

Who is FAST?

Foundation for Angelman Syndrome Therapeutics is run by a driven board of parents of individuals living with Angelman syndrome and professionals dedicated to curing AS through the funding of an aggressive research agenda.

FAST operates as an impartial organization to drive collaboration and sharing across the industry (pharmaceuticals, research institutions and other global organizations) to reach a cure faster. FAST is served by two boards: the board of directors and an independent scientific advisory board. Together, we are working hard to bring practical treatment into current medical practice as quickly as possible.

FAST is committed to finding meaningful and transformative treatments for all individuals living with AS globally, regardless of age or genotype in order to achieve our mission of finding a cure.

“The stars are aligned in favor of Angelman syndrome being amenable to a really curative kind of treatment. I think FAST has transformed the landscape of Angelman research.”

– Art Beaudet, M.D. National Academy of Sciences

