consequences of variants of unknown significance found in subjects. Here, we integrate a number of state-of-the-art technologies in the fly field to establish an in vivo pipeline to effectively study the functional impact of DNMs identified in a large cohort of ASD subjects.

RESULTS

Prioritization of ASD variants to study in Drosophila

We prioritized genes with coding DNMs identified in ASD probands from the Simons Simplex Collection (SSC) (Iossifov et al., 2014) that are conserved in Drosophila. We primarily focused on missense variants and in-frame indels because functional consequences of these variants are more difficult to predict compared with nonsense and frameshift alleles. However, we also tested a few truncating variants in single-exon genes because these transcripts escape nonsense-mediated decay. In this cohort, 1,708 ASD proband-specific de novo missense or in-frame indels were identified through whole-exome sequencing (WES) (Figure 1A), corresponding to 1,519 unique human genes. Of these, 920 fly genes corresponding to 1,032 human genes were identified. Based on multiple ortholog prediction algorithms scores, 487 human genes had no or weak ortholog candidates in Drosophila (cut off: DIOPT <4/16; Hu et al., 2011; Table S1). By overlapping these 920 Drosophila genes with available fly lines containing Minos-mediated integration cassette (MiMIC) transposons within coding introns that permit targeting of all annotated protein isoforms (“gold”; 1,732 insertions; Nagarkar-Jaiswal et al., 2015; Venken et al., 2011), we identified reagents for 122 fly genes corresponding to 143 human genes and 179 ASD proband variants from the SSC. Compared with the entire genome, the SCC subset and our study subset showed enrichment for constrained genes by assessing gene-level metrics, such as probability of loss of function intolerance (pLI) (Figure 1B; Lek et al., 2016), loss of function observed or expected (o/e) upper bound fraction (LOEUF) (Figure 1C; Karczewski et al., 2020), or missense o/e (Figure 1D; Karczewski et al., 2020).

Of the 122 fly genes that met our selection criterion, we were able to successfully generate 108 T2A-GAL4 (TG4) lines via recombinase-mediated cassette exchange (Diao et al., 2015; Gnerer et al., 2015; Lee et al., 2018). These 108 TG4 lines correspond to 128 SSC genes (some fly genes correspond to multiple human genes with variants in SSC; Figure S1). These fly lines behave as loss-of-function (LoF) alleles that simultaneously produce a GAL4 transactivator in the same temporal and spatial pattern as the gene of interest (Figure 1E; Tang et al., 2009).

To generate upstream activating sequence (UAS) human reference transgenic (Ref-Tg) and human SSC candidate variant transgenic (SSC-Tg) fly lines, we obtained human open reading frame (ORF) collections from the Mammalian Gene Collection (MGC Project Team et al., 2009) or commercial sources (Figure 1F). Out of the 143 human genes and 179 SSC-DNMs of interest that we attempted to generate, we were successful in generating 194 UAS-cDNA (106 Ref-Tgs; 88 SSC-Tgs) flies (Figures 1G and S1; Tables S2 and S3). The UAS-Ref-Tg and UAS-SSC-Tg were inserted into the same genomic docking site in the
fly genome and are generated with the same construct, only differing in the variant point mutation, allowing for direct comparison of function by controlling for positional effects.

We were able to make a complete set of TG4, UAS-Ref-Tgs, and UAS-SSC-Tgs lines for 65 fly genes corresponding to 74 human genes and 79 variants (again, some fly genes correspond to multiple human genes and multiple SSC variants are found for a small subset of human genes), which were critical to test variant function using a rescue-based humanization strategy. In summary, 302 Drosophila stocks were generated for this project as a resource for the community, and these stocks are available from the Bloomington Drosophila Stock Center (BDSC) or in the process of being transferred and registered at BDSC (please see Figure S1 for detailed screen pipeline data).

In order to determine whether there are functional differences between the Ref-Tg and SSC-Tg, we used them in combination with TG4 lines to “humanize” Drosophila genes or crossed these lines to ubiquitous and tissue-specific drivers to ectopically overexpress reference or variant human proteins and assessed them for phenotypic differences (Bellen et al., 2019; Figure 1H). If the SSC variant lacked a function a reference allele possessed, we classified it as a LoF allele (e.g., amorph and hypomorph). If the variant had some function that the reference allele did not possess, we classified it as a gain-of-function (GoF) allele (e.g., hypermorph, antimorph, and neomorph) in this study.

**Humanization of essential Drosophila genes reveal loss-of-function ASD variants**

We identified 47 of 65 lethal TG4 mutants that remained lethal when placed in trans with a corresponding deficiency line. These 47 TG4 lines correspond to 60 human ASD candidate genes for both reference and variant human cDNA transgenic fly lines (Figure S1; Table S4). To assess whether the human homolog can replace the corresponding fly genes, we determined whether UAS-Ref-Tg can rescue the lethality of lethal TG4 mutants. We assessed rescue at four temperatures (18°C, 22°C, 25°C, and 29°C) as the GAL4/UAS system is temperature dependent (Nagarkar-Jaiswal et al., 2015). We found that lethality was suppressed in 17 of 37 genes tested (46%; Figure 2A; Table S5). We next tested whether SSC-DNMs have functional consequences by comparing the rescue efficiency of UAS-Ref-Tg and UAS-SSC-Tg. We observed significant functional differences in the ability for SSC variants to rescue lethality for ABL2, CAT, CHST2, TRPM6 (two variants), and TRIP12 (Figures 2B–2D). For ABL2 and CAT, we further found that humanized flies carrying the SSC-DNM (ABL2A1099T or CATG204E) had significantly decreased lifespan compared with reference animals (Figures 2E and 2F). Overall, we found that 32% (6/19) of the tested SSC-DNMs functionally differed from the reference in vivo, all behaving as LoF alleles.

To assess whether the fly homologs of human ASD candidate genes from SSC are expressed in the central nervous system (CNS), we crossed each TG4 line to UAS-nls.mCherry (red fluorescent protein with a nuclear localization signal) and performed co-staining with neuronal (Elav) and glial (Repo) nuclear markers. We chose the anterior central brain of the fly CNS to image as it is enriched for neuronal nuclei. All five genes associated with deleterious LoF DNMs were expressed in the adult central brain (Figures 2G and S2A). Examining neuronal and glia expression throughout the central brain revealed that all five genes were found in subsets of neurons and some glia, where Abl (corresponding to ABL2)
and ctrip (TRIP12) are expressed in neurons and a wider set of glia (Figure S2B). All five genes are also expressed in the third-instar larval CNS (Figure S2C).

Humanization of viable Drosophila TG4s reveals ASD variants with altered function

While we were able to test the function of 37 human genes based on rescue of lethality as mentioned above, 61 TG4 lines corresponding to 68 SSC-ASD candidate genes were viable and did not exhibit any obvious morphological phenotypes that can be used for variant functional studies. Note that 18 of 61 TG4 lines were homozygous lethal but were viable as compound heterozygotes over a molecularly defined deficiency that covers the locus, indicating that these lines carry second site lethal mutations (Figure S1), which has been previously reported in a subset of MiMIC strains (Nagarkar-Jaiswal et al., 2015). For the 43 of 61 remaining viable TG4 lines, we attempted to generate humanized TG4; UAS-Ref-Tg and TG4; UAS-SSC-Tg lines in an appropriate genetic background. Before carrying out behavioral studies, we had to replace the X chromosome of the TG4 lines with the X chromosome from a Canton-S strain to eliminate the effect of yellow (y) and white (w) alleles that were present in the original stocks (Figure S3). We were successful in generating all the necessary strains for eight genes. Using these humanized flies in a y+ w+ background, we performed courtship assays to assess social interactions in mutant and humanized flies considering that ASD patients exhibit social deficits (Figure 3A; Liu, 2013). Fly courtship involves a complex set of neurological components involving sensory input, processing, and motor output (Guo et al., 2019). We measured the amount of time a TG4 male fly spent performing wing extensions as a proxy for courtship, as well as the amount of time spent copulating with a wild-type (Canton-S) female. In addition to quantifying these two parameters, we also measured the time flies spent moving within the test chamber to assess their locomotion and tracked grooming, a stereotypic behavior in flies that involves a complex neurocircuit (Seeds et al., 2014).

Of the eight SSC-DNMs tested, we found five variants that showed functional alterations from the reference allele in at least one of four behavioral paradigms (Figure 3B). Humanized KCND3R86P flies displayed increased movement and decreased grooming behavior when compared with the humanized reference flies (Figures 3C, S4A, and S4B). The humanized KDM2AR449K flies showed decreased time copulating compared with the humanized reference (Figures 3D and S4C–S4E). Humanized USP30P200S flies displayed decreased grooming behavior when compared with humanized reference flies (Figures 3E and S4F–S4H). While these variants have clear functional differences from the reference allele, it was difficult to classify them as clean LoF or GoF alleles. Humanized ALDH1LN900H flies displayed a significant reduction in courtship and an increase in grooming behavior when compared with the humanized reference fly or the TG4 mutant alone, potentially indicating the variant acts as some kind of GoF allele (Figures 3F, S4I, and S4J). Finally, humanized reference GLRA2 flies failed to copulate at all but still exhibited normal movement while the humanized GLRA2N136S flies were capable of copulating within the trial period similar to the TG4 mutant alone, suggesting it behaves as a LoF allele (Figures 3G, S4K, and S4L). Courtship assessment of humanized SSC-DNMs in HTR1D, SLC23A1, and MADD did not show altered function to reference (Figures S4M–S4P).
In summary, 63% (5/8) SSC-DNMs act functionally different from reference alleles using quantitative behavioral measurements in flies.

Finally, we determined the CNS expression of TG4 lines corresponding to all eight lines we were able to humanize. Surprisingly, we only detected expression of 4/8 genes in the adult (Figures 3H, S5A, and S5B) and larval CNS (Figure S5C). All are expressed in a subset of neurons, whereas GluClα (GLRA2) is also expressed in some glia (Figure S5B).

**Overexpression assays reveal ASD variants with diverse functional consequences**

We complemented our rescue-based assays by overexpressing Ref-Tg and SSC-Tg in a wild-type background using ubiquitous (tub-GAL4), eye-specific (GMR-GAL4), and wing-specific (nub-GAL4) drivers (Figure 4A). This approach has routinely been employed to discern functional differences between reference and disease-associated variant proteins in vivo, regardless of whether the phenotypic readout has similarities to the patient’s conditions (Ansar et al., 2019; Chung et al., 2020; Goodman et al., 2021; Harel et al., 2016; Huang et al., 2020; Kanca et al., 2019; Liu et al., 2018; Marcogliese et al., 2018; Post et al., 2020; Ravenscroft et al., 2021; Splinter et al., 2018). Critically, UAS-Ref-Tg and UAS-SSC-Tg are inserted into the same genomic landing site in the fly genome, and the construct only differs by the point mutation, allowing for direct functional comparison. Across the three drivers and testing 66 human genes (73 SSC variants), we found 21/73 SSC-DNMs (in 19 fly genes) showed functional alteration in phenotypic assays, 17/73 displayed phenotypes comparable to reference, and 35/73 did not produce a scorable phenotype (Table S5).

Twelve variants in eleven human genes (ATP2B2T818M, EPHA1V567L, GLRA2N136S, GRK4P385A, ITGAGR748G, IRF2BPIF30L, IRF2BPN701666*, KCND3R86P, MINK1C269R, NPFFR2M163I, PDK2R120Q, and TSC2R1557W) behaved as LoF alleles. GLRA2N136S, GRK4P385A, ITGAGR748G, KCND3R86P, MINK1C269R, NPFFR2M163I, PDK2R120Q, and TSC2R1557W were annotated as LoF alleles using a ubiquitous driver because they failed to reduce the expected viability to the extent of the corresponding reference alleles upon overexpression (Figure 4B). Notably, GLRA2N136S and KCND3R86P were also annotated as LoF alleles in our rescued based assay, showing consistency (Figures 3C and 3G). Moreover, KCND3R86P and IRF2BPN701666* variants behaved as LoF alleles when assessed with multiple drivers. KCND3R86P abolished the activity of the reference transgene, which caused lethality when overexpressed with a ubiquitous driver (Figure 4B). In the wing, KCND3R86P failed to produce a severe notching phenotype that is observed by expression of the reference transgene (Figure 4D). Ubiquitous or wing-specific overexpression of reference IRF2BPL caused lethality, whereas the IRF2BPLN701666* frameshift allele (note that IRF2BPL is a single-exon gene) does not cause any phenotype (Figures 4B and 4D). Interestingly, the missense variant, IRF2BPIF30L, behaved as a LoF using the wing driver but was indistinguishable using the ubiquitous driver, indicating it is likely to be a partial LoF allele(Figures 4B and 4D). In addition to IRF2BPL DNMs, variants in two genes (ATP2B2 and EPHA1) were identified as LoF alleles based on the wing-specific driver and assay. Wing-specific overexpression of reference ATP2B2 in the developing wing disc causes a curled wing phenotype while the ATP2B2T818M variant fails to do so (Figure 4D). Expression of reference EPHA1 caused a wing-size reduction and wing-margin...
serration, whereas the \textit{EPHA1}^{V567I} variant caused serrated wings of normal size (Figure 4D), indicating partial LoF.

Seven variants (\textit{ACE}^{Y818C}, \textit{GPC5}^{M133T}, \textit{MYH9}^{R1571Q}, \textit{PCP1024R}, \textit{SLC23A1}^{L465M}, \textit{HTR1D}^{T99N}, and \textit{BAIAP2L1}^{A481V}) behaved as GoF alleles. Flies overexpressing variant forms of \textit{ACE}, \textit{GPC5}, \textit{MYH9}, \textit{PC}, and \textit{SLC23A1} exhibited enhanced lethality when compared with reference protein (Figure 4B). The GoF nature of \textit{MYH9}^{R1571Q} was also observed in the wing-size-based assay (Figure 4D). \textit{HTR1D}^{T99N} displayed consistent stronger phenotypes compared with reference when expressed in the eye or the wing, resulting in eye size reduction and absent wing phenotype, respectively (Figures 4C and 4D). \textit{BAIAP2L1}^{A481V} caused a smaller, more crumpled wing phenotype compared with its reference allele (Figure 4D).

Intriguingly, \textit{EPHB1}^{V916M} and \textit{MAP4K1}^{M725T} exhibited conflicting results in the eye and wing; therefore, they could not be categorized as simple LoF or GoF variants. While overexpression of reference \textit{EPHB1} or \textit{MAP4K1} in the developing eye causes eye-size-reduction phenotype, SSC variant forms of either gene result in normal eyes, indicating they behave as LoF alleles in this tissue. However, the same variant transgenes for these two genes expressed in the wing result in blistered or crumpled wings, respectively, that are phenotypically stronger than the reference alleles (Figures 4C and 4D), indicating they behave as GoF alleles in this tissue. In summary, when a scorable phenotype was present, 48\% (21/44) of the SSC-DNMs tested with an overexpression strategy impact function. Furthermore, we found diverse SSC variant consequences, including 12 LoF, 7 GoF, and 2 with complex phenotypes.

While this overexpression-based screening approach was not directly investigating the function of genes in the nervous system, expression analysis revealed that most (15/19) fly genes that correspond to SSC-DNMs with a functional difference identified through our overexpression assay are expressed in the adult (Figures 3H, S6, S7, and S8A) and larval (Figure S8B) CNS (see Marcogliese et al., 2018 for \textit{Pits} [corresponding to human \textit{IRF2BP1}]). The deleterious nature of variants in three of these genes was identified in our behavioral screen (\textit{GluClalpha}, 5-HT1B, and \textit{Usp30}, Figure 3H). Four genes (\textit{CG6293}, \textit{Mhc}, \textit{Shal}, and \textit{Ance}) were not detected in the brain in our analysis, which could be because they are expressed at very low levels or may primarily function in non-neural tissues. While most genes are enriched in neuronal subpopulations, \textit{Pdk} (\textit{PDK2}) and \textit{hppy} (\textit{MAP4K1}) are enriched in glia as well. Interestingly, \textit{if ITGA8} is not detected in either neurons or glia but revealed a unique expression pattern, that may reflect its expression in tracheal cells. In addition, based on imaging with an nls.GFP that leaks into the cytoplasm, it may also be present within cells that wrap around neurons reminiscent of cells in \textit{pars intercerebralis}, a neuroendocrine organ analogous to the mammalian hypothalamus (de Velasco et al., 2007; Figure S8A). Taken together, most of the fly genes corresponding to SSC-DNMs in which we found \textit{in vivo} alterations using overexpression-based assays are expressed in the fly CNS, similar to hits from rescue-based studies.
Identification of 30 deleterious SSC variants by merging all functional data

In total, we found 29 missense and 1 frameshift SSC-DNMs that displayed functional differences when compared with their respective reference allele (for a total of 28 genes: one variant for 26 genes and two variants for two genes; Tables 1 and S5). Approximately 53% (30/57) of the SSC-DNMs exhibited functional differences compared with the reference. Intriguingly, in our study, we only found GoF variants for genes corresponding to viable TG4 fly mutants based on both rescue-based and overexpression-based assays (Figure S9A). Interestingly, while we were able to classify the variants into LoF or GoF based for most genes, we found in two cases where different assays gave different results.

When we informatically surveyed the genes and variants with functional consequences identified through our screen in comparison to other genes included in our study (variants with comparable function or those lacking a phenotype to assess) using the MARRVEL tool (Wang et al., 2017), we did not find any significant differences in gene-level metrics, such as pLI (Lek et al., 2016); LOEUF (Karczewski et al., 2020); missense o/e (Karczewski et al., 2020); pathogenicity prediction scores based on several in silico algorithms, including sorting intolerant from tolerant (SIFT) (Vaser et al., 2016), PolyPhen-2 (Adzhubei et al., 2010), and combined annotation-dependent depletion (CADD) (Kircher et al., 2014); or absence or presence of identical variant in gnomAD (Lek et al., 2016) (Tables S5 and S6; Figures S9B–S9Z). By analyzing Gene Ontology (GO) by PantherDB (Ashburner et al., 2000) and visualizing with reduce visualize gene ontology (REVIGO) (Supek et al., 2011), ASD candidate genes from SSC with deleterious variants in vivo were compared with all protein-coding genes. We found most significant enrichment for genes with GO terms for “synapse (cellular component)” (Figure S10A) and “ion binding (function)” (Figure S10B). Finally, we systematically imaged the expression pattern of 41 additional TG4 lines generated through our study to document their expression in the adult (Figure S11A) and larval (Figure S11B) CNS as a resource for the community.

Loss- or gain-of-function alleles in GLRA2 cause X-linked neurodevelopmental disorders

Many genes implicated in ASD are also associated with neurodevelopmental disorders beyond autism (Levitt and Campbell, 2009; Sullivan and Geschwind, 2019). Therefore, we asked whether additional variants in genes with disruptive SSC-DNMs could also be responsible for neurological diseases beyond ASD by identifying human subjects with rare, potentially deleterious variants that have not previously been associated with neurological disease (Chong et al., 2015; Gahl et al., 2016; Sobreira et al., 2015). Out of 28 genes in which we identified damaging SSC-DNMs, eight are associated with Mendelian diseases that have neurological presentations documented in OMIM (Amberger et al., 2019; Table 1). For one of these genes (IRF2BPL), we recently reported de novo truncating variants as the cause of a severe neurodevelopmental disorder that presents with abnormal movements, loss of speech, and seizures. This work was done in collaboration with the Undiagnosed Diseases Network (Gahl et al., 2016; Marcogliese et al., 2018). Here, aided by using the online matchmaking software, GeneMatcher (Sobreira et al., 2015), internal human genetics databases, and re-analysis of clinical exome sequencing data, we report variants in GLRA2 as a cause of a neurodevelopmental syndrome with developmental delay (DD), intellectual disability (ID), ASD, and epilepsy (Figure 1H; Table 2).
We identified rare GLRA2 variants in 13 unrelated subjects with or without autistic features. In addition to developmental and cognitive delay of variable severity, 4/13 subjects have microcephaly, 6/13 subjects have a history of epilepsy, and 10/13 subjects have ocular manifestations, including congenital nystagmus that improved with age in three of the cases (Table 2; see GLRA2 subject case histories). Glycine receptor alpha 2 (GLRA2) is an X-linked gene that encodes a subunit of a glycine-gated chloride channel (Zeilhofer et al., 2018). All female subjects harbored DNMs, including a recurring GLRA2\textsuperscript{T296M} \textit{de novo} variant found in 6/8 female subjects. The GLRA2\textsuperscript{T296M} variant was also identified in a female subject in previous large-scale developmental disorder study (Deciphering Developmental Disorders Study, 2017). The five male subjects had inherited GLRA2 variants from unaffected mothers. The mother of subject 12 has a history of learning problems. The CADD scores for all five male subjects are predicted to be damaging (Table 2). Four of the five male subjects had diagnosed or suspected ASD. A maternally inherited microdeletion of GLRA2 was previously reported in a single male patient with ASD (Pinto et al., 2010), indicating that hemizygous LoF allele of this gene in males may cause ASD. Indeed, the \textit{de novo} GLRA2\textsuperscript{N136S} variant present in the SSC in a male subject acts as a LoF allele based on overexpression studies (Figure 4B), which is supported by our behavioral assay on humanized flies, as the GLRA2\textsuperscript{N136S} variant loses the toxic effect on copulation caused by expression of the humanized reference protein (Figure 3G).

To better understand the functional consequences of variants found in our GLRA2 cohort, we generated additional transgenic flies to assay the function of p.T296M (found in six female subjects) and p.R252C (found in a male subject) variants (Figures 5A and S12A). By overexpressing reference and variant GLRA2 using a ubiquitous driver, we found that GLRA2\textsuperscript{R252C} behaves as a LoF allele (Figure 5B), similar to GLRA2\textsuperscript{N136S} (Figure 4B). In contrast, this assay did not distinguish GLRA2\textsuperscript{T296M} from the reference (Figure 5B). Given the recurrent nature of this variant, as well as structural prediction that the residue has a potential role in obstruction of the ion pore in the closed conformation (Figure S12G; Du et al., 2015; Moraga-Cid et al., 2015), we further tested GLRA2\textsuperscript{T296M} and other alleles using additional GAL4 drivers. Using a \textit{par-GAL4} that is expressed in the dorsocentral stripe in the notum, we found that GLRA2\textsuperscript{T296M}, but not the reference or any other GLRA2 variant tested, causes lethality when expressed at high levels (Figure S12B). When we expressed GLRA2\textsuperscript{T296M} at lower levels by manipulating the temperature, we found that this variant induces the formation of melanized nodules in the thorax, a phenotype that we never observe when the reference or other GLRA2 variants are overexpressed (Figures 5C, 5D, and S12B).

To further examine the functional consequences of overexpression of reference and variant GLRA2 in the nervous system, we performed electroretinogram (ERG) recordings on the fly eye expressing human GLRA2 using two distinct drivers. Pan-neuronal driver (\textit{nSyb-GAL4}; Pauli et al., 2008) allows one to express GLRA2 in both pre-synaptic photoreceptors and post-synaptic neurons in the nervous system. Using this driver, we found a significant increase in amplitudes of “OFF” transients with GLRA2\textsuperscript{T296M} (Figures 5E, 5F, S12C, and S12D). This indicates an increase in synaptic transmission (Deal and Yamamoto, 2018), supporting the finding in the notum that p.T296M behaves as a GoF allele. Interestingly, when we limited the expression of GLRA2 to pre-synaptic photoreceptors using \textit{Rh1-GAL4} (Xiong et al., 2012), we did not observe any functional difference between GLRA2\textsuperscript{T296M} and the reference.
DISCUSSION

In this study, we generated >300 fly strains that allow functional studies of human variants and homologous fly genes in vivo. These reagents can be used to study many coding variants that are being identified through next-generation sequencing efforts in the human genomics field in diverse disease cohorts beyond ASD. Our screen elucidated 30 SSC variants with functional differences compared with reference, which was over half (~53%) of the genes in which we were able to perform a comparative functional assay.

Our screen was part of a larger effort to characterize the functional consequences of missense de novo changes from the SSC dataset using different strategies. One approach was based on proteomics by performing yeast two-hybrid assays on 109 SSC-DNMs found in subjects, showing 20% of protein-protein interactions that are found in reference proteins are disrupted by variants (Chen et al., 2018). Another study reported that ~70% of 37 SSC-DNMs knocked into homologous C. elegans genes caused scorable phenotypes (Wong et al., 2019). These studies are complementary to each other because, while some variants have been identified as deleterious by more than one approach (e.g., GLRA2^N136S identified in both worm and fly screens), others are uniquely identified in one study, which could be due to technical limitations. For example, our approach utilizing human cDNA transgenes allowed us to test variant function, regardless of residue conservation in Drosophila. Of the 29 disruptive missense SSC-DNMs identified through our study, 14 affect residues that are conserved in flies and 10 in worms.

To take an unbiased approach, our gene prioritization was only based on gene-level conservation and tool availability (e.g., intronic MiMIC lines and full-length human cDNA) rather than based on gene level constraints and variant-level pathogenicity prediction scores. Hence, our study subset, although somewhat limited, can be considered a random sample of ASD-implicated genes and variants. Interestingly, we could not find any significant difference in pathogenicity prediction for disruptive variants in vivo. It should also be noted that we were limited by the availability of full-length human cDNA, which could select against genes encoding larger transcripts, for which reagents are often harder to obtain. We were able to generate 13 additional TG4 lines that were homozygous viable and successfully crossed back to a Canton-S x chromosome. We assessed their behavior phenotypes in comparison to the reference Canton-S files, which is presented in Figures S4Q–S4T as an additional resource for the community.

Of the 29 missense SSC-DNMs that had functional consequences in our assays, four were not predicted as damaging variants (CADD < 10), nine had moderate scores (CADD: 10–
20), and 16 were predicted to be disruptive (CADD > 20). Understanding how variants that are not predicted to be damaging based on state-of-the-art informatics programs impact protein function may provide guidance to improve the accuracy of in silico tools.

To study the functional consequences of SSC-DNMs, we took two conceptually different approaches (rescue-based humanization strategy and overexpression-based strategy). Indeed, the two approaches were complementary, as only two variants (GLRA2\textsuperscript{N136S} and KCND2\textsuperscript{R86P}) were detected in both screens, showing consistent LoF effects using both approaches. However, it should be noted that variant interpretation is not always straightforward for behavioral analysis. Interestingly, GRK4, NPFFR2, and PDK2 SSC-DNMs were found to be LoF variants when overexpressed ubiquitously, yet these variants were able to rescue lethality in a similar manner to their respective reference alleles (Figure 4; Table S5). This suggests that these variants are partial LoF alleles and different drivers and assays have different sensitivity. Moreover, two disruptive SSC-DNMs, EPHB1\textsuperscript{V916M} and MAP4K1\textsuperscript{M725T}, behaved as complex alleles, displaying discordant phenotypes in the eye and wing. This suggests that these variants may behave in a context-dependent manner, acting as a GoF allele in one tissue while behaving as a LoF allele in another. Thus, one functional assay may not be enough to reveal the full nature of pathogenic mechanisms, and some disease-associated variants may act differently in different tissues or cell types.

It is also important to note that a variety of factors may explain why functional differences are not observed across assays, as different GAL4 lines may have variable developmental timing, strength, and context dependency. In addition, the presence of the endogenous protein in cells and the cell-type-specific effect of exogenous protein may also contribute to our functional readout. Therefore, variant annotation in Drosophila should be supplemented by deeper characterization of the loss-of-function mutant and gene-expression studies to provide the clearest supportive evidence to a molecular diagnosis.

Starting from a single de novo hemizygous missense variant that we identified as a LoF allele in GLRA2 (p.N136S), we identified a cohort of subjects with overlapping neurodevelopmental phenotypes carrying LoF or GoF variants in this gene. Interestingly, female subjects harbored DNMs and male subjects carried maternally inherited variants in this X-linked gene, which undergoes random X inactivation in females, but not in males (Barakat and Gribnau, 2012). The X-linked status of GLRA2 may mean that variants causing reduced GLRA2 activity lead to disease in males but can be tolerated in heterozygous females. This is supported by asymptomatic mothers of male probands who had maternally inherited alleles (subjects 9–11 and 13). In contrast, GoF variants in this channel could be overrepresented in females since hyperactivation of this channel may cause neurological defects (Zhang et al., 2017). Of the eight female subjects, six carried the identical recurring DNM, p.T296M. None of the variants observed in females are present in control databases, arguing strongly that these variants are pathogenic. While the exact mechanism of how the p.T296M variant affects GLRA2 function remains unclear, the presence of melanized nodules in flies expressing this variant are indicative of an innate immune response (Dudzic et al., 2019), potentially as a result of leaky ion channel function (Feske et al., 2015). Fittingly, our structural analysis revealed that the p.T296 residue is adjacent to a critical amino acid that is likely important for keeping the ion pore in a conformationally closed state (Figure S12G).
In summary, we utilized a model organism-based *in vivo* functional genomics approach to study the functional consequence of rare genetic events in a common neurological disorder, ASD. In addition to garnering variant functional data for ASD subjects in the SSC, we leveraged this information to identify and document a rare neurological condition through matchmaking and collaboration. Such bidirectional communication and collaboration between bench scientists and clinicians greatly facilitates functional studies of human variants found in common diseases, such as ASD, and can also lead to novel discoveries that have an impact on rare-disease research.

**Limitations of the study**

Although this study revealed a number of rare variants found in ASD patients that have functional consequences, there are several caveats to recognize based on the design and assays used in this screen. First, it is not clear for the majority of the 30 hits found by the screen whether the disruptive variants are truly pathogenic and directly contribute to ASD. Identification of additional patients with similar genotypes and phenotypes will be necessary to establish a causal relationship between these variants and ASD pathogenesis. Second, we do not argue that this screen was able to identify all variants that had functional consequences, considering most variants were shown to be deleterious based on one phenotypic assay. Hence, if one performs additional assays in different biological contexts, more variants with altered function may be discovered. Third, it is not clear how many phenotypic assays are required to determine whether a variant has functional consequences and which phenotypic assays in flies are more relevant to complex human phenotypes, such as ASD. Additional studies of variants identified as having functional consequences in an invertebrate model organism should be followed up using mammalian models or human cells, tissues, and organoids to assess whether the deleterious variants affect biological processes that relate to ASD. Fourth, although the majority of the hits were alleles of genes in which the fly homolog is expressed in the nervous system, some were in genes in which we failed to detect any expression in this organ system in *Drosophila*. This could be because some of these genes are expressed at low levels that are beyond the detection limit of our assay systems; only expressed at a specific time point during neural development; expressed in the nervous system of humans, but not flies; or they may be contributing to neurological phenotypes through their function in non-neural organ systems. Additional gene expression and functional studies will be required to fully understand their mechanistic contributions to ASD. Fifth, although we were able to identify some deleterious variants that affect fly behavior, which is the most relevant phenotype to ASD out of all assays we performed in this screen, we have not been able to assess this for all variants due to technical limitations. In order to perform clean behavioral experiments, one must control the genetic background since this could be a significant confounding factor. Although we tried to eliminate some of the genetic variability by inserting the reference and variant transgenic constructs into the same genomic location and swapping out mutant chromosomes (*y w*) that are known to affect behavioral outcomes, we did not isogenize all chromosomes through multiple rounds of back crossing to facilitate the speed of our screen. Hence, additional behavioral studies performed on a more standardized genetic background (e.g., cantonized flies) will likely provide additional information regarding the role of these genes in fly behavior, which could provide additional insights into their links to ASD. Finally, we would like to emphasize...
that this work should be considered as a pilot screen rather than a comprehensive screen of de novo missense variants identified in the SSC. The genetic variants that we were able to study in our screen were limited by the availability of intronic MiMIC elements in fly genes because recombinase-mediated cassette exchange (RMCE) was the only efficient way to generate TG4 lines at the time of project initiation. With the advent of CRISPR-mediated integration cassette (CRIMIC) and KozakGAL4 insertion technologies (Lee et al., 2018; Kanca et al., 2021), virtually any fly gene is now targetable to generate a strong LoF allele with GAL4 expression that allows for subsequent humanization experiments. Therefore, future screening strategies could employ these and other emerging techniques to assess functional consequences of many rare missense variants found in ASD or other disease patients.

STAR★METHODS

RESOURCE AVAILABILITY

Lead contact—Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Shinya Yamamoto (yamamoto@bcm.edu).

Materials availability—The fly lines generated in this study have been deposited to the Bloomington Drosophila Stock Center (BDSC). Please see key resources table for unique identifiers. Please contact the Lead Contact for further information.

Data and code availability

• Confocal imaging movies of neuron/glia colocalization with TG4>UAS-nls.mCherry have been deposited at Mendeley Data and are publicly available as of the date of publication. The DOI is listed in the key resources table. De-identified GLRA2 variant data have been deposited at ClinVar. They are publicly available as of the date of publication. Accession numbers are listed in the key resources table.
• This study didn’t generate any code.
• Any additional information required to reanalyze the data reported in this work paper is available from the Lead Contact upon reasonable request.

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Generation of TG4 lines—All TG4 alleles in this study were generated by ϕC31-mediated recombination-mediated cassette exchange of MiMIC (Minos mediated integration cassette) insertion lines (Gnerer et al., 2015; Nagarkar-Jaiswal et al., 2015; Venken et al., 2011). Conversion of the original MiMIC element was performed via genetic by crossing UAS-2×EGFP, hs-Cre, vas-ϕC31, Trojan T2A-GAL4 triplet flies to each MiMIC strain and following a crossing scheme (Diao et al., 2015). 73 TG4 lines were described previously but not extensively characterized (Lee et al., 2018), while 35 lines were generated specifically for this study.

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**Generation of UAS-human cDNA lines**—The majority of reference human cDNA clones were obtained in either pDONR221 or pDONR223 donor vectors. The LR clonase II (ThermoFisher) enzyme was used to shuttle ORFs into the p.UASg-HA.attB destination vector via Gateway™ cloning. Some ORFs that were not Gateway compatible were obtained from additional sources (Table S2), amplified with flanking attB sites and cloned into pDONR223 plasmid using BP clonase II (ThermoFisher). Sequence-verified variants were generated in the DONR vectors by either site-directed mutagenesis (SDM) or High-Throughput Mutagenesis (HiTM) as previously described (Tsang et al., 2016). SDM was performed with primers generated using NEBaseChanger (Table S3) with the Q5® mutagenesis kit (NEB). Sequence-verified reference and variant ORFs in the pUASg-HA.attB destination plasmid were microinjected into ~200 embryos in one three attP docking sites (attP86Fb, VK00037 or VK00033) docking sites by ϕC31 mediated transgenesis (Bischof et al., 2007; Venken et al., 2006). The docking site of choice were selected based on the genomic locus of the corresponding fly gene. In principal, VK00037 docking site on the 2nd chromosome was used for human genes that correspond to fly genes on the X, 3rd or 4th chromosome, whereas VK00033 or attP86Fb docking site on the 3rd chromosome was used for human genes that correspond to fly genes on the 2nd chromosome.

**Fly husbandry**—Unless otherwise noted, all flies used in experiments were grown in a temperature and humidity-controlled incubator at 25°C and 50% humidity on a 12-hour light/dark cycle. Some experiments were conducted at different temperatures that are specifically indicated in the text and figures. Stocks were reared on standard fly food (water, yeast, soy flour, cornmeal, agar, corn syrup, and propionic acid) at room temperature (~22°C) and routinely maintained.

**Fly stocks used that were not generated here**—tub-GAL4 (y¹ w*; P[w=+mC]=tubP-GAL4]LL7/TM3, Sb¹ Ser¹) BDSC_5138, GMR-GAL4 (w*; P[w=+mC]=GAL4-ninaE.GMR]12) BDSC_1104, nub-GAL4 (P[GawB]nubbin-AC-62) (Calleja et al., 2000), nSyb-GAL4 (y¹ w*; P[nSyb-GAL4.S]3) BDSC_51635, Rh1-GAL4 (P[ry+i+7.2=erh1-GAL4]3, ry[506]) BDSC_8691, pnr-GAL4 (y¹ w¹¹¹8; P[w=+mW.hs]=GawB]pnr[MD237]/TM3, P[w=+mC]=UAS-y.C]MC2, Ser¹) BDSC_3039, UAS-nlsGFP (w¹¹¹8; P[w=+mC]=UAS-GFP.nls]14) BDSC_4775, and UAS-nls.mCherry (w*; P[w=+mC]=UAS-mCherry.NLS]3) BDSC_38424.

**Patient recruitment and consent**—Affected individuals were investigated by their referring physicians at local sites. Prior to research studies, informed written consent for testing and publication was obtained according to the institutional review boards (IRB) and ethics committees of each institution. Individuals who were ascertained in diagnostic testing procedures (and/or their legal guardians) gave clinical written informed consent for testing, and their permission for inclusion of their anonymized data in this cohort series. This was obtained using standard forms at each local site by the responsible referring physicians.
GLRA2 subject case histories—Subject 1 is an 8 years old female with global developmental and cognitive delay. Pregnancy was naturally conceived and uncomplicated, other than decreased fetal movements noted by the mother. She was delivered at term (39 weeks gestational age) via c/section due to breech presentation. Birth weight was 3,600 grams. Neonatal period was uneventful. There were no feeding difficulties and her growth remained within the normal limits. She was delayed with all her milestones but most significantly for speech (walked at 18 months, scribbled with a crayon at 3.5 years, first words at 24 months and combined words to sentences at 4–5 years of age). She was diagnosed with mixed expressive-receptive speech delay and received speech therapy, occupational therapy and physical therapy interventions. In school she exhibits learning problems, inattention and is below her grade level. She has a modified curriculum and is receiving resources in reading and math. There is no history of developmental regression or seizures. The medical history is otherwise significant for nystagmus that was first noted in infancy and improved with age, as well as myopia and astigmatism requiring corrective glasses. Family ethnicity is Hispanic and the family history is non-contributory. The patient had a normal brain MRI (magnetic resonance imaging) at 6 months of age. EEG (electroencephalography) at 4 years of age showed a slow and poorly formed background, indicative of mild encephalopathy, but did not detect epileptiform activity. Genetic testing included: mitochondrial DNA sequencing which detected a pathogenic variant m.13042 G>A though at heteroplasmy level of 1.9%, which was felt unlikely to explain the phenotype. CMA (chromosomal microarray) was negative. Trio whole exome sequencing (WES) detected a de novo, heterozygous variant of unknown clinical significance in GLRA2, c.887C>T, p.Thr296Met (NC_000023.10, chrX: g.14627284C>T). This variant is absent in gnomAD. More recent clinical reanalysis of exome data did not detect any other candidates that may explain the phenotype.

Subject 2 is a 6 years old female with epilepsy, developmental delay (DD), mild intellectual disability (ID) and autism spectrum disorder (ASD). Pregnancy was uncomplicated and she was delivered at term (41 weeks gestational age) via vaginal delivery with vacuum extraction. The neonatal period was uneventful. At the age of 6 months, she developed a severe epileptic encephalopathy with myoclonic seizures. Seizure control was achieved with medications, and she has been seizure-free without medications since the age of about two years old. Delayed psychomotor development was noted, most significantly for her speech with a mixed expressive-receptive speech delay (non-verbal). Her ability to concentrate is poor and she displays mood swings. The medical history is otherwise significant for nystagmus that was first noted in infancy (6 weeks old) and improved with age, and sleep disturbance. She has mild microcephaly [< 1st centile: −2.84 standard deviation (SD)] and mild bilateral cutaneous 3rd-4th syndactyly, with no other congenital anomalies. Family ethnicity is European (German/Italian) and the family history is significant for a maternal aunt that had epilepsy in adulthood but her cognitive development was normal. Brain MRI showed delayed myelination at 7 months old and a small arachnoid cyst. EEG was abnormal for bilateral synchronized, sometimes high amplitude spike/polyspike-waves-complexes, and bitemporo-occipital hints for severe functional defects with epileptic potentials. Chromosomal analysis, Angelman syndrome methylation study, epilepsy next generation sequencing (NGS) gene panel and MECP2 sequencing were negative. Trio WES...
detected a *de novo*, heterozygous variant of unknown clinical significance in *GLRA2*, c.887C>T, p.Thr296Met (NC_000023.10, chrX: g.14627284C>T). This variant is absent in gnomAD.

Subject 3 is a 5 year and 6 months old female with DD, microcephaly, abnormal eye movements and ataxic gait. Pregnancy was uncomplicated and she was born at term via c/section. Abnormal eye movements were noticed two weeks after birth, during hospitalization due to a lower respiratory tract infection. At the age of 6 months, clinical examination revealed mildly delayed developmental milestones and erratic conjugate eye movements akin to opsoclonus. At age 4 years OFC (occipitofrontal circumference) was 43 cm (<1st centile: −4.28 SD) and ophthalmological evaluation revealed alternating exotropia, for which patching therapy was initiated. Language was limited to a few words and neuropsychological evaluation documented moderate developmental delay (Bayley-III). The patient could walk unsupported with ataxic gait. At age 5 years 6 months, erratic eye movements were considerably reduced and she could walk independently but her expressive language was still limited to a few words, with delayed receptive speech and nonverbal communicative skills. Family ethnicity is European and the family history is unremarkable. Brain MRI at 6 months of age showed mild cortical atrophy with thinning of the corpus callosum. EEG, while awake and asleep, laboratory and metabolic investigations were unremarkable. Array-CGH (comparative genomic hybridization) highlighted a maternally inherited 3q25.32 duplication (chr3:157746089–158324659, hg19) that was interpreted as likely benign. Trio WES detected a *de novo* heterozygous variant in *GLRA2*, c.887C>T, p.Thr296Met (NC_000023.10, chrX: g.14627284C>T). This variant is absent in gnomAD. In addition, it detected a *de novo* variant in *CACNA1B*, c.5381C>T, p.Thr1794Met (NC_000009.11:g.141000212C>T), which is a variant of unknown significance in a gene that is linked to an autosomal recessive condition (Neurodevelopmental disorder with seizures and nonepileptic hyperkinetic movements, MIM #618497). Failure to identify a second allele in this gene reduces the likelihood that this variant is responsible for this patient’s phenotype.

Subject 4 was a female infant with seizures and severe developmental delay who passed away at 7 months of age secondary to complications of COVID-19 infection. Pregnancy was uneventful and she was born at term (40 weeks gestational age). She was noted to have focal seizures at 2–3 weeks of age, and was diagnosed with infantile spasms when she was 5 months old. At 6 months of age she was not reaching for objects, not sitting up and only making high-pitched sounds. She had borderline microcephaly with dysmorphic features including midface retrusion, apparent hypotelorism, deep set eyes, thick eyebrows, downturned corners of the mouth, and wide-spaced nipples. Family ethnicity is Hispanic and the family history is non-contributory. She had normal plasma and CSF (cerebrospinal fluid) lactate, piperolic acid and piperideine-6-carboxylate, ammonia, urine organic acids, plasma amino acids, acylcarnitine profile, and CSF amino acids. An Epilepsy gene panel was non-diagnostic. Trio WES detected a *de novo* heterozygous variant in *GLRA2*, c.887C>T, p.Thr296Met (NC_000023.10, chrX: g.14627284C>T). This variant is absent in gnomAD.

Subject 5 is a 5 years and 4 months old female with global developmental delay. She was born in Afghanistan to consanguineous parents, and there is limited information available
regarding her birth history. Pregnancy was uneventful. She walked at 2 years and is currently non-verbal. She has ptosis and oculomotor apraxia. On physical exam she is noted to have broad halluces. Brain MRI at 5 years old was normal. Metabolic testing was non-revealing. Array-CGH, molecular genetic testing of \textit{FMR1} and screening for congenital disorders of glycosylation resulted negative. WES detected a \textit{de novo}, heterozygous variant of unknown clinical significance in \textit{GLRA2}, c.887C>T, p.Thr296Met (NC_000023.10, chrX: g.14627284C>T). This variant is absent in gnomAD.

Subject 6 is a 9 months old female with developmental delay, West syndrome, microcephaly, up-beat nystagmus and myopia. She was born at term (38 gestational weeks, birth weight 2770 g (10\textsuperscript{th} percentile), OFC 34 cm (33\textsuperscript{rd} percentile)) via c/section due to breech presentation. She had postnatal respiratory distress. Abnormal eye movements were noticed at 6 weeks of age. Ophthalmologic assessment was normal with normal fixation apart from up-beat nystagmus. Clinical evaluation at 4 months of age revealed developmental delay and an OFC of 38.5 cm (3rd percentile). At approximately 7 months of age, she developed infantile spasms with an intermittent hypsarrhythmic pattern in EEG (West syndrome). Therapy with Sulthiame resulted in resolution of seizure activity and EEG normalisation. She has made developmental progress since starting treatment with antiepileptic medications. At 9 months of age she is showing mild gross motor delay. Family ethnicity is European, and the family history is non-contributory. Brain MRI, abdominal ultrasound, and metabolic screening labs (including plasma amino acids, acylcarnitine profile, CSF analysis and CSF neurotransmitters) were normal. Trio exome sequencing detected a heterozygous \textit{de novo} missense variant in \textit{GLRA2} (NM_002063.4), c.887C>T, p. Thr296Met (NC_000023.10, chrX: g.14627284C>T, hg19). This variant is absent in gnomAD.

Subject 7 is a 6 years and 7 months old female with a history of infantile spasms, epilepsy and intellectual disability. She was born at term and first presented with infantile spasms at 3 months of age. This evolved to atonic and tonic-clonic seizures as she grew up. She was delayed with all milestones (walked at 4.5 years old and remains non-verbal). She had nystagmus that improved with age and strabismus. The medical history is otherwise significant for hyperactivity, inattention and sleep disturbance. Her ethnicity is African (Senegal). Brain MRI at 3 years of age showed cortical and white matter atrophy, including vermian atrophy. EEG showed hypsarrhythmia at onset and she had normal interictal EEG afterwards. She had normal SNP (single nucleotide polymorphism)-array, negative targeted epilepsy panel and negative metabolic lab results. Trio WES identified a \textit{de novo} heterozygous variant in \textit{GLRA2} (c.140T>C, p.Phe47Ser (NC_000023.10, chrX: g.14550432T>C). This variant is absent in gnomAD.

Subject 8 is a 5 years and 3 months old female with mild developmental delay and learning disabilities. Pregnancy was naturally conceived and uncomplicated. She was born near term following premature rupture of membrane. Birth weight was 3,280g, birth length was 50cm and OFC was 33.5cm. Apgar scores was 8/9 at 1\textsuperscript{st}/5\textsuperscript{th} minutes of life. She had feeding difficulties during the neonatal period. With regards to her milestones, she was able to sit unsupported at 9 months and walked independently at 17 months. Speech was not delayed but she has difficulties in pronunciation and articulation. There is no history of
developmental regression. She has a School aide for learning difficulties but is interacting well with other children. She is receiving therapy for fine motor difficulties and is followed by a psychiatrist with a diagnosis of infantile psychosis. She also has a history of sleep disturbance (short sleep duration). The family history is non-contributory, she is the only child to her parents as a couple and has two paternal half-siblings that are healthy. Her growth is normal, with weight at 22 kg (+1.1 SD), height at 117.5 cm (+1.4 SD) and OFC 51 cm (+0.2 SD). On exam she has mild dysmorphic features, including long face, pointed chin, and overlapping 1st and 2nd toes. Although she does not have clinical seizures, EEG was abnormal for left fronto-temporal spike-waves focus which diffuses in the right frontal region, activated by sleep but not meeting criteria for Epilepsy with continuous spike-wave during sleep (CSWS). Brain MRI showed nonspecific dot-like hypersignal in FLAIR of the subcortical white matter of the frontal region. Fragile X testing was negative. Exome sequencing detected a de novo heterozygous variant in GLRA2, c.777C>G, p.Ile259Met (NC_000023.10, chrX: g.14627174C>G). This variant is absent in gnomAD.

Subject 9 is an 11 months old male with hypotonia, DD and dysmorphic craniofacial features. Pregnancy was uncomplicated, he was delivered at term (38 and 3/7 weeks gestational age) and the neonatal period was uneventful. Soon after birth dysmorphic features were noted, including an elongated face, high anterior hairline, epicanthal folds, downslanting palpebral fissures and a bulbous nose. Growth remains within the normal limits. His medical history is otherwise significant for obstructive sleep apnea and strabismus. Family ethnicity is European (Dutch) and the family history is significant for the maternal grandfather who has not further specified unexplained neurological complaints, and which could not be further investigated. Investigations for metabolic disorders, Fragile X syndrome and a SNP-array were normal. Trio WES identified a rare variant in GLRA2, c.754C>T, p.Arg252Cys (NC_000023.10, chrX: g.14627151C>T), which was inherited from mother. No other possible disease explaining variant was identified. The mother displayed skewed X chromosome inactivation (82% on two measurements). The variant was absent in the maternal uncle and the maternal grandmother, but was inherited from the maternal grandfather, who was not available for clinical investigations. His level of functioning remains unknown. This variant is present in one heterozygous female in gnomAD.

Subject 10 is a 7 years old male with epilepsy, DD with regression, and ASD. Pregnancy was uncomplicated. He was born full term via uncomplicated delivery, and was meeting his early developmental milestones. He was speaking in sentences at 2.5 years old when he started having generalized tonic-clonic seizures. He developed staring spells, ataxia, and an increased frequency of myoclonic jerks, which around the age of 6 years old were occurring 20 times per day on average, with 5–6 atonic seizures per day each lasting less than 30 seconds. Following seizure onset, he experienced developmental regression. At 3 years of age he was diagnosed with ASD. At 6 years of age his vocabulary was about 20 words, with gains in development lost following significant seizures. His ethnicity is European, and the family history is significant for a younger brother with ASD, although he has not presented with seizures. Neither mother nor father have a history of seizures or delays. At age 3, EEG depicted generalized slowing and generalized epileptiform discharges associated with myoclonic jerks. MRI showed minimal increased T2 signal intensity on the occipital
lobes that was thought to be within normal limits. Genetic testing for Fragile X syndrome, Prader-Willi/Angelman syndromes, and congenital disorders of glycosylation was normal. Additional tests, including plasma amino acids, lysosomal enzymes, and cerebral creatine deficiency were also normal. Array-CGH reported a maternally inherited 1p33 deletion of unknown significance (chr1: 48,688,391–49,922,153). The patient was enrolled to The Manton Center for Orphan Disease Gene Discovery Core protocol. Trio WES identified a maternally inherited variant in GLRA2, c.862G>A, p.Ala288Thr (NC_000023.10, chrX: g.14627259G>A). This variant is absent in gnomAD.

Subject 11 is a 35 years old male with a history of DD, learning disabilities and ASD. Pregnancy was uncomplicated and he was born at term (40 weeks gestational age). Since early childhood he showed slow movement and difficulties in motor coordination. He walked and said his first words at 24 months, and first sentences at 3 years of age. In school learning disabilities were noted, including difficulties in writing, reading, praxias, temporal orientation, calculation, drawing, and visuo-spatial organization. He graduated high school and continued to higher education, though he did not complete a degree. Neuropsychiatric assessment in adulthood was consistent with ASD and social and cognitive deficits. There is no history of seizures. The medical history is otherwise significant for environmental allergies, myopia, and astigmatism. The ethnicity is European, and the family history is non-contributory. The patient had a normal brain MRI at 29 years old. Trio WES identified a maternally inherited variant in GLRA2, c.1186C>A, p.Pro396Thr (NC_000023.10, chrX: g.14748434C>A). This variant is present in 3 heterozygous females and 1 hemizygous male in gnomAD.

Subject 12 is a 15 years old male with a history of epilepsy, DD, expressive language disorder, and ASD. Pregnancy was uncomplicated and he was born at term. He walked at 17 months and said his first words at 13 months. Following febrile convulsions at 18 months of age, more pronounced developmental regression was noted. At 7 years of age, he had focal (partial) motor seizures for which he still receives antiepileptic medications. In school he is enrolled in mainstream classes, but he has learning problems and inattention, and is behind his age-peers. Speech remains delayed, he has limited vocabulary and receives speech therapy. He is frequently agitated and has social anxiety. The medical history is otherwise significant for obesity. The ethnicity is European, and the family history is significant for learning problems in his mother. On exam, he is noted to have mild dysmorphic features, including broad face, Widow’s peak, horizontal and broad eyebrows, long prominent eyelashes, and a broad nasal tip. Brain MRI at 7 years of age showed increased signal intensity in FLAIR of the cortical matter of the parietal region. EEG at that age showed abnormal alpha waves with high polymorphic spikes with a focus in the right temporal region (consistent with the clinical presentation of partial simple seizures). Genetic testing for Fragile X syndrome, and metabolic labs were normal. Trio WES identified a maternally inherited variant in GLRA2, c.1199C>T, p.Pro400Leu (NC_000023.10, chrX: g.14748447C>T). This variant is absent in gnomAD. × chromosome inactivation study in the mother, who is mildly affected, showed moderately unbalanced × inactivation (60:40) in two independent experiments.
Subject 13 is a 6 years old male with intractable epilepsy, developmental delay, and suspected autism spectrum disorder. Pregnancy was uncomplicated and he was delivered at term (39 and 2/7 weeks gestational age, weight: 3440 g (−0.19 SD), length: 49 cm (−1.34 SD), OFC: 35 cm (−0.3 SD)). The neonatal period was uneventful. He learned to walk at the age of 21 months. He spoke his first words at 18 months, but his speech development was delayed. At the age of 4 years and 3 months he first presented with generalized tonic-clonic and tonic seizures. At most recent evaluation he was reported to have 4–5 generalized tonic-clonic or tonic seizures, and absence seizures daily (while being treated with five anti-epileptic medications simultaneously). EEG at age 5 years and 6 months showed excessive beta-activity most likely due to medication; left and right posterior and right frontal Intermittent slowing; and bilateral polyspikes during sleep with a frequency of < 1/minute three times associated with tonic-clonic seizures. Brain MRI at the same age showed no abnormalities. Concomitant to the epilepsy, he was diagnosed with DD with a focus on speech development and cognitive impairment. He is presenting with poor expressive speech (partly able to speak in sentences). The behavioral abnormalities include a short attention span, poor impulse control, sleep disturbances and recently also some compulsive traits have been reported by the mother. An early-childhood autism spectrum disorder is suspected, and an evaluation has been initiated. He visits an integrative Kindergarten with permanent one-on-one care and receives speech therapy, occupational therapy, and physiotherapy. Growth parameters at the last assessment (5 years and 11 months) were normal (weight at 0.51 SD, height at 1.12 SD and OFC at −0.72 SD). On clinical exam, he was noted to have downslanting palpebral fissures and a high forehead. There is no family history of developmental delay and epilepsy. Neonatal metabolic screening was unremarkable. Genetic testing for Fragile × syndrome as well as array-CGH, karyotype and proband exome sequencing were non-revealing. Trio exome sequencing on a research basis identified a maternally inherited hemizygous missense variant in GLRA2 c.1334G>A, p.Arg445Gln (NC_000023.10, chrX: g.14748582G>A). This variant is present in three heterozygous females in gnomAD and has not been observed in a hemizygous state. With Sanger sequencing, it was shown that the variant is absent in the unaffected maternal half-brother of subject 13.

METHOD DETAILS

Ortholog candidate identification—We utilized DIOPT (Hu et al., 2011) to determine the fly ortholog of the 1519 human genes from the SSC. DIOPT scores lower than 4/16 were excluded. If there were multiple fly paralogs with equal DIOPT scores we referred to the weighted score. If the weighted scores were equal, we chose the ortholog in which a Gold MiMIC present in the other paralogs of equal strength.

Electroretinograms (ERG)—ERG recordings on adult flies were performed on nSyb-GAL4 (Pauli et al., 2008) and Rh1-GAL4 (Xiong et al., 2012) driven UAS-GLRA2 at 5 days post-eclosion raised at 25°C in 12h light/12h dark cycle as previously described (Verstreken et al., 2003) using LabChart software (AD instruments). 4–10 flies were examined for each genotype. Recording was repeated at least 3 times per fly. Quantification and statistical analysis was performed using ANOVA followed by Bonferroni’s multiple comparison test using Prism 8.0.
Complementation test of lethality in TG4 lines—Out of the 108 TG4 mutants generated, 65 TG4 mutants were homozygous lethal. Because lethality can be caused by disruption of the gene of interest or due to second site lethal mutations carried on the same chromosome, we performed complementation test using standard methodology. For genes on the 2nd and 3rd chromosome, female heterozygous TG4 lines balanced with either SM6a or TM3, Sh, Ser, respectively, were crossed with male flies carrying a corresponding deficiency (Df) that covers the gene of interest (see Table S4). Three independent crosses were set at 25°C for each TG4 line and we determined if any TG4 flies survived to the adult stage in trans with their corresponding Df (TG4/Df). If viable, a second Df line covering the same gene was used to validate this finding to make sure the complementation is not due to some problematic Df lines. If TG4 was viable over two independent Df lines, we ascribed the lethality to a second site mutation on the TG4 chromosome. TG4 that remained lethal in trans with a Df line are be considered to be disrupting an essential gene in flies. For five genes on the X-chromosome of the fly, complementation was performed by first rescuing hemizygous TG4 males with a duplication (Dp) line obtained from BDSC (Table S4), and crossing these rescued flies to female TG4/FM7 flies. If TG4/Y; Dp/+ lines were viable, we ascribed the lethality of TG4 to the gene of interest. All Df and Dp lines were obtained from BDSC, and the specific stock used in our analysis are listed in Table S4.

Through this experiment, we found 65 TG4 mutant lines that were homozygous lethal, and 47 of 65 remained lethal when in trans with a corresponding deficiency line (Figure S1; Table S4). The 47 essential genes in D. mel corresponded to 60 SSC related human genes (Figure S1). The lethality of 18 TG4 lines corresponding to 19 human genes were due to a second site lethal mutation, potentially present in the original MiMIC line, or introduced during RMCE which has been reported previously (Nagarkar-Jaiswal et al., 2015). These TG4 lines together with viable TG4 lines are likely associated with non-essential genes in Drosophila.

Rescue of lethality in TG4 lines by UAS-human cDNA transgenes—In order to assess the ability of human reference or SSC variant cDNAs to rescue lethality observed in TG4 mutants in essential genes, we first double balanced all Df lines that fail to complement a lethal TG4 line with UAS-reference or variant cDNA lines. For genes on the 2nd chromosome, we generated Df/CyO; UAS-cDNA/(TM3, Sh, Ser) stocks. For genes on the 3rd chromosome, we generated UAS-cDNA/(CyO); Df/TM3, Sh, Ser. Heterozygous TG4/Balancer females were crossed to double balanced Df/Balancer, UAS-human cDNA males at multiple temperatures (18°C, 22°C, 25°C, 29°C) to determine rescue of lethality to adult stage. A minimum of two independent crosses were conducted at each temperature. For the five genes on the X-chromosome of the fly, we attempted rescue by crossing female TG4/FM7 flies to UAS-cDNA/(SM6a) males to generate hemizygous TG4 males that expresses human cDNA (TG4/Y; UAS-cDNA+) to test their viability. Statistical analysis was performed using Two-way ANOVA followed by Sidak’s multiple comparison test across temperature and genotype.

Lifespan assays—Lifespan analysis was performed as previously described (Chung et al., 2020). Briefly, newly eclosed flies were separated by genotype and sex and incubated at
25°C. Flies were transferred into a fresh vial every two days and survival was determined once a day. 11–49 flies were tested per group. Statistical analysis was performed using Log-rank (Mantel-Cox) test.

**Behavioral assays**—Of 61 TG4 mutants that were viable when in trans with a corresponding deficiency, 18 lines exhibited lethality in homozygous states, indicating the presence of a second site lethal mutation. Out of 43 TG4 mutants that were homozygous viable, we prioritized to study 21 TG4 mutants based on reagent availability. Courtship assay was performed as previously described (Guo et al., 2019). Due to the reported effects of the y¹ w* mutations on behavior (Krstic et al., 2013; Massey et al., 2019), we replaced the X-chromosome containing y¹ w* with the y+ w+ X-chromosome from a Canton-S strain. This Canton-S strain was provided by Dr. Hugo Bellen and has been in his stock collection since the 1980s. Please see Figure S3 for the crossing strategy. Collection of socially naive adults was performed by isolating pupae in 16 × 100 polystyrene vials containing approximately 1 mL of fly food. After eclosion, flies were anesthetized briefly with CO₂ to ensure they were healthy and lacking wing damage. Anesthetized flies were returned to their vials and allowed 24 hours to recover before testing. Courtship assays were performed in a 6 well acrylic plate with 40mm circular wells, with a depth of 3 mm and a slope of 11 degrees, as per the chamber design (Simon and Dickinson, 2010). One Canton-S virgin female (6–10 days post-eclosion), and one TG4 mutant male fly (3–5 days post-eclosion) with or without UAS-human cDNAs were simultaneously introduced into the chamber via aspiration. Recordings were taken using a Basler 1920UM, 1.9MP, 165FPS, USB3 Monochromatic camera using the BASLER Pylon module, with an adjusted capturer rate of 33 fps (frames per second). Conversion of captured images into a movie file was performed via a custom MatLab script, and tracking of flies in the movie was performed using the Caltech Flytracker (Eyjolfsdottir et al., 2014). Machine learning assessment of courtship was performed using JAABA (Kabra et al., 2013) using classifiers that scored at 95% or higher accuracy during ground-truthing trials. At least 10 animals were tested per genotype. Analysis of data was performed using Excel (Microsoft) and Prism (GraphPad). A ROUT (Q=1%) test was performed in Prism to identify outliers. Determination of significance in behavior tests was performed using the Kruskal-Wallis one-way analysis of variance and the Dunn’s multiple comparison test. P-values of 0.05 or less were considered significant.

**Overexpression assays to assess lethality and morphological phenotypes**—To detect any differences in the phenotypes induced by overexpression of reference and variant human cDNA in order to assess variant function, we crossed UAS-human cDNAs with reference or variant alleles to ubiquitous (tub-GAL4) (Guelman et al., 2006), wing (tub-GAL4) (Calleja et al., 2000) or eye (GMR-GAL4) (Mehta et al., 2005) specific drivers. In the ubiquitous expression screen, 3–4 virgin females of tub-GAL4/TM3 Sb flies were crossed to 24 males of the UAS-cDNA reference and variant at 25°C. After 3–4 days, the parents were transferred into new vials, and the new vial was placed at 29°C while the old vial was kept at 25°C, allowing us to test two temperatures simultaneously. The parents were discarded after 3–5 days. Flies were collected after most of the pupae eclosed. The total number of flies were counted and scored with the genotype of interest (i.e. tub-GAL4>UAS-cDNA) as well as all other genotypes, (i.e. genotypes with balancers). A
minimum of 10 flies were scored per experiment, though for the majority of crosses 50–100 flies were scored in this primary analysis. Viability was calculated by taking the % of observed/expected based on Mendelian ratio, and any UAS-cDNA with survival less than 70% was recorded as having scorable phenotype (lethal or semi-lethal). All of lines showing a phenotype at 29°C also showed phenotypes at 25°C, so subsequent experiments were performed at 25°C. To validate our hits, we performed the same viability assay, except each UAS-cDNA was tested at least three times to statistically validate that there is a difference between reference and variant. In addition, two independent UAS-cDNA transgenic lines established from the same construct were tested for each reference and variant. A variant was considered to have functional consequence (true hit) if both transgenic lines showed the same phenotype. In the cases where the difference is rather minor (e.g. <20% difference between survival), this was considered within the variation of the experiment paradigm, and the variant phenotype was documented. Functional study using wing or eye drivers were performed using similar strategies, but morphological phenotypes were scored instead of lethality.

**Imaging of adult fly morphology**—*Drosophila* eyes, wings and nota (dorsal thorax) were imaged after flies were frozen at −20°C for at least 24 hours. Wings for some flies were dissected in 70% EtOH and mounted onto slides for imaging. Images were obtained with the Leica MZ16 stereomicroscope equipped with Optronics MicroFire Camera and Image Pro Plus 7.0 software to extend the depth-of-field for Z-stack images.

**Expression analysis of TG4 lines in larval and adult brains**—All TG4 lines are crossed with *UAS-nls.mCherry* (3rd chromosome) or *UAS-nls.GFP* (3rd chromosome) at room temperature. Note that the nls.GFP that is being used here is prone to leak outside the nucleus while nls.mCherry is retained in the nucleus. The brains of mCherry/GFP positive third instar larvae and 3–5 days old adult flies were dissected in 1X phosphate-buffered saline (PBS). Adult brains were fixed immediately in 4% paraformaldehyde (PFA) and incubated at 4°C overnight (o/n) on a shaker. Next day these brains were post-fixed with 4% PFA with 2% Triton-X in PBS (PBST), kept in a vacuum container for an hour to get rid of the air from the tracheal tissue also make the tissue more permissive. Fixative was replaced every 10 minutes during this post-fixation step. Larval brains were fixed for 50 minutes on a rotator at room temperature. After thorough washing with PBS with 0.2% Triton (PBTX) both adult and larval brains were incubated with primary antibodies overnight (o/n) at 4°C on a shaker. The sample were extensively washed with 0.2% PBTX before secondary antibodies were applied at room temperature for 2 hours. Samples were thoroughly washed with PBST and mounted on a glass slide using Vectasheild (Vector Labs, H-1000–10). Primary antibodies used: Mouse anti-repo (DSHB: 8D12) 1:50, Rat anti-elav (DSHB: 7E8A10) 1:100, Goat anti-GFP (abcam: ab6662) 1:500. Secondary antibodies used: Anti-mouse-647 (Jackson ImmunoResearch: 715-605-151) 1:250, Anti-rat-Cy3 (Jackson ImmunoResearch: 712-165-153) 1:500. The samples were scanned using a laser confocal microscope (Zeiss LSM 880) with a 20X objective, and images were processed using ZEN (Zeiss) and Imaris (Oxford Instruments) software. Co-localization between mCherry and Elav or mCherry and Repo was performed with default thresholds in Imaris.
GeneOntology (GO) analysis—GO analysis was performed based on the PANTHER (Protein Analysis Through Evolutionary Relationships) system (http://www.pantherdb.org; date last accessed October 31, 2020 (Ashburner et al., 2000). Statistical analysis was performed by using the default PANTHER Overrepresentation Test (Released 20200728), Annotation Version and Release Date: GO Ontology database https://doi.org/10.1016/10.5281/zenodo.4033054 Released 2020-09-10 which used the Fisher’s Exact test with a false discovery rate p < 0.05.

Exome sequencing and identification of GLRA2 variants—Subjects 1, 2, 4, 5 and 8 had clinical exome sequencing at GeneDx (Gaithersburg, MD, United States), at the Praxis für Humangenetik Tubingen (Tubingen, Germany), at Baylor Genetics (Houston, TX, United States), at Pitié-Salpêtrière Hospital Genetics lab (Paris, France) and at Integragen (Evry, France), respectively. Subject 3 WES was performed at the Meyer Children’s Hospital, University of Florence, in the context of the DESIRE program and as previously described (Vetro et al., 2020). Briefly, the SureSelectXT Clinical Research Exome kit (Agilent Technologies, Santa Clara, CA) was used for library preparation and target enrichment, and paired-end sequencing was performed using Illumina sequencer (NextSeq550, Illumina, San Diego, CA, USA) to obtain an average coverage of above 80x, with 97.6% of target bases covered at least 10x. Reads were aligned to the GRCh37/hg19 human genome reference assembly by the BWA software package, and the GATK suite was used for base quality score recalibration, realignment of insertion/deletions (InDels), and variant calling (DePristo et al., 2011; McKenna et al., 2010). Variant annotation and filtering pipeline included available software (VarSeq, Golden Helix, Inc v1.4.6), focusing on nonsynonymous/splice site variants with minor allele frequency (MAF) lower than 0.01 in the GnomAD database (Karczewski et al., 2020) (http://gnomad.broadinstitute.org/), an internal healthy control database and pre-computed genomic variants score from dbNSFP (Liu et al., 2011). Subject 6 WES was performed at the Institute of Human Genetics, Technical University of Munich, Germany as described previously (Brunet et al., 2021). DNA was extracted from blood samples by the Gentra Puregene Blood Kit (Qiagen, Hilden, Germany). WES was performed using the Sure Select Human All Exon 60 Mb V6 Kit (Agilent, Santa Clara, CA, USA) for exomic enrichent according to the manufacturer’s protocol. Sequencing was performed on a NovaSeq 6000 sequencer (Illumina, San Diego, CA, USA) to an average coverage of >94X. Reads were aligned to the UCSC human reference assembly (hg19) with the BWA algorithm v.0.5.9. For detection of single-nucleotide variants (SNVs) as well as small insertions and deletions SAMtools v.0.1.19 was applied. Copy number variations (CNVs) were called with the software ExomeDepth. In-house custom Perl scripts were used for variant annotation. Variant analysis was performed using I) a recessive filter for homozygous or compound-heterozygous variants with a minor allele frequency (MAF) of <1% in our in-house database of 22,000 exome datasets, II) a filter for X-chromosomal variants with a MAF<0.1%, III) a filter for de novo variants (MAF <0.01%) and IV) a phenotype-based filter using search terms of characteristic phenotypic traits and a MAF <0.1%. CNVs were assessed using a MAF <0.01%. Current criteria for variant classification according to the American College of Human Genetics (ACMG) were used for variant interpretation (Richards et al., 2015). Subject 7 had exome sequencing at Lyon University Hospital (Lyon, France). The SeqCap EZ Medexome kit (Roche, Pleasanton, CA, USA)
was used for library preparation and target enrichment before paired-end sequencing using an Illumina instrument (NextSeq500, Illulina, San Diego, CA, USA). A mean depth of coverage of 133x was obtained with 99.0% of target bases covered at least 10x. Reads were aligned to the GRCh37/hg19 human genome reference assembly by the BWA software package, and the GATK suite was used for base quality score recalibration, realignment of insertion/deletions (InDels), and variant calling (DePristo et al., 2011; McKenna et al., 2010). Variant annotation was performed with SnpEFF and filtering pipeline focused on non-synonymous/splice site variants with minor allele frequency (MAF) lower than 0.01 in the GnomAD database (Karczewski et al., 2020) (http://gnomad.broadinstitute.org/). Subject 9 WES was performed at the Erasmus MC as previously described (Perenthaler et al., 2020). In brief, exome-coding DNA was captured with the Agilent SureSelect Clinical Research Exome (CRE) kit (v2). Sequencing was performed on an Illumina HiSeq 4000 platform with 150-bp paired-end reads. Reads were aligned to hg19 using BWA (BWA-MEM v0.7.13) and variants were called using the GATK Haplotype Caller (McKenna et al., 2010) v3.7 (https://www.broadinstitute.org/gatk/). Detected variants were annotated, filtered and prioritized using the Bench lab NGS v5.0.2 platform (Agilent technologies).

Subject 10 WES and data processing were performed by the Genomics Platform at the Broad Institute of MIT and Harvard with an Illumina Nextera or Twist exome capture (~38 Mb target), and sequenced (150 bp paired reads) to cover >80% of targets at 20x and a mean target coverage of >100x. WES data was processed through a pipeline based on Picard and mapping done using the BWA aligner to the human genome build 38. Variants were called using Genome Analysis Toolkit (GATK) Haplotype Caller package version 3.5 (McKenna et al., 2010) (https://www.broadinstitute.org/gatk/). WES for subjects 11 and 12 was performed in collaboration with the Autism Sequencing Consortium (ASC) at the Broad Institute on Illumina HiSeq sequencers using the Illumina Nextera exome capture kit. Exome sequencing data was processed through a pipeline based on Picard and mapping done using the BWA aligner to the human genome build 37 (hg19). Variants were called using Genome Analysis Toolkit (GATK) Haplotype Caller package version 3.4 (McKenna et al., 2010) (https://www.broadinstitute.org/gatk/). Variant call accuracy was estimated using the GATK Variant Quality Score Recalibration (VQSR) approach. High-quality variants with an effect on the coding sequence or affecting splice site regions were filtered against public databases (dbSNP150 and gnomAD V2.0) to retain (i) private and clinically associated variants; and (ii) annotated variants with an unknown frequency or having minor allele frequency <0.1%, and occurring with a frequency <2% in an in-house database including frequency data from > 1,500 population-matched WES. The functional impact of variants was analyzed by CADD V1.3, Mendelian Clinically Applicable Pathogenicity V1.0 (Jagadeesh et al., 2016; Kircher et al., 2014), and using InterVar V0.1.6 to obtain clinical interpretation according to American College of Medical Genetics and Genomics/Association for Molecular Pathology 2015 guidelines (Li and Wang, 2017).

Trio exome sequencing of subject 13 and his parents was performed at the Institute of Human Genetics, University of Leipzig Medical Center, in the context of the research project “Genetics of rare disorders”. Testing in a research setting was approved by the ethics committee of the University of Leipzig (224/16-ek and 402/16-ek). Library preparation was done using the Nextera DNA Flex Pre-Enrichment Library Prep with Illumina Nextera DNA...
UD Indexes by Illumina. Target enrichment was achieved by using the Human Core Exome hybridization probes from Twist Bioscience. Paired-end Next-Generation-Sequencing was then performed on a NovaSeq 6000 Instrument using an S1 Reagent Kit (300 cycles) by Illumina. Analysis of the raw data was performed using the software Varfeed (Limbus, Rostock). Variants were annotated and prioritized using the software Varvis (Limbus, Rostock). Rare variants (minor allele frequency below 1% in the general population) were prioritized based on inheritance mode, impact on protein, clinical relevance in variant databases and in silico prediction. GeneMatcher (Sobreira et al., 2015) (https://genematcher.org/) assisted in the recruitment of Subjects 2, 3 and 5–13.

**SDS-PAGE/Western blot**—Five heads of nSyb-GAL4 UAS-GLRA2 reference and variant flies aged for 5 days post eclosion were lysed in 30μL NETN buffer (50mM Tris pH 7.5, 150mM NaCl, 0.5% NP-40, 1 mM EDTA) with an electric douncer for 10 seconds for three times on ice. 30μL of 2x Laemmli Sample Buffer (Bio-Rad) with 10% 2-mercaptoethanol was added to the lysis and incubated on ice for 10 min. Samples were boiled at 95°C and spun at 14,000 RPM for 5 minutes at 4°C. The soluble fraction was loaded onto a standard SDS-PAGE gel. PVDF (polyvinylidene difluoride) membrane activated for 1 minute with 100% methanol. After running and wet transfer, the membrane was blocked in 5% skim milk for 1 hour. The membrane was incubated (overnight, shaking, at 4°C) with mouse anti-HA (HA.11, 1:1,000, 901501, BioLegend) and mouse anti-Actin (C4) (1:50,000, MAB1501, EMD Millipore) primary antibodies in 3% BSA (bovine serum albumin), followed by 10 minute washes (3 times) with 1% Triton-X in Tris-buffered saline (TBST). We incubated this with goat anti-mouse HRP-conjugated (1:15000, 115-035-146, Jackson ImmunoResearch) secondary antibody in skim milk. The membrane was washed three times with 1% TBST and detected with Western Lightning™ Chemiluminescence Reagent Plus (perkinelmerNEL104001EA) ECL solution using the Bio-Rad ChemiDoc MP imaging system.

**Structural biological analysis of GLRA2 patient variants**—Protein residues that corresponds to GLRA2 patient variants were mapped onto the crystal protein structure of GLRA1 protein in Protein Data Bank (PDB, ID: 4X5T) (Burley et al., 2019) using the PyMOL (https://pymol.org/) (Yuan et al., 2016) because GLRA1 and GLRA2 are highly homologous proteins (85% similarity, 78% identity and 3% gaps) based on DIOPT (Hu et al., 2011).

**Image generation**—Cartoon images in Figure 1H were generated with BioRender.com.

**QUANTIFICATION AND STATISTICAL ANALYSIS**

All statistical analysis was performed with GraphPad Prism version 8. Significance was defined as *p<0.05, **p<0.01, ***p<0.001, ****p<0.0001.

**Gene and variant level statistics**—Gene level constraints for Figures 1B–1D (pLI, LOEUF, missense O/E) for all genes (Global, n=19704), the SSC subset (n=1493), and the study subset (n=143) were based on metric availability from gnomAD and analyzed by ANOVA followed by Dunn’s multiple comparison test. The same test was used in Figure
S9B–S9Y for gene and variant analysis. The contingency graph for variant consequences (Figure S9A) grouped by variants corresponding to lethal (essential genes) or viable (non-essential genes) TG4 mutants was analyzed by Chi square, df (20, 3, p<0.0002).

**Rescue-based and overexpression-based survival**—For humanized rescue of TG4 lethality (Figures 2B–2D), 3 independent crosses were set per genotype and a minimum of n>50 flies were quantified for each cross. Statistical analyses were performed by ANOVA followed by Sidak’s multiple comparisons test. For lethality caused by overexpression with Tub-GAL4 (Figures 4B, 5B, and S12B), a minimum of 3 independent crosses were set with two independent UAS-transgenic lines. 50–100 flies (a minimum of 10 if overexpression caused survival defects) were scored. Statistical analyses were performed by unpaired t test.

**Lifespan analysis**—Lifespan analysis of humanized TG4 lines at 25°C (Figures 2E and 2F) were analyzed by Log rank (Mantel-Cox) test with a minimum of 11–49 flies for each genotype from three independent crosses.

**Behavior**—Fore courtship analysis (Figures 3C–3G and S4A–S4T), at least 10 animals were tested per genotype. Analysis of data was performed using Excel (Microsoft) and Prism (GraphPad). A ROUT (Q=1%) test was performed in Prism to identify outliers. Data was analyzed with Kruskal-Wallis ANOVA followed by Dunn’s multiple comparison test.

**Assessment of melanized nodules**—Melanized nodules formed on the notum of flies expressing GLRA2$^{T296M}$ driven by pnr-GAL4 (Figure 5D) were analyzed by ANOVA followed by Tukey’s multiple comparison test. 3 independent crosses were set and minimum of 50 flies examined for presence of melanized nodules per genotype.

**Electroretinograms**—Electrophysiological field potential recordings (ERG) of nSyb-GAL4 or Rh1-GAL4 flies expressing UAS-GLRA2 constructs (Figures 5E–5H and S12C–S12F) were analyzed by ANOVA followed by Bonferroni’s multiple comparison test. 4–10 flies were examined for each genotype. Recording was repeated at least 3 times per fly.

**Supplementary Material**

Refer to Web version on PubMed Central for supplementary material.

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Highlights

- We generate and characterize >300 (TG4 and cDNA) *Drosophila* mutants and transgenics
- Humanization and overexpression strategies to functionally assess ASD variants *in vivo*
- ASD variant data in flies help identify *GLRA2*-related neurodevelopmental disorders
- Basic and clinical collaboration facilitates variant testing and disease gene discovery
Figure 1. Gene and variant prioritization, resource generation, and screening outline

(A) Criteria to prioritize ASD candidate genes and variants for this study. (B–D) Gene level constraints from control individuals (gnomAD) for (B) probability of loss of function intolerance (pLI), (C) loss of function observed or expected (o/e) upper bound fraction (LOEUF), and (D) missense o/e. **p < 0.01, ***p < 0.001, and ****p < 0.0001.

(E) Schematic depicting generation and effect of TG4 lines on gene function.

(F) Schematic illustrating generation of UAS-human cDNA constructs.

(G) Total number of Drosophila reagents generated for this study.

(H) Screening paradigms using both humanization and overexpression strategies to assess SSC-DNM function.
Figure 2. Assessment of SSC-DNM function through humanization of essential fly genes

(A) Rescue of lethality to adult stage by TG4-driven UAS-reference human cDNA and subsequent comparison of reference and variant cDNA.

(B–D) Observed/expected Mendelian ratios for rescue of humanized TG4 mutants across different temperatures. Three independent crosses were set per genotype, and n > 50 flies were quantified for each cross. Statistical analyses were performed by ANOVA followed by Sidak’s multiple comparisons test. Error bars are +SEM (standard error of mean).

(E and F) Lifespan analysis of humanized TG4 lines at 25°C. Survival comparisons obtained by log rank (Mantel-Cox) test with a minimum of 11–49 flies for each genotype from three independent crosses.
(G) Single focal confocal images showing expression pattern of UAS-nls.mCherry driven by TG4 (red) in the anterior of the central adult brain. Bottom two rows depict the 5× zoom from dotted white box of co-localization between TG4 reporter and Elav (neurons) or Repo (glia) in cyan. Co-localization is depicted in white. Scale bars represent 50 μm. Dotted magenta lines outline of the brain. *p < 0.05, **p < 0.01, ***p < 0.001, and ****p < 0.0001.
Figure 3. Assessment of SSC-DNM function through humanization of viable TG4 lines and behavioral analysis

(A) Analysis pipeline used to evaluate *Drosophila* behavior.

(B) SSC-DNMs in which variants display significant differences in time spent performing a specific behavior (courtship, copulation, movement, or grooming) when compared with reference humanized flies.

(C–G) The number of frames male flies spent performing courtship (single-wing extensions), copulating, moving within the chamber, or grooming during a 30-min test period. The red line represents the average number of frames a *Canton-S* (control) male spends performing the same task. n = 10–40 flies were used per genotype. Statistical analysis was performed by non-parametric Kruskal-Wallis one-way ANOVA and the Dunn’s multiple comparison test.
(H) Single focal confocal images showing expression pattern of UAS-nls.mCherry driven by TG4 (red) in the anterior of the central adult brain. Bottom two rows depict the 5× zoom from dotted white box of co-localization between TG4 reporter and Elav (neurons) or Repo (glia) in cyan. Co-localization is depicted in white. Scale bars represent 50 μm. *p < 0.05, ***p < 0.001, ****p < 0.0001, and ns, not significant. Error bars are ± SEM.
Figure 4. Variant assessment by overexpression of reference and SSC-DNMs

(A) Phenotypes observed upon overexpressing the reference and variant cDNAs using a ubiquitous driver (tub-GAL4) at 25°C, an eye-specific driver (GMR-GAL4) at 29°C, or a wing-specific driver (nub-GAL4) at 25°C. Black denotes there was no phenotype (NP), purple there was a comparable phenotype (CP), or red there was a functional difference (FD).

(B) Quantification of viability upon overexpression of reference or variant human cDNAs using a ubiquitous driver for genes where the variants showed a functional difference. Minimum of three independent crosses were set with two independent UAS-transgenic lines. We scored 50–100 flies (a minimum of 10 if overexpression caused survival defects). Statistical analyses were performed by unpaired t test.
(C and D) Representative optical sections of eyes and wings for variants with a functional difference using eye-specific (GMR-GAL4) and wing-specific (nub-GAL4) drivers, respectively, at 25°C. *p < 0.05, **p < 0.01, ***p < 0.001, and ****p < 0.0001. Error bars are +SEM.
Figure 5. GLRA2<sup>T296M</sup> found in female probands acts as a GoF allele while GLRA2<sup>R252C</sup> and GLRA2<sup>N136S</sup> found in male probands behave as LoF alleles

(A) Schematic diagram of domain structure of GLRA2 and the relative positions of subject variants functionally assessed in Drosophila.

(B) Mendelian ratios upon overexpression of the GLRA2 reference or variant human cDNAs using a ubiquitous driver (tub-GAL4). A minimum of three independent crosses were set.

(C and D) Representative images and quantification of melanized nodules formed on the notum of flies expressing GLRA2<sup>T296M</sup> driven by a dorsocentral thorax-specific (pnr-GAL4) driver at 25°C.

(E–H) Representative traces of ERG and quantification of “OFF”-transient amplitude (blue bracket) in animals expressing GLRA2 pan-neuronally (both pre-synaptic photoreceptors and post-synaptic laminar neurons; nSyb-GAL4) or only in the pre-synaptic photoreceptors.
(Rh1-GAL4). ERG was analyzed by ANOVA followed by Bonferroni’s multiple comparison test. Four to ten flies were examined for each genotype. Recording was repeated at least three times per fly. **p < 0.01, ***p < 0.001, and ****p < 0.0001. Error bars are +SEM.
# Table 1.

Identification of 30 SSC-DNMs with functional consequences

| H. sap gene | pLI (LOEUF) | Missense O/E | OMIM disease | SSC variant | CADD | D. mel gene | TG4 lethality | Functional assay | SSC-CV consequence |
|-------------|-------------|--------------|---------------|-------------|------|-------------|---------------|------------------|-------------------|
| **ABL2**    | 0 (0.58)    | 0.81         | -             | p.A1099T    | 28.2 | *Abl*       | yes           | RB               | LoF               |
| **ACE**     | 0 (1.08)    | 1.07         | 267430 (AR)   | PY818C      | 7.5  | *Ance*      | no            | OE               | GoF               |
| **ALDH1L1** | 0 (0.78)    | 0.93         | -             | EN900H      | 9.8  | *CG8665*    | no            | RB               | GoF?              |
| **ATP2B2**  | 1 (0.15)    | 0.54         | 601386 (AR)   | PT818M      | 33.0 | *PMCA*      | yes           | OE               | LoF               |
| **BAIAP2L1**| 0 (0.65)    | 0.90         | -             | PA481V      | 17.8 | *IRSp53*    | no            | OE               | GoF               |
| **CAT**     | 0 (1.05)    | 1.01         | 614097 (AR)   | PG204E      | 28.1 | *Cat*       | yes           | RB               | LoF               |
| **CHST2**   | 0.02 (0.81) | 0.66         | -             | p.R52P      | 12.8 | *CG31637*   | yes           | RB               | LoF               |
| **EPHA1**   | 0 (1.04)    | 0.97         | -             | p.V567I     | 1.3  | *Eph*       | no            | OE               | LoF               |
| **EPIBI1**  | 1 (0.26)    | 0.73         | -             | PV916M      | 34.0 | *Epih*      | no            | OE               | complex           |
| **GLRA2**   | 0.97 (0.30) | 0.43         | -             | PN136S      | 25.1 | *GluCka*    | no            | RB, OE           | LoF               |
| **GPC5**    | 0 (1.08)    | 1.09         | -             | PM133T      | 24.3 | *daily*     | no            | OE               | GoF               |
| **GRK4**    | 0 (1.09)    | 1.09         | -             | PP358A      | 26.0 | *Gprk2*     | yes           | OE               | LoF               |
| **HTR1D**   | 0 (1.30)    | 0.98         | -             | p.T99N      | 19.4 | *5-HT1B*    | no            | OE               | GoF               |
| **IRF2BPL** | 0.84 (0.41) | 0.90         | 618088 (AD)   | p.F30L, p.N701fs | 24.8 | *Pits*     | yes           | OE               | LoF               |
| **ITGA8**   | 0 (0.66)    | 1.03         | 191830 (AR)   | PR748C      | 35.0 | *if*        | yes           | OE               | LoF               |
| **KCND3**   | 0.99 (0.28) | 0.48         | 607346 (AD)   | p.R86P      | 32.0 | *Shal*      | no            | RB, OE           | LoF               |
| **KDM2A**   | 1 (0.04)    | 0.43         | -             | PR449K      | 5.7  | *Kdm2*      | no            | RB               | ?                 |
| **MAP4K1**  | 0.99 (0.29) | 0.59         | -             | PM725T      | 21.3 | *hppy*      | no            | OE               | complex           |
| **MINK1**   | 1 (0.13)    | 0.60         | -             | PC269R      | 26.8 | *msn*       | yes           | OE               | LoF               |
| **MYH9**    | 1 (0.09)    | 0.71         | 603622 (AD)   | p.R1571Q    | 35.0 | *Mbc*       | no            | OE               | GoF               |
| **NPFFR2**  | 0 (1.13)    | 1.20         | -             | PM163I      | 13.3 | *SIFaR*     | yes           | OE               | LoF               |
| **PC**      | 0.01 (0.43) | 0.69         | 266150 (AR)   | PP1042R     | 24.6 | *PCB*       | no            | OE               | GoF               |
| **PDK2**    | 0 (0.92)    | 0.63         | -             | PR120Q      | 25.3 | *Pdk*       | yes           | OE               | LoF               |
| **SLC23A1** | 0.02 (0.54) | 0.71         | -             | PL465M      | 17.9 | *CG6293*    | no            | OE               | GoF               |
| **TRIP12**  | 1 (0.06)    | 0.60         | 617752 (AD)   | PR1643Q     | 36.0 | *ctrip*     | yes           | RB               | LoF               |
| H. sapiens gene | pLI (LOEUF) | Missense O/E | OMIM disease | SSC variant | CADD | D. melanogaster gene | TG4 lethality | Functional assay | SSC-CV consequence |
|----------------|-------------|-------------|--------------|-------------|------|---------------------|--------------|-----------------|------------------|
| TRPM6 | 0 (0.45) | 0.86 | 602014 (AR) | p.T2011P | 12.2 | Tripm | yes | RB | LoF |
|  |  |  | | p.A641E | 17.8 |  |  | RB |  |
| TSC2 | 1 (0.07) | 1.03 | 613254 (AD) | p.R1557W | 16.0 | gG | yes | OE | LoF |
| USP30 | 0 (0.66) | 0.76 | - | PP200S | 14.7 | Usp30 | no | RB | ? |

List of all human genes and corresponding SSC variants determined to have a functional difference across all assays in this study. GoF, gain of function; LoF, loss of function; OE, overexpression; RB, rescue-based.

*a Nervous system disorder

*b Known lethal mutants
Table 2.

Salient features of subjects with GLRA2 variants

| Subject | GLRA2 variant (hg19, NM_001118886.1) | Inheritance | CADD score | Gender | Age at most recent evaluation (years) | Developmental delay/intellectual disability | Hypotonia/incoordination | Autism spectrum disorder | Inattention/hyperactivity | Sleep disturbance | Microcephaly | Ocular features | Epilepsy | EEG findings | EEG findings |
|---------|--------------------------------|-------------|------------|--------|--------------------------------------|-----------------------------------------------|------------------------|-------------------------|-----------------------|----------------|-------------|----------------|----------|--------------|--------------|
| 1       | c.887C>T, p.Thr296Met           | de novo     | 27         | female | 6.7                                  | yes                                           | no                     | no                      | yes                   | no             | yes         | myopia, astigmatism, and nystagmus (improved with age) | no         | slow background suggestive of mild encephalopathy |
| 2       | c.887C>T, p.Thr296Met           | de novo     | 27         | female | 6.5                                  | yes                                           | no                     | no                      | yes                   | no             | yes         | alternating exotropia, borderline oculomotor apraxia, ptosis | yes        | normal |
| 3       | c.887C>T, p.Thr296Met           | de novo     | 27         | female | 5.5                                  | yes                                           | no                     | no                      | no                    | no             | yes         | nystagmus (improved with age) | no         | normal |
| 4       | c.887C>T, p.Thr296Met           | de novo     | 27         | female | 0.5                                  | yes                                           | no                     | no                      | no                    | no             | no          | none                   | yes        | normal |
| 5       | c.887C>T, p.Thr296Met           | de novo     | 27         | female | 5.4                                  | yes                                           | no                     | no                      | no                    | no             | no          | oculomotor apraxia, ptosis | no         | normal |
| 6       | c.887C>T, p.Thr296Met           | de novo     | 27         | female | 0.8                                  | yes                                           | no                     | no                      | no                    | no             | no          | alternating exotropia, borderline oculomotor apraxia, ptosis | no         | normal |
| 7       | c.140T>C, p.Phe47Ser            | de novo     | 27         | female | 6.7                                  | yes                                           | no                     | no                      | no                    | no             | no          | nystagmus (starting 6 weeks after birth) | no         | normal |
| 8       | c.777C>G, p.Ile259Met           | de novo     | 27.8       | female | 5                                    | yes                                           | yes                    | yes, incoordination | no                    | no             | no          | normal                   | yes        | normal |
| 9       | c.862G>A, p.Arg252Cys           | de novo     | 31         | male   | 7                                    | yes, with regression                           | yes                    | yes, ataxia             | no                    | yes            | no          | myopia, astigmatism | no         | normal |
| 10      | c.754C>T, p.Arg252Cys           | de novo     | 27.2       | male   | 15                                   | yes                                           | yes                    | yes                     | yes                   | no             | yes         | nystagmus (improved with age) | no         | normal |
| 11      | c.618G>A, p.Ala288Thr           | de novo     | 20.9       | male   | 5                                    | yes                                           | yes, with regression                           | yes                    | yes                     | yes                   | no             | yes         | none                   | no         | normal |
| 12      | c.1186C>T, p.Pro396Leu          | de novo     | 22.3       | male   | 5                                    | yes                                           | yes                    | yes                     | yes                   | no             | yes         | nystagmus (starting 6 weeks after birth) | no         | normal |
| 13      | c.1199C>T, p.Pro400Leu          | de novo     | 33         | male   | 5.9                                  | yes                                           | yes                    | yes                     | yes                   | no             | yes         | nystagmus (improved with age) | no         | normal |
| 14      | c.1334G>A, p.Arg445Gln          | de novo     | 34         | male   | 5                                    | yes                                           | yes                    | yes                     | yes                   | no             | yes         | nystagmus (improved with age) | no         | normal |

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| Subject | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 | 9 | 10 | 11 | 12 | 13 |
|---------|---|---|---|---|---|---|---|---|---|----|----|----|----|
| Brain MRI findings | normal | delayed myelination, a small arachnoid cyst | mild cortical atrophy, thinning of corpus callosum | normal | normal | normal | cortical and white-matter atrophy, including vermian atrophy | increased signal intensity in FLAIR of the subcortical white matter of the frontal region | not performed | minimally increased T2 signal intensity on the occipital lobes | normal | increased signal intensity in FLAIR of the cortical matter of the parietal region | normal |

CADD, combined annotation-dependent depletion; EEG, electroencephalography; MRI, magnetic resonance imaging.
# KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Antibodies**      |        |            |
| anti-elav           | DSHB Cat# 7E8A10 | RRID:AB_528218 |
| anti-Repo           | DSHB Cat# 8D12 | RRID:AB_528448 |
| anti-mouse-647      | Jackson ImmunoResearch Labs Cat# 715-605-151 | RRID:AB_2340863 |
| anti-rat-Cy3        | Jackson ImmunoResearch Labs Cat# 712-165-153 | RRID:AB_2340667 |
| anti-GFP            | Abcam Cat# ab6662 | RRID:AB_305635 |
| anti-HA             | BioLegend Cat# 902301 | RRID:AB_2565018 |
| anti-actin          | Millipore Cat# MAB1501 | RRID:AB_2223041 |

| **Experimental models: Organisms/strains** | | |
|-------------------------------------------|--------|------------|
| UAS-2xEGFP, hs-Cre, vas-dpC31             | Diao et al., 2015 | N/A |
| Tub-GAL4                                  | BDSC   | RRID:BDSC_5138  |
| nubbin-GAL4                               | BDSC   | RRID:BDSC_51635 |
| GMR-GAL4                                  | BDSC   | RRID:BDSC_1104  |
| UAS-nls.GFP                               | BDSC   | RRID:BDSC_4775  |
| UAS-nls.mCherry                           | BDSC   | RRID:BDSC_38424 |
| nSyb-GAL4                                 | BDSC   | RRID:BDSC_51635 |
| Rh1-GAL4                                  | BDSC   | RRID:BDSC_6691  |
| pmr-GAL4                                  | BDSC   | RRID:BDSC_3039  |
| CG4562 Tg4                                 | BDSC   | RRID:BDSC_76740 |
| Abl Tg4                                    | BDSC   | RRID:BDSC_67429 |
| Adk1 Tg4                                   | Lee et al., 2018 | N/A |
| CG7470 Tg4                                 | BDSC   | RRID:BDSC_76749 |
| CG8665 Tg4                                 | BDSC   | RRID:BDSC_66811 |
| RhoGAP19 D Tg4                             | BDSC   | RRID:BDSC_76687 |
| osa Tg4                                    | Lee et al., 2018 | N/A |
| PMCA Tg4                                   | BDSC   | RRID:BDSC_76741 |
| Cat Tg4                                    | BDSC   | RRID:BDSC_76660 |
| Cep135 Tg4                                 | BDSC   | RRID:BDSC_66853 |
| CG31637 Tg4                                | BDSC   | RRID:BDSC_76647 |
| CIC-a Tg4                                  | BDSC   | RRID:BDSC_66801 |
| CLIP-190 Tg4                               | BDSC   | RRID:BDSC_66834 |
| Dh44-R2 Tg4                                | BDSC   | RRID:BDSC_66865 |
| h Tg4                                      | BDSC   | RRID:BDSC_76724 |
| arm Tg4                                    | BDSC   | RRID:BDSC_66903 |
| mbc Tg4                                    | BDSC   | RRID:BDSC_66840 |
| spg Tg4                                    | BDSC   | RRID:BDSC_76205 |
| CG17684 Tg4                                | Lee et al., 2018 | N/A |
| shot Tg4                                   | BDSC   | RRID:BDSC_76760 |

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| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| fne        TG4       | BDSC   | RRID:BDSC_77796 |
| dom        TG4       | BDSC   | RRID:BDSC_76192 |
| Eph        TG4       | BDSC   | RRID:BDSC_66800 |
| fly        TG4       | BDSC   | RRID:BDSC_76736 |
| Gclc        TG4       | BDSC   | RRID:BDSC_76654 |
| daddy       TG4       | BDSC   | RRID:BDSC_66830 |
| GluRIB      TG4       | BDSC   | RRID:BDSC_76135 |
| Nindar2     TG4       | BDSC   | RRID:BDSC_76705 |
| Gpkl2       TG4       | BDSC   | RRID:BDSC_66828 |
| Leep        TG4       | BDSC   | RRID:BDSC_77798 |
| Pits        TG4       | BDSC   | RRID:BDSC_77731 |
| if          TG4       | BDSC   | RRID:BDSC_66867 |
| Lpt         TG4       | BDSC   | RRID:BDSC_76714 |
| wh          TG4       | BDSC   | RRID:BDSC_76189 |
| Lrchn       TG4       | BDSC   | RRID:BDSC_77756 |
| LRPFj       TG4       | BDSC   | RRID:BDSC_76640 |
| Rah3-GEF     TG4       | BDSC   | RRID:BDSC_76623 |
| hppy        TG4       | BDSC   | RRID:BDSC_67447 |
| mbl         TG4       | BDSC   | RRID:BDSC_66779 |
| msn          TG4       | BDSC   | RRID:BDSC_76204 |
| Mhc          TG4       | BDSC   | RRID:BDSC_76653 |
| cl            TG4       | BDSC   | RRID:BDSC_76720 |
| Nlg3        TG4       | BDSC   | RRID:BDSC_76134 |
| Cad99C       TG4       | BDSC   | RRID:BDSC_67483 |
| a            TG4       | BDSC   | RRID:BDSC_76725 |
| plj          TG4       | BDSC   | RRID:BDSC_77693 |
| Peczo        TG4       | BDSC   | RRID:BDSC_66558 |
| ptx1         TG4       | BDSC   | RRID:BDSC_67497 |
| l(1)G0289    TG4       | BDSC   | RRID:BDSC_67467 |
| CG31211      TG4       | BDSC   | RRID:BDSC_76718 |
| Spn          TG4       | This study | N/A |
| CG6767       TG4       | BDSC   | RRID:BDSC_77797 |
| otk          TG4       | BDSC   | RRID:BDSC_76759 |
| Lar          TG4       | BDSC   | RRID:BDSC_67451 |
| Pxn          TG4       | BDSC   | RRID:BDSC_66850 |
| CG5521       TG4       | BDSC   | RRID:BDSC_76180 |
| emp          TG4       | BDSC   | RRID:BDSC_66904 |
| sdk          TG4       | BDSC   | RRID:BDSC_76628 |
| retm         TG4       | BDSC   | RRID:BDSC_66816 |
| SemaSc       TG4       | BDSC   | RRID:BDSC_77809 |

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| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| CG6293 TG4          | BDSC   | RRID:BDSC_76761 |
| CG18304 TG4         | BDSC   | RRID:BDSC_76128 |
| rol5 TG4            | BDSC   | RRID:BDSC_76150 |
| CG7744 TG4          | BDSC   | RRID:BDSC_76662 |
| Tipm1 TG4           | BDSC   | RRID:BDSC_77748 |
| Nipped-A TG4        | BDSC   | RRID:BDSC_76723 |
| gll TG4             | BDSC   | RRID:BDSC_67515 |
| Tisp TG4            | BDSC   | RRID:BDSC_66798 |
| unc890 TG4          | BDSC   | RRID:BDSC_76686 |
| Usp35 TG4           | BDSC   | RRID:BDSC_76704 |
| bchs TG4            | BDSC   | RRID:BDSC_76762 |
| CG6225 TG4          | BDSC   | RRID:BDSC_76769 |
| Yip1d1 TG4          | BDSC   | RRID:BDSC_67492 |
| Anco TG4            | BDSC   | RRID:BDSC_76676 |
| Ace TG4             | BDSC   | RRID:BDSC_76688 |
| Aldh-III TG4        | BDSC   | RRID:BDSC_77692 |
| CG33298 TG4         | BDSC   | RRID:BDSC_76700 |
| IRSp53 TG4          | BDSC   | RRID:BDSC_67637 |
| Best1 TG4           | BDSC   | RRID:BDSC_76671 |
| tok TG4             | BDSC   | RRID:BDSC_76679 |
| CASK TG4            | BDSC   | RRID:BDSC_76631 |
| Ddr TG4             | This study | N/A |
| cv-c TG4            | This study | N/A |
| CRMP TG4            | This study | N/A |
| CG11594 TG4         | This study | N/A |
| GluClalpha TG4      | BDSC   | RRID:BDSC_77841 |
| Galphao TG4         | This study | N/A |
| 5-HT1B TG4          | BDSC   | RRID:BDSC_76668 |
| ltp-r83A TG4        | This study | N/A |
| Shal TG4            | This study | N/A |
| Kdm3a TG4           | This study | N/A |
| Pve TG4             | BDSC   | RRID:BDSC_76657 |
| beta-Man TG4        | This study | N/A |
| dxp TG4             | This study | N/A |
| Sme TG4             | BDSC   | RRID:BDSC_76743 |
| Ndg TG4             | BDSC   | RRID:BDSC_76768 |
| Nos TG4             | BDSC   | RRID:BDSC_76766 |
| SIFaR TG4           | BDSC   | RRID:BDSC_76670 |
| NetB TG4            | BDSC   | RRID:BDSC_76730 |
| PH4alphaEFB TG4     | BDSC   | RRID:BDSC_76678 |

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| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| PCB Tg4             | BDSC   | RRID:BDSC_66832 |
| Pdk Tg4             | BDSC   | RRID:BDSC_77785 |
| PKD Tg4             | BDSC   | RRID:BDSC_76706 |
| Rim Tg4             | This study | N/A |
| CG9098 Tg4          | BDSC   | RRID:BDSC_76763 |
| ctrip Tg4           | BDSC   | RRID:BDSC_76764 |
| CG1815 Tg4          | This study | N/A |
| UAS-ABCC4           | BDSC   | RRID:BDSC_78481 |
| UAS-ABCC4.M276V     | BDSC   | RRID:BDSC_92726 |
| UAS-ABCC5           | BDSC   | RRID:BDSC_78508 |
| UAS-ABCC5.R697W     | BDSC   | RRID:BDSC_92728 |
| UAS-ABCC5.T1046M    | BDSC   | RRID:BDSC_92727 |
| UAS-ABL2.A1084T     | BDSC   | RRID:BDSC_92729 |
| UAS-ABL2            | BDSC   | RRID:BDSC_78453 |
| UAS-ACE.Y818C       | BDSC   | RRID:BDSC_92730 |
| UAS-ACE             | This study | N/A |
| UAS-ACHE            | BDSC   | RRID:BDSC_78466 |
| UAS-ACHE.G151R      | BDSC   | RRID:BDSC_92732 |
| UAS-ACHE.P548L      | BDSC   | RRID:BDSC_92731 |
| UAS-AK1             | BDSC   | RRID:BDSC_78462 |
| UAS-AK1.558L        | BDSC   | RRID:BDSC_92733 |
| UAS-ALDH18A1.D703H.HA | BDSC   | RRID:BDSC_92734 |
| UAS-ALDH18A1.HA     | BDSC   | RRID:BDSC_78488 |
| UAS-ALDH111.N900H   | BDSC   | RRID:BDSC_92735 |
| UAS-ALDH111         | This study | N/A |
| UAS-ALDH3A1.F402L.HA | BDSC   | RRID:BDSC_92926 |
| UAS-ALDH3A1.HA      | BDSC   | RRID:BDSC_78497 |
| UAS-ATP10A          | This study | N/A |
| UAS-ATP2B2.T818M    | BDSC   | RRID:BDSC_92952 |
| UAS-ATP2B2          | This study | N/A |
| UAS-ATP2B4          | BDSC   | RRID:BDSC_78455 |
| UAS-BAIAP2L1.A481V  | BDSC   | RRID:BDSC_92922 |
| UAS-BAIAP2L1        | BDSC   | RRID:BDSC_78446 |
| UAS-BCHE.HA         | This study | N/A |
| UAS-BCHE.F433V.HA   | This study | N/A |
| UAS-BEST3           | This study | N/A |
| UAS-BEST3.R130S     | This study | N/A |
| UAS-BMP1.G927S.HA   | BDSC   | RRID:BDSC_92931 |
| UAS-BMP1.HA         | BDSC   | RRID:BDSC_77944 |
| UAS-CAMK2A          | This study | N/A |
| UAS-CAMK2A.E183V    | This study | N/A |
| UAS-CARS.HA         | BDSC   | RRID:BDSC_79001 |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| UAS-CARS.N348S.HA  | BDSC   | RRID:BDSC_92918 |
| UAS-CASK            | This study | N/A |
| UAS-CAT.HA          | BDSC   | RRID:BDSC_78471 |
| UAS-CAT.G204E.HA    | This study | N/A |
| UAS-CEP135.HA       | BDSC   | RRID:BDSC_78458 |
| UAS-CEP135,S947P.HA | This study | N/A |
| UAS-CHST2.HA        | BDSC   | RRID:BDSC_78472 |
| UAS-CHST2.R52PHA    | This study | N/A |
| UAS-CLCNKB.HA       | BDSC   | RRID:BDSC_77934 |
| UAS-CLCNKB.M176L.HA | BDSC   | RRID:BDSC_92910 |
| UAS-CLIP2.G13W.HA   | BDSC   | RRID:BDSC_92934 |
| UAS-CLIP2.HA        | BDSC   | RRID:BDSC_78461 |
| UAS-CSAD.HA         | BDSC   | RRID:BDSC_78498 |
| UAS-CSAD.A411V.HA   | This study | N/A |
| UAS-CTNNB1          | BDSC   | RRID:BDSC_78496 |
| UAS-CTNNB1.T551M    | This study | N/A |
| UAS-DDR2.B          | BDSC   | RRID:BDSC_78483 |
| UAS-DLC1.HA         | BDSC   | RRID:BDSC_78089 |
| UAS-DPP6            | This study | N/A |
| UAS-DPYSL2.HA       | BDSC   | RRID:BDSC_77938 |
| UAS-DPYSL2.R496C.HA | BDSC   | RRID:BDSC_92946 |
| UAS-DPYSL3.HA       | BDSC   | RRID:BDSC_77939 |
| UAS-DPYSL3.V139I.HA | BDSC   | RRID:BDSC_92945 |
| UAS-ELAVL3          | BDSC   | RRID:BDSC_78505 |
| UAS-ELAVL3.L186P    | BDSC   | RRID:BDSC_92912 |
| UAS-EP400.B         | BDSC   | RRID:BDSC_78493 |
| UAS-EPHA1.HA        | BDSC   | RRID:BDSC_77931 |
| UAS-EPHA1.V567L.HA  | BDSC   | RRID:BDSC_92963 |
| UAS-EPHB1           | BDSC   | RRID:BDSC_78473 |
| UAS-EPHB1.V916M     | BDSC   | RRID:BDSC_92932 |
| UAS-EPT1            | BDSC   | RRID:BDSC_78485 |
| UAS-EPT1.H82N       | This study | N/A |
| UAS-EXD2.E513D.HA   | BDSC   | RRID:BDSC_92942 |
| UAS-EXD2.HA         | BDSC   | RRID:BDSC_77933 |
| UAS-FGGY            | BDSC   | RRID:BDSC_78091 |
| UAS-GCLC.HA         | This study | N/A |
| UAS-GCLC.R128W.HA   | This study | N/A |
| UAS-GLRA2.HA        | BDSC   | RRID:BDSC_77954 |
| UAS-GLRA2.N136S.HA  | BDSC   | RRID:BDSC_92915 |
| UAS-GNA01.HA        | BDSC   | RRID:BDSC_79003 |
| UAS-GPCS.HA         | BDSC   | RRID:BDSC_77936 |
| UAS-GPCS.M133T.HA   | BDSC   | RRID:BDSC_92913 |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| UAS-GRIA1.HA        | BDSC   | RRID:BDSC_78474 |
| UAS-GRIA1.R218H.HA  | BDSC   | RRID:BDSC_92930 |
| UAS-GRIK5           | BDSC   | RRID:BDSC_77957 |
| UAS-GRK4.HA         | BDSC   | RRID:BDSC_78503 |
| UAS-GRK4.P385A.HA   | BDSC   | RRID:BDSC_92916 |
| UAS-HTR1D.HA        | This study | N/A |
| UAS-HTR1D.T99N.HA   | This study | N/A |
| UAS-IGF2R.HA        | BDSC   | RRID:BDSC_78454 |
| UAS-IRF2BPL         | BDSC   | RRID:BDSC_78509 |
| UAS-IRF2BPL.F30L    | This study | N/A |
| UAS-IRF2BPL.N701X   | This study | N/A |
| UAS-ITGA8.HA        | BDSC   | RRID:BDSC_78501 |
| UAS-ITGA8.R748C.HA  | BDSC   | RRID:BDSC_92924 |
| UAS-JUPHA           | BDSC   | RRID:BDSC_78500 |
| UAS-JUPN690.S.HA    | This study | N/A |
| UAS-KCND3.HA        | BDSC   | RRID:BDSC_78475 |
| UAS-KCND3.R86PHA    | BDSC   | RRID:BDSC_92921 |
| UAS-KCNH8.HA        | BDSC   | RRID:BDSC_78470 |
| UAS-KDM2A           | BDSC   | RRID:BDSC_78476 |
| UAS-KDM2A.R449K     | BDSC   | RRID:BDSC_92927 |
| UAS-KDR             | BDSC   | RRID:BDSC_78451 |
| UAS-KDR.D1171N      | BDSC   | RRID:BDSC_92955 |
| UAS-LAMA2           | BDSC   | RRID:BDSC_79000 |
| UAS-LRCH4           | BDSC   | RRID:BDSC_78477 |
| UAS-LRCH4.V42M      | BDSC   | RRID:BDSC_92928 |
| UAS-MADD            | BDSC   | RRID:BDSC_78457 |
| UAS-MADD.R514C      | BDSC   | RRID:BDSC_92933 |
| UAS-MANBA           | BDSC   | RRID:BDSC_77960 |
| UAS-MAP4K1.HA       | BDSC   | RRID:BDSC_77946 |
| UAS-MAP4K1.M725T.HA | This study | N/A |
| UAS-MBNL1.HA        | BDSC   | RRID:BDSC_78467 |
| UAS-MBNL1.V45A.HA   | BDSC   | RRID:BDSC_92908 |
| UAS-MEGF11          | BDSC   | RRID:BDSC_78460 |
| UAS-MEGF11.R911C    | This study | N/A |
| UAS-MINK1.HA        | BDSC   | RRID:BDSC_78489 |
| UAS-MINK1.C269R.HA  | This study | N/A |
| UAS-MYH9.HA         | BDSC   | RRID:BDSC_79002 |
| UAS-MYH9.R1571Q.HA  | This study | N/A |
| UAS-NCOR1           | BDSC   | RRID:BDSC_78486 |
| UAS-NCOR1.P569S     | BDSC   | RRID:BDSC_92953 |
| UAS-NID2            | This study | N/A |
| UAS-NLGN1           | This study | N/A |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| UAS-NLGN1.H795Y     | BDSC   | RRID:BDSC_92936 |
| UAS-NLGN3.R195W     | This study | N/A |
| UAS-NOS3            | This study | N/A |
| UAS-NPFR2           | BDSC   | RRID:BDSC_78478 |
| UAS-NPFR2.M163I     | BDSC   | RRID:BDSC_92917 |
| UAS-NR2F1           | BDSC   | RRID:BDSC_77959 |
| UAS-NR2F1.R404H     | This study | N/A |
| UAS-NTN1.A449D      | BDSC   | RRID:BDSC_92958 |
| UAS-NTN1            | BDSC   | RRID:BDSC_78495 |
| UAS-NTN5            | BDSC   | RRID:BDSC_78507 |
| UAS-P4HA2.G153E.HA  | BDSC   | RRID:BDSC_92938 |
| UAS-P4HA2.HA        | BDSC   | RRID:BDSC_77935 |
| UAS-PD2             | BDSC   | RRID:BDSC_77928 |
| UAS-PDC1            | This study | N/A |
| UAS-PDGFRB          | BDSC   | RRID:BDSC_92941 |
| UAS-PDGFRB.A366T.HA | This study | N/A |
| UAS-PDK2.HA         | BDSC   | RRID:BDSC_77949 |
| UAS-PDK2.R190Q      | BDSC   | RRID:BDSC_92962 |
| UAS-PDK2.R190Q.HA   | BDSC   | RRID:BDSC_77919 |
| UAS-PEAR1           | BDSC   | RRID:BDSC_77806 |
| UAS-PEAR1.T824I     | BDSC   | RRID:BDSC_92911 |
| UAS-PDLC1           | This study | N/A |
| UAS-PDLC1.R42Q      | BDSC   | RRID:BDSC_78499 |
| UAS-PRKD1           | BDSC   | RRID:BDSC_78494 |
| UAS-PRKD1.R441W     | BDSC   | RRID:BDSC_92950 |
| UAS-PITX1.HA        | BDSC   | RRID:BDSC_78497 |
| UAS-PITX1.L242F.HA  | BDSC   | RRID:BDSC_92914 |
| UAS-PITX1.L242F.HA  | BDSC   | RRID:BDSC_78404 |
| UAS-PITX1.L242F.HA  | BDSC   | RRID:BDSC_92909 |
| UAS-PRPS1L1         | This study | N/A |
| UAS-PRPS1L1.G61D    | BDSC   | RRID:BDSC_92929 |
| UAS-PTK7.R570Q      | This study | N/A |
| UAS-PTPRF           | BDSC   | RRID:BDSC_92404 |
| UAS-PTPRF.S334R     | BDSC   | RRID:BDSC_92937 |
| UAS-PXDN.HA         | BDSC   | RRID:BDSC_77955 |
| UAS-PXDN.R643Q      | BDSC   | RRID:BDSC_92951 |
| UAS-RALGAPA1        | BDSC   | RRID:BDSC_78449 |
| UAS-RALGAPA1.LL1769L| This study | N/A |
| UAS-RMS2.HA         | BDSC   | RRID:BDSC_77937 |
| UAS-SCARB2.HA       | BDSC   | RRID:BDSC_77929 |
| UAS-SCARB2.V173A.HA | This study | N/A |
| UAS-SDK2.HA         | BDSC   | RRID:BDSC_78487 |
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| UAS-SEC14L5         | BDSC   | RRID:BDSC_78092 |
| UAS-SH2D3C.HA       | BDSC   | RRID:BDSC_78492 |
| UAS-SH2D3C.R227Q.HA | This study | N/A |
| UAS-SLC23A1         | BDSC   | RRID:BDSC_78459 |
| UAS-SLC23A1.L465M   | BDSC   | RRID:BDSC_92943 |
| UAS-SLC8A2          | BDSC   | RRID:BDSC_78490 |
| UAS-SLC8A2.G792R    | This study | N/A |
| UAS-SLC04A1         | BDSC   | RRID:BDSC_78479 |
| UAS-SLC04A1.V679I   | BDSC   | RRID:BDSC_92920 |
| UAS-SOGA3.HA        | BDSC   | RRID:BDSC_78469 |
| UAS-SOGA3.R873P.HA  | This study | N/A |
| UAS-SRCAP           | BDSC   | RRID:BDSC_78450 |
| UAS-SRCAP.2137del   | This study | N/A |
| UAS-SRCAP.G1937S    | This study | N/A |
| UAS-TANC2           | BDSC   | RRID:BDSC_78452 |
| UAS-TANC2.H1689R    | BDSC   | RRID:BDSC_92925 |
| UAS-TRIP12          | BDSC   | RRID:BDSC_78518 |
| UAS-TRIP12.R1643Q   | BDSC   | RRID:BDSC_92956 |
| UAS-TRPM1           | BDSC   | RRID:BDSC_78517 |
| UAS-TRPM1.F794L     | BDSC   | RRID:BDSC_92954 |
| UAS-TRPM6.HA        | BDSC   | RRID:BDSC_77958 |
| UAS-TRPM6.A641E.HA  | BDSC   | RRID:BDSC_92939 |
| UAS-TRPM6.T2011P.HA | BDSC   | RRID:BDSC_92940 |
| UAS-TRPM7.HA        | BDSC   | RRID:BDSC_78447 |
| UAS-TRPM7.T379A.HA  | BDSC   | RRID:BDSC_92923 |
| UAS-TSC2.HA         | BDSC   | RRID:BDSC_78465 |
| UAS-TSC2.R1557W.HA  | BDSC   | RRID:BDSC_92949 |
| UAS-TSC2.R548M.HA   | BDSC   | RRID:BDSC_92947 |
| UAS-TULP4           | BDSC   | RRID:BDSC_92408 |
| UAS-USP30.HA        | BDSC   | RRID:BDSC_78480 |
| UAS-USP30.P200S.HA  | BDSC   | RRID:BDSC_92957 |
| UAS-YIPF5.HA        | BDSC   | RRID:BDSC_82197 |
| UAS-ZMYND8.HA       | BDSC   | RRID:BDSC_77945 |

Deposited data

- **TG4 imaging videos**: This paper: Mendeley Data [https://doi.org/10.1016/10.17632/64gz799sb.1](https://doi.org/10.1016/10.17632/64gz799sb.1)

- **Human GLRA2 variant p.Thr296Met**: ClinVar SCV002055997
- **Human GLRA2 variant p.Phe475Ser**: ClinVar SCV002056017
- **Human GLRA2 variant p.Ile259Met**: ClinVar SCV002056018
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Human GLRA2 variant p.Arg252Cys | ClinVar | SCV002056022 |
| Human GLRA2 variant p.Ala288Thr | ClinVar | SCV002056021 |
| Human GLRA2 variant p.Pro396Thr | ClinVar | SCV002056020 |
| Human GLRA2 variant p.Pro400Leu | ClinVar | SCV002056019 |
| Human GLRA2 variant p.Arg445Gln | ClinVar | SCV002056023 |

Recombinant DNA

| REAGENT or RESOURCE | SOURCE |
|---------------------|--------|
| pUASg-HA.attB | Drosophila Genomics Resource Center, DGRC_1423 |

Software and algorithms

| REAGENT or RESOURCE | SOURCE |
|---------------------|--------|
| Prism8 | Graph Pad | https://www.graphpad.com |
| Zen Blue | Zeiss | https://www.zeiss.com/microscopy/us/products/microscope-software/zen-lite.html |
| Zen Black | Zeiss | https://www.zeiss.com/microscopy/us/products/microscope-software/zen-lite.html |
| Snapgene | Snapgene | https://www.snapgene.com |
| Imaris | Imaris | https://imaris.oxinst.com/ |
| Lab Chart | Ad instruments | https://www.adinstruments.com/products/labchart |