

NEWLY DIAGNOSED GUIDE



Life doesn't end when FTD is diagnosed. AFTD provides information and resources to help you adjust to the changes ahead. Please contact the **AFTD HelpLine** with questions at **866-507-7222** or **info@theaftd.org**.

LEARN ABOUT THE DISEASE

- ☐ Confirm the diagnosis.
- ☐ Learn about FTD symptoms, genetics, and what you might expect over the course of the disease.
 - AFTD's website (www.theaftd.org) is a great place to start.
 - Double-check information found online. Use websites you can trust and confirm with experts.
 - AFTD recommends meeting with a genetic counselor to learn the benefits and limitations of genetic testing.
- ☐ Sign up for AFTD's newsletters to keep informed about the latest FTD news.
- ☐ Plan to attend the AFTD Education Conference. AFTD offers modest travel grants to help with costs if needed.

CREATE YOUR CARE TEAM

- ☐ Identify professionals (primary care physician, neurologist, psychiatrist, case manager/social worker) and coordinate their services.
- ☐ Obtain copies of diagnostic evaluations for your records. Organizing paperwork helps future healthcare providers.
- ☐ Keep a log or journal that includes:
 - Changes in behavior. AFTD has a **Behavioral Tracker** for caregivers and a **Temperature Tracker** for persons with FTD.
 - Medications started or discontinued.
 - Issues you have or questions you want to ask at your next doctor's appointment.
- ☐ If appropriate, consult an occupational therapist (OT), physical therapist (PT), or speech therapist for evaluation and techniques to maximize abilities.
- ☐ Visit www.theaftd.org to find an in-person or virtual support group for either care partners/caregivers or persons with FTD.
- ☐ **Keep a list of tasks family, friends and neighbors can help with. When they offer, say "yes!"**

FOCUS ON WELLNESS AND A POSITIVE DAILY ROUTINE

- ☐ Follow a daily routine to structure the day that includes a healthy diet and regular exercise.
- ☐ Keep tabs on your mental health and don't be afraid to reach out for help.
- ☐ Stay active with friends and interests. Adapt activities according to strengths and needs.
- ☐ If needed, apply for AFTD's Comstock Grants—these can be for care partner respite, conference travel, or for a quality-of-life stipend for the person with FTD.

ADDRESS SAFETY ISSUES REGULARLY AND MAKE CHANGES BEFORE A CRISIS OCCURS

- ☐ Keep home environment safe and equipped to reduce risk of falls. Occupation or physical therapy can help you access your home for safety.
- ☐ Where judgment is impaired, monitor bank accounts, investments and online activity; change access as needed to protect assets.
- ☐ Use GPS monitoring or similar device if getting lost is a risk.
- ☐ Learn the laws where you live regarding driving privileges.
- ☐ If behaviors warrant, notify local law enforcement of diagnosis.

ADDRESS LEGAL AND FINANCIAL ISSUES

- ☐ Consult an elder law attorney to help you complete legal documents (power of attorney, living will, trusts, etc.).
- ☐ Plan transition from employment, if still working.
- ☐ Review long-term financial plan and options if care needs increase (assisted living, nursing home facilities).
- ☐ Apply for Social Security Disability Insurance using the Compassionate Allowances program, which includes FTD as an eligible diagnosis.
- ☐ Veterans may be able to obtain benefits from the VA – [check here](#) for eligibility.
- ☐ Research day programs and long-term care facilities early for optimal planning.

PARTICIPATE IN RESEARCH

- ☐ Join the FTD Disorders Registry (www.FTDRegistry.org).
- ☐ Participate in the ALLFTD study to support development of treatments.
- ☐ There are studies for care partners, caregivers, persons with FTD and their children. *Some clinical trials are now actively recruiting persons with FTD meeting specific criteria, with more expected on the horizon.*
- ☐ Consider **brain donation** to confirm diagnosis and advance research. Arrangements must be made well in advance.
- ☐ Visit the **Studies Seeking Participants** page on AFTD's website to learn more.

The Association for Frontotemporal Degeneration

theaftd.org | HelpLine: 1.866.507.7222 | info@theaftd.org

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