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Profile Function

chief of service

Publications (58)

Wendebourg M, Weigel M, Weidensteiner C, Sander L, Kesenheimer E, Naumann N, Haas T, Madoerin P, Braun N, Neuwirth C, Weber M, Jahn K, Kappos L, Granziera C, Schweikert K, Sinnreich M, Bieri O, Schlaeger R. Cervical and thoracic spinal cord gray matter atrophy is associated with disability in patients with amyotrophic lateral sclerosis. *Eur J Neurol* 2024:e16268.

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Projects (18)

A Phase 3, Randomized, Double-blind, Placebo-controlled, Parallel, Multicenter Study to Evaluate the Safety and Efficacy of ALXN1720 in Adults with Generalized Myasthenia Gravis <i>Clinical Studies - Feb 1, 2024 - Apr 4, 2027</i> <i>Ongoing</i>
Sicherheit und Wirksamkeit von ION363 bei Patienten mit genetischer ALS mit Fused in Sarcoma Mutation (FUS-ALS), Phase 1-3 <i>Clinical Studies - Apr 3, 2023 - Apr 3, 2028</i> <i>Ongoing</i>
TEAR und EARLY ALS <i>Fundamental Research - Sep 1, 2021 - Mar 31, 2024</i> <i>Automatically Closed</i>
Study to Assess the Safety, Tolerability, Pharmacokinetics, and Effect on Disease Progression of BIIB078 Administered to Previously Treated Adults C9ORF72-Associated Amyotrophic Lateral Sclerosis (ALS) <i>Clinical Studies - Apr 28, 2020 - Jul 28, 2023</i> <i>Automatically Closed</i>
ALXN1210-ALS-308: An Efficacy and Safety Study of Ravulizumab in ALS Patients <i>Clinical Studies - Mar 30, 2020 - Oct 17, 2021</i> <i>Aborted</i>
SMARtCare: Longitudinal data assessment from patients with spinal muscular atrophy <i>Clinical Studies - Dec 9, 2019 - Dec 31, 2030</i> <i>Ongoing</i>
Arimoclomol in Amyotrophic Lateral Sclerosis - Open Label Extension Trial <i>Clinical Studies - Sep 19, 2019 - Jul 2, 2021</i> <i>Aborted</i>
A Phase 1 Multiple-Ascending-Dose Study to Assess the Safety, Tolerability, and Pharmacokinetics of BIIB078 Administered Intrathecally to Adults with C9ORF72-Associated Amyotrophic Lateral Sclerosis <i>Clinical Studies - Sep 10, 2018 - Nov 17, 2021</i> <i>Completed</i>
Arimoclomol in Amyotrophic Lateral Sclerosis <i>Clinical Studies - Jul 31, 2018 - Dec 18, 2020</i> <i>Aborted</i>

Methodologische Studie neuartiger elektrophysiologischer, physikalischer und bildgebender Messgrößen zur Beurteilung der Krankheitsprogression bei amyotropher Lateralsklerose

Clinical Studies - May 1, 2016 - Jun 1, 2018

Automatically Closed

Methodology Study of Novel Outcome Measures to Assess Progression of ALS

Clinical Studies - Apr 2, 2016 - Aug 14, 2019

Completed

Survival, Trigger and Risk, Epigenetic, eNviromental and Genetic Targets for motor neuron Health (STRENGTH)

Clinical Studies - Jan 1, 2015 - Dec 31, 2017

Completed

Randomized placebo-controlled crossover trial with quinine sulfate for the treatment of cramps in amyotrophic lateral sclerosis (ALS)

Clinical Studies - Nov 1, 2013 - Mar 26, 2020

Automatically Closed

Swiss Registry for Neuromuscular Disorders

Clinical Studies - Jan 1, 2012 - Dec 30, 2030

Ongoing

A novel neurophysiological measurement (MUNIX) in neuromuscular disorders

Clinical Studies - Mar 26, 2011 - Mar 26, 2020

Automatically Closed

Validierung der Awaji-Kriterien

Clinical Studies - Mar 20, 2011 - Mar 20, 2014

Completed

PEG: Perkutane Endoskopische Gastrostomie-Register für Patienten mit amyotropher Lateralsklerose (ALS) in Deutschland und Schweiz. Multizentrische Beobachtungsstudie

Clinical Studies - Dec 15, 2010 - Dec 14, 2014

Completed

Autopsy study in ALS

Clinical Studies - Jan 1, 1999 - Jan 1, 2003

Automatically Closed

