

# Resource Summary Report

Generated by [ASWG](#) on Jul 31, 2026

## w[1118]; Fmr1[Delta50M]/TM6B, Tb[1]

RRID:BDSC\_6930

Type: Organism

### Proper Citation

RRID:BDSC\_6930

### Organism Information

**URL:** <https://n2t.net/bdsc:6930>

**Proper Citation:** RRID:BDSC\_6930

**Description:** Drosophila melanogaster with name w[1118]; Fmr1[Delta50M]/TM6B, Tb[1] from BDSC.

**Species:** Drosophila melanogaster

**Notes:** Donor: Kendal Broadie, Vanderbilt University

**Affected Gene:** Fmr1, Tb, w

**Genomic Alteration:** Chromosome 1, Chromosome 3

**Catalog Number:** 6930

**Database:** Bloomington Drosophila Stock Center (BDSC)

**Database Abbreviation:** BDSC

**Availability:** available

**Alternate IDs:** BDSC\_6930, BL6930

**Organism Name:** w[1118]; Fmr1[Delta50M]/TM6B, Tb[1]

**Record Creation Time:** 20240911T222204+0000

**Record Last Update:** 20260725T194839+0000

### Ratings and Alerts

No rating or validation information has been found for w[1118]; Fmr1[Delta50M]/TM6B, Tb[1].

No alerts have been found for w[1118]; Fmr1[Delta50M]/TM6B, Tb[1].

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## Data and Source Information

**Source:** [Integrated Animals](#)

**Source Database:** Bloomington Drosophila Stock Center (BDSC)

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## Usage and Citation Metrics

We found 8 mentions in open access literature.

**Listed below are recent publications.** The full list is available at [ASWG](#) .

Gurijala A, et al. (2025) Interaction between neuromuscular junction metabolic requirements in fragile X syndrome and glycogen storage disease models. Disease models & mechanisms, 18(8).

Geng J, et al. (2023) Deregulation of ER-mitochondria contact formation and mitochondrial calcium homeostasis mediated by VDAC in fragile X syndrome. Developmental cell, 58(7), 597.

Leahy SN, et al. (2023) FMRP activity and control of Csw/SHP2 translation regulate MAPK-dependent synaptic transmission. PLoS biology, 21(1), e3001969.

Golovin RM, et al. (2021) Neuron-Specific FMRP Roles in Experience-Dependent Remodeling of Olfactory Brain Innervation during an Early-Life Critical Period. The Journal of neuroscience : the official journal of the Society for Neuroscience, 41(6), 1218.

Kennedy T, et al. (2020) Genetic background mutations drive neural circuit hyperconnectivity in a fragile X syndrome model. BMC biology, 18(1), 94.

Kim N, et al. (2019) BMP-dependent synaptic development requires Abi-Abl-Rac signaling of BMP receptor macropinocytosis. Nature communications, 10(1), 684.

Russo A, et al. (2019) Wnd/DLK Is a Critical Target of FMRP Responsible for Neurodevelopmental and Behavior Defects in the Drosophila Model of Fragile X Syndrome. Cell reports, 28(10), 2581.

Kennedy T, et al. (2017) Fragile X Mental Retardation Protein Restricts Small Dye Iontophoresis Entry into Central Neurons. The Journal of neuroscience : the official journal of the Society for Neuroscience, 37(41), 9844.