

SCITrialsFinder.net - SCI Trials of Rehabilitation and Technological Interventions to Improve Functional Outcomes

Revised January 31, 2023 - Listing of 170 Trials

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
NCT05664646	VA Office of Research and Development	Autonomic Effects of Stimulation in SCI	Age 35-50 years SCI level C3-T1 AIS A-B SCI > 1 year	Not yet recruiting	20	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: Single (Participant)	Technology	General health	TSCS F/U 2 years	July 2023	December 27, 2022	December 27, 2022	Bronx, NY, USA
NCT05665998	Ecole Polytechnique Fédérale de Lausanne	Brain Controlled Spinal Cord Stimulation in Participants With Cervical Spinal Cord Injury for Upper Limb Rehabilitation	Age 18-75 years SCI level C1-C8 AIS A-D Traumatic SCI > 6 months	Not yet recruiting	3	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General health	ARAT GRASSP CUE-T ISNSCI F/U 12 months	March 2023	December 27, 2022	December 27, 2022	
NCT05659121	National University Hospital, Singapore	Improving Mobility Via Robotic Exoskeletons in Local Rehabilitation Settings in Singapore	Age 21-90 years FAC 0-3	Recruiting	400	Phase: Not Applicable, Primary Purpose: Health Services Research, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Rehabilitation	Standing/walking mobility	10 MWT 6MWT COVS EQ-5D WISCI F/U 6 months	November 2018	December 21, 2022	December 21, 2022	Singapore, Singapore
NCT05644522	Shirley Ryan AbilityLab	Nomad P-KAFO Study	Age 18-89 years NO Non-correctable knee varus/valgus in excess of 15 degrees	Not yet recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Standing/walking mobility	10MWT 6MWT BBS FGA TUG ABC WHOQOL EQ5D-5L NPRS F/U 3 months	January 2023	December 9, 2022	December 9, 2022	
NCT05644171	National Neuroscience Institute	RESTORES Trial: RESToration Of Rehabilitative Function With Epidural Spinal Stimulation	Age > 21 years AIS A-B SCI level T2-L1 SCI > 1 year	Not yet recruiting	3	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General health	F/U 2 years	December 2022	December 9, 2022	December 9, 2022	
NCT05645003	Afyonkarahisar Health Sciences University	Transcranial Magnetic Stimulation Therapy in Neuropathic Painful Spinal Cord Injury Patients	Age 20-70 years	Recruiting	60	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor)	Technology	Pain	SF-36 BDI F/U 6 weeks	November 2022	December 9, 2022	December 9, 2022	Afyonkarahisar, Turkey
NCT05643313	ABLE Human Motion S.L.	Clinical Investigation on Feasibility and Usability of the ABLE Exoskeleton Device for Individuals With Spinal Cord Injury to Perform Skills for Home and Community Environments	Age 18-70 years AIS A-B SCI level T2-L1 SCI > 6 months	Enrolling by invitation	10	Phase: Not Applicable, Primary Purpose: Device feasibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Rehabilitation	Standing/walking mobility	10MWT 6MWT QUEST 2.0 SCIM TUG WISCI WHOQOL-BREF F/U 11 weeks	November 2022	December 8, 2022	December 29, 2022	Badalona, Spain
NCT05637775	I.R.C.C.S. Fondazione Santa Lucia	DISCloser: Improving Arm Sensorimotor Functions After Spinal Cord Injury Via Brain-Computer Interface Training	Age 18-70 years AIS A-D SCI level C1-T1 SCI 30-90 days	recruiting	30	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single (Outcomes Assessor)	Technology	Arm/hand function	GRASSP ISCI SCIM F/U 48 hours	November 2022	December 5, 2022	December 5, 2022	Rome, Italy
NCT05626816	MetroHealth Medical Center	Acute Genital Nerve Stimulation for Neurogenic Bowel Dysfunction in Individuals Living With Spinal Cord Injury	Age > 18 years SCI level C1-T12 AIS A-D SCI > 6 months NO cardiac pacemaker, implanted defibrillator or other implanted FES device	Recruiting	52	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single (Participant)	Technology	Bowel health	SCI-QOL Bowel Management Difficulties ARM outcomes SCI Bowel function basic dataset F/U 3 weeks	January 2023	November 25, 2022	January 5, 2023	Cleveland, OH, USA
NCT05619146	Shirley Ryan AbilityLab	Arm and Leg Cycling for Accelerated Recovery From SCI	Age 18 - 65 years Traumatic SCI level C1-T10 AIS C-D SCI > 1 year NO Significant other disease (ex: cardiological or heart disease, renal, hepatic, malignant tumors, mental or psychiatric disorders) that would prevent participants from fully engaging in study procedures NO implanted pacemakers NO tremor	Not yet recruiting	5	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Single Group Assignment, Masking: None (open label)	Rehabilitation	Standing/walking mobility	10 MWT 6MWT MAS BBS EMG WISCI F/U 120 days	December 2022	November 16, 2022	November 16, 2022	Chicago, IL, USA
NCT05615766	University of Liverpool	Assessment of a Robotic Exoskeleton for Upper Limb Rehabilitation	Age > 18 years SCI level C5-C8 AIS C-D	Not yet recruiting	9	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Single Group Assignment, Masking: Single (Investigator)	Technology	Arm/Hand Function	MAS GRASSP SCIM COPM F/U 12 weeks	March 2023	November 14, 2022	November 14, 2022	
NCT05605912	Sint Maartenskliniek	Myosuit in Incomplete Spinal Cord Injury	Age > 18 years AIS C-D NO other neurological disease NOT taller than 195 cm or smaller than 150 cm NO psychiatric background	Recruiting	34	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Rehabilitation	Standing/walking mobility	10 MWT 6MWT EQ-5D SCI-FAP F/U 23 weeks	October 2022	November 4, 2022	November 4, 2022	Ubbergen, Netherlands

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NCT05601661	The University of Texas Health Science Center, Houston	Safety and Feasibility of Paired Vagus Nerve Stimulation With Rehabilitation for Improving Upper Extremity Function in People With Cervical Spinal Cord Injury	Age > 18 years SCI level C1-C8 AIS B-D Traumatic SCI > 12 months NO history of prior injury to vagus nerve	Recruiting	8	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Triple (Participant, Investigator, Outcomes Assessor)	Technology	General health	TRI-HFT CUE-Q SCIM SCI-QoL GRASSP F/U 132 days	February 2023	November 1, 2022	January 25, 2023	Houston, TX, USA
NCT05593237	University of California, San Francisco	Transcranial Magnetic Stimulation for Chronic Neuropathic Pain	Age 18 - 80 years meeg criteria for chronic neuropathic pain NO implanted device	Recruiting	20	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Crossover Assignment Masking: Single (Participant)	Technology	Pain	• BDI • PSQI • MPQ • WHOQOL-BREF F/U 6 months	April 2022	October 25, 2022	January 6, 2023	San Francisco, CA, USA
NCT05589415	University of Texas Rio Grande Valley	Targeted HD-IDCS to Improve Upper Limb Rehabilitation in SCI	Age 18 - 75 years SCI level C2-T1 AIS C-D SCI > 18 months NO history of seizures NO other neurological impairment or condition	Recruiting	24	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Double (Participant, Outcomes Assessor)	Technology	Arm/Hand Function	• GRASSP • MEP • EMG • MVC F/U 14 days	November 2020	October 21, 2022	October 21, 2022	Harlingen, TX, USA
NCT05563103	Spaulding Rehabilitation Hospital	Combination Therapy to Improve SCI Recovery.	Age 18 - 70 years SCI level C2-L2 AIS C-D SCI >12 months NO active implanted devices	Not yet recruiting	60	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Triple (Participant, Care Provider, Outcomes Assessor)	Rehabilitation	Standing/walking mobility	• 10MWT • 6MWT • TUG • NPRS • CVLT 12 weeks	October 2022	October 3, 2022	October 3, 2022	Chicago, IL, USA Cambridge, MA, USA
NCT05558254	Spinal Cord Injury Centre of Western Denmark	ROBERT® as an Intervention to Enhance Muscle Strength After Spinal Cord Injury	Age ≥18 years AIS C-D SCI <12 months NO previous cerebral injury	Recruiting	12	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Outcomes Assessor)	Technology	Standing/walking mobility	• MVC • EMG F/U 8 weeks	September 2022	September 28, 2022	September 28, 2022	Viborg, Denmark
NCT05550987	University of Rzeszow	The Use of Modern Technologies in Neurorehabilitation	Age 18-75 years SCI level C1-S5	Not yet recruiting	120	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: None (Open Label)	Rehabilitation	Standing/walking mobility	• 10 MWT • Barthel Index • BBS • LOSSCI • TCT • TUG • WHOQOL-BREF F/U immediately after intervention	November 2022	September 22, 2022	November 8, 2022	
NCT05544175	Charles University, Czech Republic	NEUROModulation Pain Therapy in Combination With Intensive Physiotherapy	Age 18-60 years SCI level C1-S5	Not yet recruiting	10	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Single Group Assignment Masking: None (Open Label)	Technology	Standing/walking mobility	• 10MWT • 6MWT • MAS • BBS • TUG • SCIM • WISCI & WISCI II • WHODAS 2.0 F/U 12 months	October 2022	September 16, 2022	October 7, 2022	
NCT05522920	NHS Greater Glasgow and Clyde	Spinal Stimulation for Chronic Complete Tetraplegia	Age ≥18 years AIS A SCI level C3-C7 SCI ≥ 1 year	Not yet recruiting	12	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Single Group Assignment Masking: None (Open Label)	Technology	Arm/Hand Function	• AIS • MAS • GRASSP F/U 38 weeks	August 2022	August 31, 2022	August 31, 2022	
NCT05504200	Royal National Orthopaedic Hospital NHS Trust	Transcutaneous Spinal Cord Stimulation With Bladder and Pelvic Floor Muscle Training	Age > =18 years SCI >= L5 SCI > 6 months Neurogenic detrusor overactivity	Recruiting	25	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Outcomes Assessor)	Technology	Bladder Health	• EQ-5D-5L • NBDS • Qualiveen • EMG F/U 14 weeks	March 2022	August 17, 2022	August 17, 2022	Stanmore, UK
NCT05503316	University Hospital, Ghent	The Roll of Balance Confidence in Gait Rehabilitation in Persons With a Lesion of the Central Nervous System	Age 18-70 years NO other neurological condition	Recruiting	42	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: None (open label)	Rehabilitation	Standing/walking mobility	• 10MWT • ABC • HADS F/U 2 weeks	44805	August 16, 2022	September 10, 2022	Ghent, Belgium
NCT05473676	University of Calgary	Robotic Walking for Children Who Cannot Walk	Age 4 and older (children) Unable to walk independently due to pediatric onset, non-progressive central nervous system disorder or injury (ie. cerebral palsy or acquired brain injury)	Recruiting	12	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Single Group Assignment Masking: None (open label)	Technology	Standing/Walking Mobility	• COPM • ECAB • EQ-5D-Y • CarerQOL • CP-CHILD F/U 28 weeks	July 2022	July 26, 2022	August 11, 2022	Calgary, Canada

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NCT05473689	University Health Network, Toronto	Outcomes Post Treatment: Impact on Motor Impairment of Sleep Efficiency in SCI (OPTIMISE SCI Trial)	Age 18 and older SCI level C2-T12 AIS A-D SCI < 30 days NOT being treated for sleep apnea prior to the spinal cord impairment onset NOT having concomitant diseases of the central nervous system NOT having Neuromuscular diseases NOT having known history of primary hypersomnia or secondary hypersomnia of any cause except for SRBDs (e.g., hypothyroidism, moderate or severe iron deficiency anemia, infections, depression, kidney failure, chronic fatigue syndrome, neurodegenerative diseases, and myotonic dystrophy) NOT having epilepsy	Recruiting	66	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Outcomes Assessor)	Rehabilitation	General Health	• AIS • DASS-21 • FSS • PHQ-9 • SCIM F/U 6 months	August 2022	July 26, 2022	November 1, 2022	Toronto, Canada
NCT05472584	Washington University School of Medicine	Spinal Cord Stimulation and Training	Age 16 - 65 years SCI level C4-T9 AIS A-D SCI > 1 year NOT having a concomitant neurologic disease (traumatic brain injury (TBI), multiple sclerosis (MS), stroke or peripheral neuropathy) NOT having depression, anxiety or cognitive disorder	Recruiting	120	Phase: Not Applicable Primary Purpose: Basic Science Intervention Model: Crossover Assignment Masking: None (open label)	Technology	Arm/hand function	• MEP F/U 4 weeks	May 2023	July 25, 2022	January 31, 2023	Saint Louis, MS, USA
NCT05470478	VA Office of Research and Development	BCI Optimization for Veterans With Paralysis	Age 18 - 80 years	Not yet recruiting	2	Phase: Not Applicable Primary Purpose: Other Intervention Model: Single Group Assignment Masking: None (open label)	Technology	General Health	• IBCI F/U 1 month	January 2023	July 22, 2022	July 22, 2022	Providence, RI, USA
NCT05465486	Aristotle University Of Thessaloniki	NeuroSuitUp: Neurorehabilitation Through Synergistic Man-machine Interfaces	Age 14 and older SCI level C4-L2 AIS A-C SCI > 2 years NOT have Other neurological condition that has a possibility to significantly affect the neurological status of the participants (or) the ability to control a BCI (or) the neurophysiological recordings NOT have multiple Sclerosis NOT have amyotrophic Lateral Sclerosis NOT have parkinson's disease NOT have refractory Epilepsy NOT have Cardiac deficiency NOT have Pulmonary deficiency	Not yet recruiting	20	Phase: Not Applicable Primary Purpose: Basic Science Intervention Model: Parallel Assignment Masking: None (open label)	Technology	Arm/hand function	• SCIM F/U 1 year	August 2022	July 19, 2022	July 19, 2022	Thessaloniki, Greece
NCT05462925	University Health Network, Toronto	Prediction of Muscle Responsiveness to FES Therapy	Age 18 and older SCI level C1-C8 SCI > 6 months tolerate to be in a seating position for at least 1 hour NOT have previous history of any other neuromuscular disorder or conditions that may affect motor response	Recruiting	10	Phase: Not Applicable Primary Purpose: Other Intervention Model: Single Group Assignment Masking: None (open label)	Technology	Arm/hand function	• sEMG F/U 10 weeks	July 2022	July 18, 2022	July 25, 2022	Toronto, Canada
NCT05433064	Buddhist Tzu Chi General Hospital	The study aims to examine the plausible interventional mechanisms underlying the effects of epidural spinal cord stimulation.	Age 20-70 years AIS A-D SCI > 1 year MUST be planning to receive an epidural electrical stimulator	Recruiting	12	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Single Group Assignment Masking: None (Open Label)	Technology	Standing/Walking Mobility	• 10MWT • AIS • MAS • BBS • TUG • WHOQOL F/U 25 months	May 2020	June 27, 2022	September 8, 2022	Hualien City Taiwan
NCT05429736	Shepherd Center, Atlanta GA	Determine if moderate-intensity, motor skill training can improve walking-related outcomes among persons with SCI and to determine if the addition of non-invasive spinal stimulation will result in greater improvements in function compared to training alone.	Age 18-70 years SCI C3 - T12 AIS C,D SCI >= 3 months	Recruiting	40	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor)	Technology	Standing/Walking Mobility	• 10MWT • 2 MWT • SCI-SET • FES • BBS • SCATS F/U 27 days	March 2022	June 23, 2022	November 23, 2022	Atlanta, GA, USA
NCT05425238	Riphah International University	investigate the effects of low load Blood Flow Resistance exercise to improve strength and transfer in lower cervical spinal cord injury	Age 15-50 years SCI level C5-C8 subacute/chronic SCI AIS A-B NO other neurologic conditions	Recruiting	16	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Outcomes Assessor)	Rehabilitation	Arm/Hand function	• Pain • Hand-Held Myometry / Dynamometry • QIF-SF • MAS F/U 6 weeks	May 2022	June 21, 2022	June 21, 2022	Lahore, Pakistan
NCT05423600	Virginia Commonwealth University	The purpose of the current study is to evaluate whether a home-based, telehealth-supported intervention combining Blood Flow Restricted Exercise (BES) and Transspinal Stimulation (TS) will improve motor and functional abilities greater than BES+sham TS in persons with chronic, incomplete tetraplegia.	Age 18-70 years SCI level >= C8 Traumatic SCI > 1 year AIS B-D NO implanted pacemaker	Recruiting	44	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Participant)	Technology	Arm/Hand function	• hand functional activities • grip strength • tactile perception • heart rate variability F/U 20 weeks	June 2022	June 21, 2022	June 29, 2022	Richmind, VA, USA

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NCT05422716	Kessler Foundation	This study aims to expand the knowledge and capacity for neuromodulation to improve the debilitating effects of severe spasticity (spasms, tonic muscle activity and/or clonus) in persons with spinal cord injury (SCI).	Age >= 18 years meet the criteria for an intrathecal baclofen pump	Not yet recruiting	20	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Participant)	Technology	Spasticity	• AIS • MAS • PSFS • EMG • Neuromuscular recovery scale F/U 8 months	July 2022	June 16, 2022	June 16, 2022	
NCT05411692	Riphah International University	yield the long term effects of priming augmented functional electrical stimulations to enhance the tenodesis function of patients with spinal cord injury.	Age 18 - 50 years Sci level C6-C7 AIS D SCI<18 months lack of active palmer and lateral grasp function (except tenodesis grasp function) NO implants NO history of epilepsy NO cardiovascular problems	Recruiting	26	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Outcomes Assessor)	Technology	Arm/Hand function	• SCIM • GRASSP • AIS • Hand-Held Myometry / Dynamometry F/U 6 weeks	March 2022	June 9, 2022	June 9, 2022	Lahore, Pakistan
NCT05390853	Montecatone Rehabilitation Institute S.p.A.	Active tDCS Versus Sham tDCS for Upper Limb Recovery in Incomplete Tetraplegic Patients	Age 18-74 years AIS B-D Sci level C3-C7 Traumatic SCI NO implanted devices NO history of epilepsy NO mechanical ventilation NO history of psychotic disorders	Not yet recruiting	30	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Triple (Participant, Investigator, Outcome Assessor)	Technology	Arm/hand function	• MAS • GRASSP F/U 10 weeks	September 2022	May 25, 2022	May 31, 2022	Imola, BO, Italy
NCT05386537	Kessler Foundation	Combining Wearable Robotic Orthosis With Visual and Haptic Feedback to Enhance the Recovery of Upper Extremity Motor Function and ADL in Persons With Acute SCI	Age 18-80 years AIS C-D <3 months after SCI ability to activate forearm muscles medically stable NO excessive spasticity in elbow NO history of neurological disorder other than SCI NO severe psychiatric disorders NO history of tinnitus	Recruiting	20	Phase: Early Phase 1 Primary Purpose: Treatment Intervention Model: Factorial Assignment Masking: Single (Participant)	Technology	Arm/hand function	• Range of motion arms and hands • sEMG • GRASSP F/U 10 weeks	April 2021	May 23, 2022	May 27, 2022	West Orange, NJ, USA
NCT05376449	HealthPartners Institute	The Effect of an Adaptive Exercise Program on Chronic Inflammation in Spinal Cord Injury	Age 18-70 >6 months after SCI adequate range of motion in elbow and wrist NO cognitive impairment NO causes for shortness of breath NO severe/unstable/uncontrolled autonomic dysreflexia	Recruiting	20	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: None (Open Label)	Rehabilitation	General health	• Inflammatory biomarkers (CRP, IL-6, TNF-alpha) F/U 12 weeks	May 2022	May 17, 2022	June 21, 2022	Saint Paul, MN, USA
NCT05369520	University of British Columbia	Noninvasive Spinal Cord Stimulation for Recovery of Autonomic Function After Spinal Cord Injury	Age 18-60 SCI level C1-T6 AIS A-B Traumatic SCI >1 year impaired lower urinary tract, bowel, sexual function NO History of cardiovascular, respiratory, bladder, or renal disease unrelated to SCI or presence of hydronephrosis or presence of obstructive renal stones NO history of seizures/epilepsy or recurring headache NO history of GI atresia or stenosis NO implanted metal (other than dental)	Not yet recruiting	30	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: None (Open Label)	Technology	General health	• sEMG • HUITT • DARS • ARM • CBF • VFT • SCW F/U 30 weeks	January 2023	May 11, 2022	October 19, 2022	Vancouver, BC, Canada
NCT05337982	Ohio State University	Treadmill walking in combination with electrical stimulation applied to the trunk and legs	• Age 18-85 years • AIS A-D • SCI level C1-T10 • Discharged from inpatient rehabilitation • NO other neurological condition • NO implanted medical devices F/U: 16 weeks	Recruiting	49	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double (Care Provider, Outcomes Assessor)	Technology	Standing/Walking/Mobility	• 10 MWT • 6MWT • ISNCSCI • SCIM • NPSI • ISAFSCI • strength • range of motion • AIS F/U 16 weeks	August 2022	April 20, 2022	July 28, 2022	Columbus, OH, USA
NCT05333770	Kessler Foundation	Evaluate the safety, efficacy and feasibility of high-frequency repetitive transcranial magnetic stimulation (HF-rTMS) in patients with subacute spinal cord injury.	Age 18-55 years inpatient: SCI < 8 weeks outpatient: SCI >= 6 months SCI level C2-C6 AIS A-D NO pacemaker NO metal in the skull NO history of seizures or brain injury	Recruiting	20	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	• GRASSP • Neuro Recovery Scale [NRS] F/U: 6 months	August 2022	April 19, 2022	October 14, 2022	West Orange, NJ, USA
NCT05321017	Medical University of South Carolina	Effect of transcranial magnetic stimulation (TMS) on upper extremity function.	Age > =18 years SCI level at or above C6 SCI > 6 months weak wrist extension NO cognitive impairment NO implanted device	Recruiting	5	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	• Arat • MAS • BBT • GRASSP • MMT • FMA F/U 2.5 months	October 2021	April 11, 2022	April 11, 2022	Charleston, SC, USA
NCT05317832	Temple University	Evaluate a sensor-enabled, just-in-time adaptive intervention (JITAI) strategy to increase and sustain physical activity levels among individuals with spinal cord injury (SCI) in their communities.	Age 18-75 SCI >= 6 months SCI C5 or below	Not yet recruiting	196	Phase: Phase 2, Primary Purpose: Prevention, Intervention Model: Parallel Assignment, Masking: Single (Participant)	Alternative/Complementary	General Health	• Physical activity: Para-SCI • Pain • Quality of Life F&U 24 weeks	February 2023	April 8, 2022	December 12, 2022	Philadelphia, PA, USA

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NCT05305118	Icahn School of Medicine at Mount Sinai	Safety and feasibility of applying Transcutaneous spinal cord stimulation in acute SCI to prevent the development of BP instability	<ul style="list-style-type: none"> Age 18-89 years SCI <= 1 year 	Recruiting	50	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Single Group Assignment: Masking: None (Open Label)	Technology	General health	<ul style="list-style-type: none"> Sit-up Test - Blood Pressure (BP) Severity of Dizziness Scale Sit-up Test - Heart Rate (HR) Sit-Up Test - Cerebral Blood Flow velocity (CBFv) Pain Numeric Rating Scale (NPRS) International SCI Pain Basic Data Set (ISCIPBDS) Spinal Cord Injury Pain Instrument (SCIPI) Autonomic Dysreflexia (AD) Symptoms Survey Orthostatic Hypotension (OH) Symptoms Survey Number of treatment sessions Number of exercise sessions in which TSCS is omitted during therapy Upper Extremity Muscle Strength (UEMS) Testing Lower Extremity Muscle Strength (LEMS) Testing 	March 2022	March 31, 2022	January 9, 2023	New York, NY, USA
NCT05301336	SpineX Inc.	Evaluate the effectiveness and safety of SCONE neuromodulation therapy after 12 weeks of therapy in comparison to inactive sham control in improving symptoms of Neurogenic Lower Urinary Tract Dysfunction	<ul style="list-style-type: none"> Age 18-70 years SCI level C3 - T8 AIS A-D chronic SCI NOT rely on indwelling catheter NO implanted Neuromodulator Not received botox injections in the bladder within the last 12 months Not have a symptomatic bladder infection NO congestive heart failure NO other concurrent neurological disease 	Recruiting	130	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Single (Participant)	Technology	Bladder Health	<ul style="list-style-type: none"> NBSS Urinary Incontinence PGL-I 	May 2022	March 29, 2022	June 9, 2022	Downey, CA, USA San Diego, CA, USA Denver, CO, USA Washington, USA
NCT05267951	University of Washington	Assess the efficacy of non-invasive (transcutaneous) closed-loop electrical spinal cord stimulation for recovery of upper limb function (Aim 1) and spasticity (Aim 2) following spinal cord injury.	<ul style="list-style-type: none"> Age: 21-70 years SCI level >= C8 AIS C-D SCI >= 1 year 	Recruiting	9	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Single (Outcome Assessor)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> GRASSP Capabilities of Upper Extremity Test Grip and Pinch Force Modified Ashworth Scale 	October 2022	March 7, 2022	October 17, 2022	Seattle, Washington, USA
NCT05265377	MarsBionics	analyse the usability of the STELO modular exoskeleton in people with ABI and SCI	<ul style="list-style-type: none"> Age 18-85 years SCI level NS AIS NS 	Recruiting	20	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Single Group Assignment: Masking: None (Open Label)	Rehabilitation	General health	<ul style="list-style-type: none"> Heart rate Oxygen saturation Blood pressure Skin integrity Pain (Visual Analogic Scale) Fall prevalence Kinematic gait analysis Muscular activation Functional Ambulation Categories WISCI Timed Up and Go (TUG) 6MWT 10MWT Device malfunction User perception of the device (QUEST 2.0) Participant's satisfaction of the device 	September 2022	March 3, 2022	September 22, 2022	Madrid, Spain
NCT05262816	Second Affiliated Hospital of Wenzhou Medical University	The purpose of this study is to clarify the therapeutic effects of different acupoints on different types of bladder	<ul style="list-style-type: none"> Age 18-80 years SCI level suprasacral AIS NS voiding dysfunction 	Recruiting	34	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Single (Outcome Assessor)	Technology	Bladder health	<ul style="list-style-type: none"> Perianal surface electromyography Detrusor overactivity Bladder compliance Maximum cystometric capacity Maximum urinary flow rate Maximum detrusor pressure 	August 2021	March 2, 2022	March 2, 2022	Wenzhou, Zhejiang, China
NCT05255679	University of Alberta	Effect of early FES cycling on muscle wasting, pain or spasticity after spinal cord injury	<ul style="list-style-type: none"> Age 18-85 years SCI C1-L5 AIS A-C acute SCI 	Recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single (Participant)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> ISNCSCI quality of life PHQ-9 WISCI 	April 2021	February 24, 2022	February 24, 2022	Edmonton, Alberta, Canada
NCT05249595	North Carolina State University	Electromyography-ultrasound imaging-based technique to sense residual voluntary strength in ankle muscles for individuals with neuromuscular disorders	<ul style="list-style-type: none"> Age 18-64 years SCI level: Cervical, thoracic, or lumbar AIS C-D sub-acute or chronic SCI 	Recruiting	25	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> volitional effort kinematic reaction forces muscle activation 	February 2020	February 16, 2022	February 21, 2022	Raleigh, NC, USA
NCT05214378	VA Office of Research and Development	The purpose of this study is to determine if bladder emptying can be achieved using stimulation of the sacral roots at certain frequencies.	<ul style="list-style-type: none"> Age ≥ 18 years SCI C1-L5 Already have a sacral anterior root stimulation (SARS) device implanted 	Enrolling by invitation	8	Phase: Early Phase 1, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Bladder health	<ul style="list-style-type: none"> Urethral sphincter pressure in response to stimulation compared to pressure in absence of stimulation during bladder emptying 	July 2022	January 28, 2022	July 20, 2022	Cleveland, OH, USA
NCT05210166	University of Castilla-La Mancha	Transcutaneous Spinal Cord Stimulation Combined With Robot-assisted Therapy in Incomplete Spinal Cord Injury Patients.	<ul style="list-style-type: none"> Age 18-80 years AIS C-D SCI level C4-T11 SCI 2-6 months 	Recruiting	30	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double (Participant, Outcome Assessor)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 10 MWT 6 MinWT LEMS Modified Ashworth Test Contraction strength (dynamometry) Motor evoked potentials SCIM TUG WISCI II 	March 2021	January 27, 2022	January 27, 2022	Toledo, Spain

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
NCT05200091	Centre for Interdisciplinary Research in Rehabilitation of Greater Montreal	4-week mixed training paradigm consisting of explosive strength training and balance perturbation training efficacy on balance control during standing and locomotion, and to understand the mechanisms of neuroplasticity	<ul style="list-style-type: none"> Age ≥ 18 years SCI: C5-T12 AIS: C-D SCI ≥ 6 months be able to stand without support for 15 seconds 	Recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> EMG responses following balance perturbation Ankle muscle strength Transcranial magnetic stimulation (TMS) Modulation of the H reflex 	July 2018	January 20, 2022	January 20, 2022	Montréal, QC, Canada
NCT05196204	Centre for Interdisciplinary Research in Rehabilitation of Greater Montreal	Evaluate trunk task-oriented training (12 weeks) combined with function electrical stimulation (FES/T-TOT) efficacy on sitting balance and functional independence, and to understand the mechanisms of neuroplasticity that would improve functional independence following FES/T-TOT	<ul style="list-style-type: none"> Age 18-65 years SCI C6-T10 AIS A-D SCI ≥ 6 months be able to sit without support for 15 seconds NOT wear a corset 	Recruiting	45	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> Function in Sitting Test Modified Functional Reach Test (lateral and anterior trunk flexion) EMG activation pattern of electromyographic activity of muscles in the trunk Center of pressure excursion Modification of the excitability of the reticulospinal pathway using the acoustic startle response Modification of the excitability of the vestibulospinal pathway using galvanic vestibular stimulation 	May 2021	January 19, 2022	January 19, 2022	Montréal, QC, Canada
NCT05183152	University of Texas at Austin	Non-invasive Brain-computer Interfaces (BCI) for Control of Assistive Devices	<ul style="list-style-type: none"> Age 18-80 years 	Recruiting	40	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Arm/hand function	<ul style="list-style-type: none"> BCI command delivery accuracy fMRI activation for different imagined movements Stability and separability of Motor Imagery features Motor-evoked potential amplitude Electroencephalography functional connectivity 	June 2021	January 10, 2022	December 5, 2022	Austin, TX, USA
NCT05180227	VA Office of Research and Development	Targeted Transcutaneous Stimulation of the Spinal Cord to Restore Autonomic Cardiovascular Health in Veterans With Spinal Cord Injury	<ul style="list-style-type: none"> Age 21-70 years Traumatic SCI SCI: at or above T6 AIS: A,B or C SCI ≥ 1 year 	Recruiting	10	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Single Group Assignment Masking: Single (Participant)	Technology	General health	<ul style="list-style-type: none"> Electrode placement Amplitude of TSCS to normalize blood pressure Systolic blood pressure Concentration of Norepinephrine Velocity of Arterial stiffness Concentration of Renin Concentration of Aldosterone 	April 2022	January 6, 2022	April 26, 2022	West Orange, NJ, USA; Bronx, New York, USA
NCT05178056	University of Louisville	Spinal Cord Epidural Stimulation (scES) will be administered by a multi-electrode array (5-6-5 Spicity/TM electrode, MEDTRONIC, Minneapolis, MN, USA) previously implanted in the epidural space over the dorsum of the spinal cord. Additionally, respiratory training sessions are performed (eighty 45-minute sessions during 16 weeks)	<ul style="list-style-type: none"> Age ≥ 18 years AIS A SCI ≥ 24 months have at least a 15%-deficit in pulmonary function outcomes 	Recruiting	30	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Parallel Assignment, Masking: Single	Technology	General health	<ul style="list-style-type: none"> Maximum Inspiratory Pressure Maximum Expiratory Pressure Surface electromyography (sEMG) Magnitude Surface electromyography (sEMG) Similarity Index Forced Vital Capacity Forced Expiratory Volume in 1 second Baroreflex sensitivity Baroreflex Effectiveness Index 	December 2021	January 5, 2022	January 9, 2023	Louisville, KY, USA
NCT05176327	The University of Hong Kong	Assess the effects of exoskeleton training on neurogenic bowel disorders in spinal cord injury/ disease.	<ul style="list-style-type: none"> Age ≥ 18 years SCI C4-L3 AIS D SCI ≥ 12 months have a bowel opening via anal route or stoma 	Recruiting	10	Phase: Early Phase 1, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Bowel health	<ul style="list-style-type: none"> Neurogenic Bowel Dysfunction (NBD) score Defaecation time (DT) Frequency of bowel incontinence episodes 	January 2022	January 4, 2022	May 18, 2022	Hong Kong
NCT05167032	University of Minnesota	This is a mechanistic Phase I randomized pilot clinical trial in 16 adults with SCI/D. The investigators will compare the effects of Cognitive Multisensory Rehabilitation (CMR) vs. adaptive fitness on sensorimotor function.	<ul style="list-style-type: none"> Age 18-75 years 	Recruiting	16	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double (Investigator, Outcome Assessor)	Rehabilitation	Arm/Hand function	<ul style="list-style-type: none"> ISNCSCI Neuromuscular Recovery Scale (NRS) Spinal Cord Injury Functional Index/Assistive Technology (SCI-FI/AT) Pain Basic Set Sleep Quality Index Anxiety Patient Health Questionnaire (PHQ-9) World Health Organization Quality of Life Instruments (WHOQOL-BREF) Body Awareness Rating Questionnaire Kinesthetic and Visual Imagery Questionnaire (KVIQ) Moorong Self-Efficacy Scale (MSES) Tampa Scale for Fear of Re-Injury Patient-Specific Functional Scale Craig Handicap Assessment and Reporting Technique-Short Form (CHART-SF) 	June 2022	December 22, 2021	June 22, 2022	Minneapolis, Minnesota, USA
NCT05163639	Columbia University	Spinal cord associative plasticity (SCAP) is a combined cortical and spinal electrical stimulation technique developed to induce recovery of arm and hand function in spinal cord injury.	<ul style="list-style-type: none"> Age 18-80 years 	Recruiting	92	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Arm/Hand function	<ul style="list-style-type: none"> hand muscle response to brain stimulation during combined brain and spinal stimulation pinch force 	September 2021	December 20, 2021	June 3, 2022	New York, NY, USA
NCT05157282	VA Office of Research and Development	Examine the behavioral and physiological effects of TESS on upper-limb muscles after cervical SCI; and 2) Maximize the recovery of reaching and grasping potential by using tailored TESS in a task-specific manner with motor training.	<ul style="list-style-type: none"> Age 18-70years SCI > 1 year SCI level C8 and above 	Recruiting	86	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Single (Participant)	Technology	Arm/Hand function	<ul style="list-style-type: none"> GRASSP MEPs 	January 2023	December 14, 2021	January 27, 2023	Chicago, IL, USA; Hines, IL, USA
NCT05142943	University of Valencia	The objective of this study is to analyze the effectiveness of visual illusion therapies in combination with conventional exercises on the symptoms and signs related to different pathologies of the nervous and musculoskeletal system that affect the upper limb.	<ul style="list-style-type: none"> Age 18-99 years AIS C-E 	Recruiting	80	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double (Participant, Outcome Assessor)	Rehabilitation	Arm/Hand function	<ul style="list-style-type: none"> motor function and motor skills hand coordination Upper limb isometric force muscle activation Pain Muscle tone Spasticity Quality of life Independence (SCIM) 	July 2022	December 3, 2021	November 30, 2022	Valencia, Spain

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NCT05141487	VA Office of Research and Development	This project will demonstrate the feasibility of a new type of nerve stimulation-triggered sacral neuromodulation-to treat NDO in Veterans. A wireless bladder sensor will be inserted into the bladder to transmit a feedback signal enabling stimulation from a percutaneous lead.	<ul style="list-style-type: none"> Age >18 years SCI > 6 months tolerate lying prone > 1 hour 	Recruiting	NA	Phase: Not Applicable, Primary Purpose: NA, Intervention Model: NA, Masking: NA	Technology	Bladder Health	<ul style="list-style-type: none"> Time between voids during conditional SNM Detrusor contraction period during conditional SNM Voided urine volume during conditional SNM Urinary incontinence symptom F/U: 2 weeks	October 2022	December 2, 2021	November 16, 2022	Cleveland, OH, USA
NCT05128994	Battelle Memorial Institute	The objective of this study is to advance personalized, portable, and non-invasive hand-grasp neuro-orthoses that restore naturalistic grasp functions for those with tetraplegia due to spinal cord injury (SCI), designed around their needs and preferences.	<ul style="list-style-type: none"> Age ≥ 18 years ≥ 12 months post-SCI and medically/neurologically stable SCI level C1-C8 Unable to grasp objects independently with both hands (tetraplegia) 	Recruiting	8	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand function	<ul style="list-style-type: none"> Feasibility of FES sleeve as a functional orthosis for activities of daily living Quadriplegia Index of Function (QIF) System Usability Scale (SUS) GRASSP Psychosocial Impact of Assistive Devices Scale (PIADS) Electromyography (EMG) signal metrics Number of serious adverse events F/U: 10 weeks	April 2022	November 22, 2021	April 7, 2022	Columbus, OH, USA
NCT05115149	Skolkovo Institute of Science and Technology	The research will jointly use a prototype neurorehabilitation orthosis, in which a robotic device moves a paralyzed arm at the command of a non-invasive brain-computer interface to perform a game life-like task augmented using a virtual-reality display, as well as an electrical stimulation device that activates the spinal cord and/or muscles of the paralyzed arm.	<ul style="list-style-type: none"> Age 18-70 years SCI: C3-C7 AIS: A or B After a first occurred acute cerebrovascular accident or in the recovery period after injury of the cervical and upper thoracic spinal cord. Early or late rehabilitation period of acute cerebrovascular accident by the type of ischemic stroke or the consequences of SCI in the late recovery period 	Recruiting	60	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double (Participant, Outcome Assessor)	Technology	Arm/Hand function	<ul style="list-style-type: none"> Fugl-Meyer scale for the upper limb Action Research Arm Test Accuracy of BCI tasks Rivermead Mobility Index Ashworth Spasticity Scale The Capabilities of Upper Extremity Test Spinal Cord Independence Measure III American Spinal Injury Association Impairment Scale 36-Item Short Form Survey F/U: 6 weeks	October 2021	November 10, 2021	November 10, 2021	Multicenter, Russian Federation
NCT05111093	Ecole Polytechnique Fédérale de Lausanne	The HemON study aims to evaluate the safety and preliminary efficacy of ARC-IM Therapy (Epidural Electrical Stimulation) to improve hemodynamic management and trunk control in people with sub-acute or chronic spinal cord injury (>= 1 month post injury) between C3 and T6 inclusive, who suffer from orthostatic hypotension.	<ul style="list-style-type: none"> Age > 18 years SCI C3-T6 (inclusive) SCI ≥ 1month Confirmed orthostatic hypotension 	Recruiting	16	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General health	<ul style="list-style-type: none"> Occurrence of Serious Adverse Events and Adverse Events Orthostatic head-up tilt test Trunk stability measurement Modified Ashworth Scale (MAS) SCIM III F/U: 26 months	November 2021	November 8, 2021	October 4, 2022	Lausanne, Vaud, Switzerland
NCT05094752	Assistance Publique - Hôpitaux de Paris	Effect of tendon vibrations with the Vibramov™ system on spasticity and sensorimotor recovery in tetraplegic and high paraplegic patients.	<ul style="list-style-type: none"> affiliated to the French social security system Age > 18 years traumatic SCI at or above T6 	Recruiting	40	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double (Participant, Investigator)	Technology	Spasticity	<ul style="list-style-type: none"> modified Asworth scale (MAS) Muscle reaction in the modified Tardieu scale (MTS) Spinal cord assessment tool for spastic reflexes Pain ISNCSCI SCIM Montreal cognitive assessment (MoCA) F/U: 1 year	October 2012	October 26, 2021	October 26, 2021	Le Kremlin-Bicêtre, France
NCT05094362	Medical University of South Carolina	Validate the capacity of a reflex training system to change the size of the targeted reflex.	<ul style="list-style-type: none"> Age >= 18 years SCI level above T11 SCI > 1 year ability to ambulate at least 10 meters with/without assistive device 	Not yet recruiting	25	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 10MWT 6MWT WISCI MAS Berg Balance Scale FIM SCI-QOL muscle strength Tardieu Scale F/U: 3 months	March 2023	October 26, 2021	May 12, 2022	Charleston, South Carolina, USA
NCT05091463	University of Louisville	Multi-modal training combining activity-based locomotor training and transcutaneous spinal stimulation (ABLT+scTS) to improve sitting posture and trunk control in children with a chronic spinal cord injury.	<ul style="list-style-type: none"> Age 3-12 years SCI > 1year SCI level T10 or above moderate to severe trunk control NO baclofen pump 	Enrolling by invitation	12	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Trunk control Modified Function Reach Center of Pressure displacement Angular Excursions of the trunk F/U: 3 months	August 2021	October 25, 2021	January 10, 2023	Louisville, KY, USA
NCT05095454	Kristin Zhao, PhD	A study comparing short-term delivery of epidural spinal stimulation versus transcutaneous spinal stimulation.	<ul style="list-style-type: none"> Age >=22 years traumatic SCI at or above T10 AIS A-D SCI >= 1 year 	Recruiting	8	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Kinematics EMG Foot pressure SEPs MEPs ISNCSCI Bowel Function Bladder function Sexual function Spasticity trunk stability F/U: 5 months	March 2022	October 17, 2021	October 4, 2022	Rochester, MIN, USA
NCT05071885	University of Manitoba	The purpose of this research proposal is the further development and validation of a multipurpose plug-n-play rehab gaming system for use in community centers, in particular, at First Step Wellness center	<ul style="list-style-type: none"> Age 20-70 years 6months < SCI > 2 years Actively extend at least ten degrees at the metacarpophalangeal and interphalangeal joints, extend ten degrees at the wrist and had at least 30 degrees of active flexion, extension of the elbow and shoulder 	Not yet recruiting	64	Phase: Early Phase 1, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Rehabilitation	Arm/Hand function	<ul style="list-style-type: none"> Computer game-based Upper Extremity (CUE) assessment of manual dexterity Wolf motor function test (WMFT) F/U: 10 weeks	October 2021	October 8, 2021	October 8, 2021	Manitoba, Canada

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NCT05044923	Aaron Phillips	Targeted Epidural Spinal Stimulation. Epidural implantation of two lead electrodes over the dorsal aspect of the spinal cord through two laminotomies. Two implantable pulse generators will be connected to the lead electrodes and implanted in the upper buttocks of the participant.	<ul style="list-style-type: none"> Age 18-70y Level C3-T6 AIS A-B SCI > 1y Confirmed orthostatic hypotension and autonomic dysreflexia 	Recruiting	8	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Occurrence of Adverse Events and Serious Adverse Events Orthostatic head-up tilt test Daily stimulation log Echocardiogram (Ejection Fraction, Strain) Vascular ultrasound Autonomic Dysfunction Following Spinal Cord Injury (ADFSCI) ISNCSCI Respiratory function evaluation (Volume, Flow) Quality of life questionnaire (WHOQOL-BREF) F/U: 9 months	December 2021	September 16, 2021	December 21, 2021	Calgary, Alberta, Canada
NCT05035823	Synchron Medical, Inc	COMMAND Early Feasibility Study (EFS)	<ul style="list-style-type: none"> Age 21-75 SCI level C1-C8 severe quadriplegia Have a study partner No allergy to nickel NO history of pulmonary embolism No psychiatric or psychological disorder 	Recruiting	6	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General health	<ul style="list-style-type: none"> Serious AEs F/U 12 months	April 2022	September 5, 2021	December 9, 2022	Pittsburgh, PA, USA
NCT04994886	Jocelyne Bloch	Targeted Epidural Spinal Stimulation to manage blood pressure instability	<ul style="list-style-type: none"> Age 18 to 70 yrs SCI level C3 and T6 AIS A or B SCI > 1 year confirmed orthostatic hypotension and autonomic dysreflexia NO mental illness 	Recruiting	8	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Adverse Events and Serious Adverse Events ISNCSCI Modified Ashworth Scale (MAS) Trunk stability Respiratory function evaluation Orthostatic head-up tilt test Autonomic Dysfunction Following Spinal cord injury (ADFSCI) Quality of Life questionnaire WHOQOL-BREF Daily Stimulation Log (DSL) F/U: 7 months	June 2021	August 6, 2021	May 26, 2022	Lausanne, Vaud, Switzerland
NCT04977284	University of British Columbia	Non-surgical spinal cord stimulation (DSTR, Digitimer) will be applied and electrical activity of muscles will be recorded.	<ul style="list-style-type: none"> Age: 18-65 years Resident of British Columbia SCI > 1y Traumatic SCI Level above T10 AIS A or B NOT be ventilator dependent NO recent treatment with OnabotulinumtoxinA into the detrusor muscle (relative contraindication). NO Intrathecal baclofen pump 	Not yet recruiting	5	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Activation of spinal circuits (presence of motor evoked potentials or free run surface electromyography) Safety and efficacy during Urodynamic Testing (UDS) Safety and efficacy during Anorectal Manometry (ARM) F/U: 4 weeks	September 2021	July 26, 2021	July 26, 2021	No known locations
NCT04977037	The University of Texas Health Science Center, Houston	Transcranial direct current stimulation (tDCS) electrodes will be placed over the primary motor cortex and delivered 20 minutes. Immediately after stimulation ceases, participants will continue with unilateral repetitive arm and finger exercises. Exercise difficulty will gradually be increased and adjusted per participant's tolerance.	<ul style="list-style-type: none"> Age: 18-70years AIS B-D SCI > 6m Level C1-C8 NO contradiction to tDCS NO chronic use of neuroactive medication (e.g., neurostimulants, anticonvulsants, or antidepressants); 	Recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double	Rehabilitation	Arm/Hand Function	<ul style="list-style-type: none"> Graded Redefined Assessment of Strength, Sensibility and Prehension (GRASSP) Adherence with the therapy (attended sessions; drop-outs) Feasibility of home intervention Grip Strength Spinal Cord Injury Independence Measure (SCIM III)-Self Care Incidence of adverse events F/U: 1 month	July 2021	July 26, 2021	July 26, 2021	Houston, TX, USA
NCT04973852	The University of Texas Health Science Center, Houston	High Intensity Training Using Overground Exoskeletons (Ekso). The locomotor training will be performed with a focus on cardiovascular training parameters	<ul style="list-style-type: none"> Age > 18years AIS C or D NO uncontrolled spasticity NO use of mechanical ventilation for respiratory support 	Recruiting	20	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Percentage of Heart Rate Reserve Achieved During HIT Gait Training Session Self Selected Gait Speed as Assessed by the 10 Meter Walk Test (10MWT) Fast Gait Speed as Assessed by the 10 Meter Walk Test (10MWT) Walking Endurance as Assessed by the 6 Minute Walk Test (6MWT) Seated Dynamic Reach as Assessed by the Modified Functional Reach Test Spatial-Temporal Gait Parameters as assessed by the GAITRite pressure map (step length, stride length, single support, double support, swing time, stance time) Metabolic Expenditure during 10MWT and 6MWT, as Assessed by Oxygen Consumption F/U: 12 weeks	August 2021	July 21, 2021	January 21, 2022	Houston, TX, USA
NCT04969042	Luming Li	Implanted of a spinal cord stimulation device (Pins Medical G122 RS) which could give closed-loop functional stimulation in the epidural space	<ul style="list-style-type: none"> Age 18-70 years AIS A, B, C or D Level T10 and above SCI > 3 months Cognitive impairment 	Recruiting	10	Phase: Phase 1, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> WISCI III 10-meter Walking Test Weight Bearing Capacity F/U: 8 months	August 2021	July 20, 2021	July 20, 2021	Beijing, China
NCT04965727	Jocelyne Bloch	Deep brain stimulation (DBS) surgery: Implanting the lead electrodes (Medtronic SenSight Directional Lead) in the right and left lateral hypothalamus through craniotomy and an implantable pulse generator (Medtronic Model B35200 Percept™ PC) in the upper part of the pectoralis major (under the clavicle).	<ul style="list-style-type: none"> Age: 18-65 years AIS C or D Traumatic SCI SCI > 12 months NO cognitive/brain damage NO epilepsy NO use of an intrathecal baclofen pump NO implanted cardiac device such as pacemaker or defibrillator 	Recruiting	3	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Occurrence of all SAEs and AEs Lower Extremity Motor Strength Walking Index for Spinal Cord Injury (WISCI II) Walking speed (10MWT/6MWT) F/U: 6 months	June 2021	July 16, 2021	October 28, 2021	Lausanne, Switzerland

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NCT04921592	Kessler Foundation	transcutaneous stimulation of the upper extremities	<ul style="list-style-type: none"> Age ≥ 18 years SCI ≥ 6 months cervical neurologic level of injury NOT be ventilator dependent. NO history of fractures history of illicit drug abuse history of cardiac, respiratory, bladder, renal or other medical disorder unrelated to spinal cord injury NO implanted pump (i.e., baclofen pump, pain pump, etc) 	Not yet recruiting	36	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Neuro recovery scale F/U 6 months	June 2021	July 10, 2021	June 14, 2021	No known locations
NCT04914975	Swiss Paraplegic Centre Nottwil	Neuromuscular electrical stimulation will be conducted at home or at the SPC for half an hour about 30 minutes before the usual bowel emptying time. Four adhesive electrodes are attached to the abdominal wall for the neuromuscular electrical stimulation. The abdominal muscles are stimulated in such a way that activation occurs, i.e. the muscle alternately contracts and relaxes again. The stimulation sessions will be documented with a defined protocol. In addition, the stool consistency is rated according to the Bristol Stool Form Scale.	<ul style="list-style-type: none"> Traumatic and non-traumatic SCI; > 1 year SCI C2 - L5 AIS A/B/C/D Age ≥ 18 years NO bladder stimulator NO Autonomic dysreflexia by application of ES of the abdominal wall Pregnancy: test in women of childbearing age (15 - 49 years) NO Opioid use NO inflammatory bowel disease NO cancerous tissue in abdominal region NO inability to follow the study, e.g. mental-health problems, language problems, dementia etc. 	Recruiting	20	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Bowel Health	<ul style="list-style-type: none"> Neurogenic Bowel Dysfunction Score Corn Test Bristol stool form scale Qualiveen Short Form ISAFSCI F/U 24 weeks	July 2021	July 7, 2021	February 17, 2022	Nottwil, Switzerland
NCT04910204	University Health Network, Toronto	FES Therapy combined with task-specific training (FEST-TST). Description: The FEST-TST protocol consists of a 1-hour session, 3 to 5 days a week, for up to 12 weeks (40 sessions total) in addition to conventional occupational and physical therapies according to the standard of care.	<ul style="list-style-type: none"> Age > 18 years subacute SCI (<3 months) AIS C or D cervical SCI 	Not yet recruiting	18	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> SCIM UEMS, UESS GRASSP needle EMG Nerve conduction studies SSEPs MEPs fMRI laboratory assessments: BDNF, NfT-3 	February 2022	July 2, 2021	June 2, 2021	Toronto, Canada
NCT04902482	University College, London	The iCycle Mark 3 is a cycle ergometer, designed to be used by people with SCI while they are seated in their own wheelchairs. The purpose of iCycle is to stimulate the leg muscles in the correct phase for cycling while motivating the person with a Virtual Reality cycling event, perhaps a race, to try to use their muscles.	<ul style="list-style-type: none"> Age ≥ 18 years SCI above T12 Incomplete SCI >12 months post injury Marginal walker defined as able to rise from a chair, stand for 10 seconds and walk >10 steps without human help (but may use an aid including parallel bars). NO cardiac pacemaker NO pressure sores/skin problems NO pregnancy NO implanted metal work at electrode site (< 3/12) 	Recruiting	6	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> ISNCSCI motor score Trunk Impairment Scale (TIS) score Walking Index for Spinal Cord Injury (WISCI II) score 6-minute walk test (6MWT) distance Threshold of evoked responses to Transcranial Magnetic Stimulation (TMS) Audio/transcribed feedback from participants from semi-structured interview F/U: 60 min	January 2022	May 26, 2021	August 12, 2022	Stanmore, Middlesex, UK
NCT04894734	Nandan Lad, M.D., Ph.D.	Epidural electrical stimulation (EES), also known as spinal cord stimulation (SCS), is a common FDA-approved therapy for chronic neuropathic pain of trunk and limb.	<ul style="list-style-type: none"> Age between 18-80 years Traumatic SCI Level T1-T12 Chronic pain (i.e., Pain >3 for > 3 months, on > 50% of the days) 	Recruiting	30	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Triple	Technology	General Health	<ul style="list-style-type: none"> Multidimensional Pain Inventory (MPI)-SCI average activity score Motor recovery as measured by EMG Motor recovery as measured by dynamometry Pain as measured by 10-point Numeric Rating Scales (NRS) Quality of Life (QOL) as measured by the Patient-Reported Outcomes Measurement Information System (PROMIS) 29 Number of prescriptions written as measured by Electronic Health Record abstraction Overall improvement as measured by Guy/Farrar Patient Global Impression of Change (PGIC) scale Motor recovery as measured by the Total American Spinal Injury Association (ASIA) motor score Motor recovery as measured by the ASIA impairment grades Independence of activities of daily living (ADLs) as measured by the Spinal Cord Independence Measure (SCIM) survey Limb movement as measured by the Ashworth spasticity scale Bladder control using urodynamics Motor recovery as measured by Transcranial Magnetic Stimulation Motor Evoked Potentials (TMS MEPs) Sensory recovery as measured by Somatosensory Evoked Potentials (SSEPs) F/U: 9 months	September 2022	May 20, 2021	November 25, 2022	Durham, NC, USA

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
NCT04889092	VA Office of Research and Development	Exercise training will consist of blood flow restriction exercise. Specifically blood pressure cuff will be wrapped around the most proximal portion of the thigh and inflated to a pressure that is 80% of the pressure required to completely occlude femoral blood flow. With the cuff inflated the subject will perform a series of knee extension/flexion exercise protocol. This consists of 30 reps, 15 reps, 15 reps and 15 reps all separated by 1 minute of recovery. This will be performed 20 times over 9 weeks. Comparison to traditional resistance exercise	<ul style="list-style-type: none"> Age 18-70 years AIS C-D SCI > 1 year Level between C3-L1 NO pregnancy NOT ventilator-dependent NO chronic inflammatory disease 	Recruiting	22	Phase: Not Applicable, Primary Purpose: Supportive Care, Intervention Model: Crossover Assignment, Masking: Single	Rehabilitation	General Health	<ul style="list-style-type: none"> Muscle cross sectional area (CTscan) Muscle strength (biodes dynamometer) Vascular endothelial function (flow mediated dilation) Muscle fatigue resistance (% decrease in maximal voluntary torque following fatigue protocol) Muscle volume (CTscan) Thrombin / antithrombin complex (blood samples) Prothrombin fragment 1.2 (marker of coagulation) Vessel stiffness in the lower limbs (pulse wave velocity) Interleukin (IL)-Beta (marker of inflammation) Neutrophil-platelet aggregates Hypoxia-inducible factor 1-alpha (HIF-1) Vascular endothelial growth factor (VEGF) Reactive hyperemia (NIRS and Doppler/ultrasound imaging) Interferon (marker for inflammation) C-reactive protein (CRP, marker for inflammation) Interleukin 6 (marker for inflammation) NETosis F/U: 56 days	July 2021	May 17, 2021	August 16, 2022	Cleveland, OH, USA
NCT04883463	University of California, Los Angeles	Epidural electrical stimulation implant weekly sessions for 21 months.	<ul style="list-style-type: none"> Age 18-75 years SCI > 1 year Non-progressive SCI Level C2-C7 AIS A-B Ventilator dependency NO history of severe autonomic dysreflexia 	Recruiting	15	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Incidence of Treatment-Emergent Adverse Events as assessed by CTCAE v4 ISNCSCI motor and sensory function Resting Spontaneous Respiratory Activity Blood Pressure (mm/Hg) Evaluating Pulmonary Function Throughout the Duration of the Study F/U: 2.5 years	September 2021	May 12, 2021	January 26, 2023	Los Angeles, CA, USA
NCT04881565	University Health Network, Toronto	Reactive balance training. During each one-hour session, participants will experience 40-50 perturbations during standing and/or walking activities. The perturbations will be applied in any direction. To create a perturbation, the researcher will apply unexpected pushes or pulls to a safety harness at waist level. The perturbation will be sufficient in magnitude to elicit a stepping response from the participant. Throughout the session, participants will complete challenging balance tasks, customized to their ability level. Balance tasks will be organized into five categories: stable, quasi-mobile, mobile, unpredictable and participant-selected.	<ul style="list-style-type: none"> Age ≥ 18 years Traumatic SCI AIS C-D Level ≥ T12 SCI > 1 year Able to stand for >30 seconds without upper limb support or assistance No contraindications to functional electrical stimulation (FES) 	Recruiting	22	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> Berg Balance Scale mini-Balance Evaluation Systems Test Activities-specific Balance Confidence Scale Falls Efficacy Scale - International Isometric strength of lower extremity muscle groups Proprioception of the ankle joints Lean-and-Release Test Tracking falls F/U: 6 months	September 2021	May 11, 2021	November 2, 2022	Toronto, Canada
NCT04879862	University of Louisville	Epidural stimulation + stand training. Participants will be encouraged to stand for as long as possible throughout the training session, with the goal to stand for 60 minutes with the least amount of assistance. Epidural stimulation + step training. Participants will step at various body weight load and speed. Participants will take a break and rest at any time they feel the need to during the session. Epidural stimulation + bladder capacity training. Daily training for capacity in a supervised on-site lab setting. Epidural stimulation + bladder voiding efficiency training. Voiding without catheterization will be attempted.	<ul style="list-style-type: none"> Age ≥ 18 years Non-progressive SCI SCI > 1 year Level T1-T10 Inability to stand and step independently Unable to voluntarily move all individual joints of the legs NO current anti-spasticity medication NOT have received botox injections in the prior six months Bladder dysfunction as a result of SCI NOT ventilator dependent 	Recruiting	16	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Ambulation Bladder storage/voiding F/U: 3 months	April 2022	May 10, 2021	August 15, 2022	Louisville, KY, USA
NCT04781621	Baylor Research Institute	Participants will wear an exoskeleton suit and receive robotic gait training with a physical therapist for 90 minutes each week. Robot gait training will include standing and walking activities while wearing a robot suit.	<ul style="list-style-type: none"> Age 16 - 70 years incomplete SCI Acute/Subacute SCI NO Traumatic Brain Injury NO degenerative diagnoses 	Recruiting	128	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> Walking Index for Spinal Cord Injury - II (WISCI-II) Gait speed via 10-Meter Walk Test (10MWT) Spinal Cord Independence Measure (SCIM) Numerical Pain Rating Scale (NPRS) Fatigue Severity Scale (FSS) Penn Spasm Frequency Scale (PSFS) Patient Health Questionnaire - 9 (PHQ-9) General Anxiety Disorder (GAD-7) International Spinal Cord Injury Quality of Life Basic Data Set Heart Rate (HR) Ratings of Perceived Exertion (RPE) Number of Steps Patient Perceptual Survey F/U within one week before discharge	April 2021	May 4, 2021	July 2, 2021	No known locations

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
NCT04858178	Spaulding Rehabilitation Hospital	This study looks to characterize autonomic nervous system dysfunction after spinal cord injury and identify the potential role that transcutaneous spinal cord stimulation may play at altering neuroregulation. experiments will utilize multiple parallel recordings to identify how the autonomic nervous system is able to inhibit and activate sympathetic signals.	<ul style="list-style-type: none"> • Age 18-30 years • traumatic SCI • SCI >=1 year • AIS A • SCI level T1-T6 • NO history of cardiovascular disease, hypertension, neurologic disorders, diabetes • NO pregnancy 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Diagnostic, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> • Valsalva Maneuver Phase II • Muscle sympathetic nerve activity • Heart rate • Blood pressure • Galvanic skin response • Autonomic Dysfunction Following Spinal Cord Injury questionnaire • Composite Autonomic Symptom Score F/U 3 months	February 2022	April 26, 2021	October 3, 2022	Charlestown, MA, USA
NCT04855812	Kessler Foundation	To evaluate the usefulness of an upper extremity assistive device, called (MyoPro) in improving upper extremity activities in people with incomplete spinal cord injury (iSCI)	<ul style="list-style-type: none"> • Age 18-80 years • AIS C-D • SCI >= 1 year • SCI level C1-C8 • NO history of other neurologic disorder or recurrent epilepsy, seizure or convulsion • NO history of treated ringing in the ears known as tinnitus or severe hearing problems 	Recruiting	24	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Crossover Assignment, Masking: Quadruple	Rehabilitation	Arm/Hand Function	<ul style="list-style-type: none"> • Range of motion of hands and arms via double sticky tape. • Muscle strength measurement on the participant's skin via double stick tape. • GRASSP • Spasticity measurement (MAS) • CUE-Q F/U 10 weeks	October 2019	April 22, 2021	May 24, 2022	East Hanover, NJ, USA
NCT04849676	NHS Greater Glasgow and Clyde	Neurofeedback is a neuromodulatory intervention that does not require applying external stimuli. It relies on displaying participants' brain activity in real time, with the aim of modifying it. Electrical activity (EEG) will be recorded from participants and displayed in real time on a computer screen, in the form of three bars. They will be asked to increase the middle bar, while keeping the side bars low. The intervention requires training to achieve voluntary control of the targeted brain activity.	<ul style="list-style-type: none"> • Aged > 18 years • AIS C-D • SCI <= 6 months and SCI level C3-C7 • SCI <= 6 months and SCI level T1-L1 • SCI >= 1 year and SCI level C3-C7 and live inside Greater Glasgow and Clyde area • SCI >= 1 year and SCI level T1-L1 and live inside Greater Glasgow and Clyde area • NO history of epilepsy 	Recruiting	20	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Spasticity	<ul style="list-style-type: none"> • Modified Ashworth Scale • Functional improvements • SCIM F/U 4 weeks	August 2021	April 19, 2021	April 18, 2022	Glasgow, UK
NCT04821635	UGECAM Rhône-Alpes	Training on a rower with solicitation of the electrostimulated lower limbs and upper limbs	<ul style="list-style-type: none"> • Paraplegia with traumatic origin • AIS A-B • SCI >= 12 months • NO other associated neurological pathologies (stroke, peripheral neuropathy, myopathy, head trauma, ...) 	Recruiting	35	Phase: Not Applicable, Primary Purpose: Prevention, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Rehabilitation	General Health	<ul style="list-style-type: none"> • VO2 peak at 6 months (L/min) • Ashworth scale • osteotendinous reflexes • Goal Attainment Scaling (GAS) • AIS F/U 9 months	April 2021	March 29, 2021	March 29, 2021	Saint-Didier-aux-Monts, France
NCT04809987	University of Valencia	Virtual Gait vs. Physical Exercise	<ul style="list-style-type: none"> • AIS C, D or E 	Recruiting	80	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> • Gait: 10 meters Walking Test • Functionality: FallSkip • Strength: Load Cell • Spasticity: MyotonPRO Secondary • Neuropathic Pain: Brief Pain Inventory • Muscle Activation: EMG F/U 10 minutes	December 2020	March 22, 2021	March 2, 2022	Valencia, Spain
NCT04807764	City University of New York	40 daily sessions of 30 minutes of non-invasive high frequency (e.g. 30 Hz) transcutaneous transspinal stimulation during standing or lying followed by 30 minutes of assisted stepping robotic gait training. Before and after training standardized clinical and neurophysiological tests will be used to assess recovery of sensorimotor function.	<ul style="list-style-type: none"> • Age 18-70 years • AIS C-D • SCI level >T10 • SCI > 6 months • NO peripheral neuropathy • NO degenerative neurological disorders of the spine or spinal cord • NO cochlear implants, pacemaker, implanted infusion device, and/or implanted stimulators of any type • NO history of seizures 	Recruiting	45	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> • Plasticity of spinal neuronal networks (soleus H-reflex following posterior tibial and common peroneal nerves stimulation) • Plasticity of corticospinal networks (recording responses to single-pulse transcranial magnetic stimulation; TMS) • Ambulatory function (2-min walk and 10-meter timed test) • Balance (BESTest clinical assessments) • Autonomic function (bowel, bladder, and sexual function) F/U 4 years	August 2021	March 19, 2021	May 31, 2022	Staten Island, NY, USA
NCT04798378	Thomas Jefferson University	Upper extremity orthosis and functional stimulation system (Neurosleeves) for restoration of independent arm function	<ul style="list-style-type: none"> • Age >= 4 years • weakness in one or both arms such that wrist flexion and wrist extension are 3/5 on the Manual Muscle Testing Scale • SCI >= 6 months • fluent in English • NO visual impairment such that following visually-guided instructions would be challenging even with ordinary corrective lenses • NO untreated psychiatric or neurologic disturbances • NO implanted medical device in the body (such as cardiac pacemaker, implanted defibrillator, metallic device) • NO history of seizure or epilepsy • history of alcohol or other substance use 	Recruiting	20	Phase: Not Applicable, Primary Purpose: Supportive Care, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> • Canadian Occupational Performance Measure (COPM) score • Action Research Arm Test (ARAT) • Motricity Index score • ABILHAND-Kids questionnaire score • Box and Blocks score F/U 8 weeks	April 2020	March 15, 2021	December 5, 2022	Philadelphia, PA, USA Wilmington, DW, USA
NCT04782947	United States Department of Defense	Compare the impact of Exoskeletal assisted walking (EAW)+Epidural stimulation (ES) following improving lower extremity muscle quality compared to those who will only undergo EAW+ ES without conducting resistance training on motor recovery, cardio-metabolic health and bladder control in persons with complete SCI.	<ul style="list-style-type: none"> • Age 18-60 years • SCI >= T10 • AIS A-B 	Recruiting	20	Phase:Phase 2/3, Primary Purpose: Health Services Research, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> • 10MWT • F/U: 12 months 	February 2021	March 2, 2021	February 3, 2022	Richmond, VA, USA

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NCT04780470	University of Helsinki	6 weeks of technological-assisted upper extremity rehabilitation	<ul style="list-style-type: none"> • AIS C-D • SCI level C2-C8 • SCI since 1 - 5 years • ability to sit • motivation and ability to participate in intensive rehabilitation 3 times per week for 6 week period. • NO memory disorder 	Enrolling by invitation	60	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Single	Technology	Arm/Hand Function	<ul style="list-style-type: none"> • Action Research Arm Test (ARAT) scores • Goal Attainment Scaling (GAS) scores • Upper Extremity Motor Scores (ASIA-UEMS) • Grip strength • Pinch strength • Upper extremity active range of motion • SCIM-SR scores F/U 6 months	April 2021	February 18, 2021	November 9, 2022	Helsinki, Finland
NCT04759976	University of Bern	Robotic motor training: performing motor tasks with upper-limb robotic devices.	<ul style="list-style-type: none"> • Age ≥18 years • NO serious medical or psychiatric disorder 	Recruiting	250	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> • Change in kinematic performance assessed by the robot • Change in kinetic performance assessed by the robot • Spatial analysis of changes in evoked potentials • Change in virtual reality (VR) embodiment • Spatial analysis of changes in Task-Based Brain Connectivity • Change in Motivation as assessed by Intrinsic Motivation Inventory (IMI) • Change in Cognitive Load as assessed by National Aeronautics and Space Administration (NASA) (Raw) Task Load Index • System Usability as assessed by System Usability Scale (SUS) F/U 1-2 days after the training	January 2019	February 18, 2021	May 26, 2022	Bern, Switzerland
NCT04755699	Northwell Health	Transcutaneous Electrical Stimulation: administration of various electrical pulses being delivered to muscles and/or the spinal column with an investigational neurostimulator to evoke various limb movements in order to improve functional movement.	<ul style="list-style-type: none"> • Age 18 - 75 yrs • limited ability or no ability to use at least one hand • SCI > 1 year • History of epilepsy • Chronically-implanted electronic medical device (e.g. deep brain stimulator, epidural stimulator, cardiac pacemaker, vagus nerve stimulator, or other) • Abnormalities of the arms/hands, legs/feet, or spinal column that would prevent electrical stimulation • Ventilator dependence • History of serious mood or thought disorder • Pregnant women • Prisoners 	Recruiting	28	Phase: Early Phase 1, Primary Purpose: Device Feasibility, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> • Device Feasibility • Muscle Activation of the Targeted Limbs F/U 12 months	December 2020	February 16, 2021	December 29, 2022	Manhasset, NY, USA
NCT04736849	Peter J. Grah	Percutaneous epidural and dorsal root stimulation	<ul style="list-style-type: none"> • Age > 22years • SCI level T10 and higher • AIS A-D • SCI > 1 year • NO history of frequent and/or severe autonomic dysreflexia • NO history of seizure disorder 	Not yet recruiting	32	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> • Kinematics • Electromyography/Electroneurography • Foot pressure • Somatosensory evoked potentials • Transcranial magnetic stimulation motor evoked potentials • AIS • Neurogenic Bowel Dysfunction Score. • Neurogenic Bladder Symptom Score • Sexual Health Inventory for Men (SHIM) / International Index for Erectile Function (IIEF) • Female Sexual Function Index (FSFI) • Spinal Cord Injury Spasticity Evaluation Tool (SCI-SET) • User Experience Questionnaire (UEQ) • WISCI II • Modified 6 Minute Walk Test • Modified Timed Up and Go test F/U 4 weeks	April 2021	February 3, 2021	October 5, 2022	No known locations
NCT04735887	University of Stellenbosch	The SADL-eM includes three elements essential to the intervention, namely: knowledge, skills, and advice. The manual includes 92 A5 pages with six detailed sections: an Introduction and five chapters: (1) Rehabilitation team, (2) Activities of Daily Living, (3) Assistive devices, (4) Home environment adaptation, and (5) Knowledge guide. The SADL-eM uses text and illustrative pictures that are carefully selected for contextual relevance. The manual is simple, easy, and suitable for people with a non-medical background.	<ul style="list-style-type: none"> • AIS A, B, and C • Age 18 - 65 yrs • SCI ≤ 6 months. • Sufficient comprehension (read/write) of the Arabic language. • communication and/or cognitive disorders such as global aphasia and memory deficit. • disturbed level of awareness such as coma or lethargy. • other cause(s) of disability in addition to SCI such as stroke or amputation. • progressive disease or a psychiatric condition that would interfere with study • mechanical ventilator dependency 	Not yet recruiting	132	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	General Health	<ul style="list-style-type: none"> • SCIM-SR • SCIM-III F/U 6 weeks	February 2021	February 3, 2021	February 3, 2021	Gaza Strip

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NCT04727866	Bronx VA Medical Center	Transcutaneous spinal direct current stimulation (tsDCS) - coronal, 20 minutes of tsDCS will be delivered at 66% of maximum tolerated intensity with cathode over C5-C7 transverse process on target side, anode over C5-C7 transverse process on non-target side. Also over -T1-T4 posteriorly, anode over -C5-T1 anteriorly, and over -C3-C5 posteriorly, anode over -C5-T1 anteriorly	<ul style="list-style-type: none"> Age 18-75 yrs SCI > 12 months SCI Level C1-C8 Score of 2, 3, or 4 (out of 5) on manual muscle testing of elbow flexion, wrist extension, wrist flexion, finger extension, finger flexion, or finger abduction in left or right hand NO multiple spinal cord lesions NO history of seizures; NO ventilator dependency or patent tracheostomy site NO History of stroke, brain tumor, or brain abscess NO History of implanted brain/spine/nerve stimulators, aneurysm clips, ferromagnetic metallic implants, or cardiac pacemaker/defibrillator 	Recruiting	20	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Intervention Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Motor evoked potential (MEP) amplitudes H-reflex amplitudes Muscle dynamometry Intracortical inhibition and facilitation <p>F/U immediately after procedure</p>	January 2021	January 27, 2021	March 17, 2022	The Bronx, NY, USA
NCT04726059	University of British Columbia	Therapeutic Transcutaneous Spinal Cord Stimulation (TSCS) during Activity-Based Therapy (ABT) using the Lokomat exoskeleton (Hocom). All participants (both arms) will train 3 times per week for 12 weeks with a target to reach 20 minutes of balance training and 30 minutes of gait training in each session.	<ul style="list-style-type: none"> Resident of British Columbia, Canada with active provincial medical services plan Age 18-60 yrs Chronic traumatic SCI (non-progressive, with complete motor paralysis) SCI Level at or above T6 SCI > 1 year AIS A or B NO ventilator dependency NO intrathecal baclofen pump NO implanted metal in skull or pacemakers, stimulators, medication in the trunk NO fractures, metal implants, recurring headaches, pressure sores, epilepsy/seizure NO severe anemia (Hgb<8 g/dl) or hypovolemia 	Recruiting	12	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Quadruple	Technology	Standing/Walking Mobility	<ul style="list-style-type: none"> Corticospinal excitability (motor evoked potentials) Attempted voluntary contraction Seated and standing balance control Blood pressure variability Cardiac outcomes Dysautonomia Autonomic dysreflexia Bladder function based on the neurogenic bladder symptom score Bladder function based on the incontinence - quality of life questionnaire Bowel function based on the neurogenic bowel dysfunction score Sexual function based on the international index of erectile function questionnaire Sexual function based on the female sexual function index Quality of life based on the short form (SF-36) health survey Quality of life based on fatigue severity scale Quality of life based on spinal cord injury; spasticity evaluation tool <p>F/U 14 weeks</p>	July 2022	January 27, 2021	November 28, 2022	Vanvouver, BC, Canada
NCT04726891	University of Alabama at Birmingham	Movement-2-Music + Social Networking Support: 12-week pilot study of the SMART-HEALTH intervention in 30 individuals with SCI	<ul style="list-style-type: none"> Age > 18 yrs SCI > 1 year Wheelchair User Able to use arms for exercise Sedentary (<60 minutes of exercise/week) NO Cognitive Impairment (Folstein's Mini-Mental State Exam Score < 24) NO Depression (Centers for Epidemiological Studies Depression Scale Score > 16) NO poorly controlled blood pressure (SBP > 159 or DBP > 95 mmHg) 	Recruiting	44	Phase: Not Applicable, Primary Purpose: Prevention, Intervention Model: Factorial Assignment, Masking: Triple	Rehabilitation	General Health	<ul style="list-style-type: none"> Physical Activity (PARA-SCI) Grip strength Respiratory functioning (Peak Expiratory Volume) Blood pressure Heart rate Physical activity self-efficacy scale Outcomes expectations for exercise scale Demographics Sleep disturbance Depression Pain intensity Anxiety <p>F/U 12 weeks</p>	June 2021	January 27, 2021	June 30, 2022	Birmingham, AL, USA
NCT04699474	Hopital du Sacre-Coeur de Montreal	Leg cycling in bed using motorized ergometer	<ul style="list-style-type: none"> Blunt traumatic SCI SCI Level C0 to L2 AIS A-C Spine surgery performed within 48 hours of injury NO moderate or severe brain injury NO pre-existing neurological disorder NO complete spinal cord transection 	Recruiting	45	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking Mobility	<ul style="list-style-type: none"> Ambulation (independent walking) Neurological recovery Spinal Cord Independence Measure Spasticity Secondary Health related quality of life (WHO-QOL-BREF) <p>F/U 6 months</p>	January 2021	January 7, 2021	August 25, 2021	Montréal, QC, Canada
NCT04688229	Shirley Ryan AbilityLab	Hummingbird hand training device plus standard of care rehabilitation. The Hummingbird is a comprehensive hardware and software platform that isolates the hand, wrist, and forearm in a neutral and comfortable position. The platform also comprises therapeutic software to train exploratory, individuated, and inter-digit complex finger movements by coupling force and torque output to NeuroAnimation physics-based virtual animals in engaging therapeutic experiences.	<ul style="list-style-type: none"> Age 18-75 yrs Traumatic SCI within 1-2 months Cervical injury at C8 or above (tetraplegia) AIS A, B, C and D No uncontrolled medical problems including pulmonary, cardiovascular or orthopedic disease No ongoing cord compression or a syrinx in the spinal cord or who suffer from a spinal cord disease such as spinal stenosis, spina bifida or herniated cervical disk. 	Not yet recruiting	24	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Graded and Redefined Assessment of Strength, Sensibility, and Prehension (GRASSP) measure Action research arm test (ARAT) Box and Block test Maximum voluntary force (MVF) Average strength index Individuation index <p>F/U 3 months</p>	October 2021	December 29, 2020	September 21, 2021	Chicago, IL, USA
NCT04641793	Shirley Ryan AbilityLab	Motion and EMG Control. Two methods (direct and indirect) for integrating motions and EMG signals	<ul style="list-style-type: none"> Age 16-65 yrs SCI Level C3-6 AIS A-C NO metal in head with the exception of dental work or any ferromagnetic metal elsewhere in the body Personal history of epilepsy (untreated with one or a few past episodes), or treated patients 	Recruiting	60	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Parallel Assignment, Masking: Single	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Time to task completion Muscle activity (EMG) Cortico spinal connectivity (motor evoked potentials, MEPs) <p>F/U: 1 week</p>	January 2020	November 24, 2020	November 24, 2020	Chicago, IL, USA

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
NCT04632290	Ecole Polytechnique Fédérale de Lausanne	STIMO-BSI system. Bilateral implantation of epidural electrocorticography devices. The decoded motor intentions are driving the implanted spinal cord stimulation system. Brain-controlled spinal cord stimulation is used for training and rehabilitation to recover voluntary movements.	<ul style="list-style-type: none"> Having completed the main phase of the STIMO study (NCT02936453) AIS A-D Level T10 and above Focal spinal cord disorder caused by either trauma or epidural, subdural or intramedullary bleeding SCI ≥ 12 months NO epilepsy NO spinal stenosis NO use of an intrathecal Baclofen pump. NO active implanted cardiac device such as pacemaker or defibrillator 	Enrolling by invitation	3	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Safety Measure (adverse events, device deficiencies) WISCI II score 10mWT Weight bearing capacity SCIM III score 6MWT Time Up and Go Maximum Voluntary Contraction ISNCSCI score Modified Ashworth Scale Berg Balance Scale Gait Analysis WHOQOL-BREF BCI performance measures (accuracy) Upper Limb Neurobiomechanics ECoG signal stability Somatosensory-evoked potentials (SSEP) <p>F/U: Through study completion, an average of 1 year</p>	July 2021	November 17, 2020	November 5, 2021	Lausanne, Switzerland
NCT04604951	University of British Columbia	Non-invasive transcutaneous spinal cord stimulation	<ul style="list-style-type: none"> Resident of British Columbia (Canada) with active provincial medical services plan 18-65 years of age Chronic traumatic SCI (non-progressive, with complete motor paralysis) at or above the T6 spinal segment. vSCI ≥ 1 year, at least 6 months from any spinal surgery. Documented presence of bladder dysfunction (NDO during UDS) Documented presence of bowel or sexual dysfunction AIS A-B NOT Ventilator dependent NO Intrathecal baclofen pump. NO implanted metal in trunk or spinal cord under the sites of application of electrodes (between anode and cathode) for those who are allocated to receive TCSCS. 	Recruiting	40	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Blood pressure (number of potential adverse cardiovascular events, i.e. autonomic dysreflexia) Surface electromyography (EMG, motor threshold for skeletal muscles known to be involved in lower urinary tract and bowel control) Urodynamics parameters Mean maximum resting anorectal pressure Frequency of urinary incontinence Neurogenic bladder symptoms Frequency of fecal incontinence Neurogenic bowel symptoms Participant's sexual function and satisfaction with their overall sexual life (International Index of Erectile Function, Female Sexual Function Index, semi-structured qualitative interview) <p>F/U: 25 weeks</p>	May 2022	October 27, 2020	November 30, 2022	Vancouver, BC, Canada
NCT04577573	VA Office of Research and Development	Functional prototype of an instrumented glove (Cognition glove) to alert the user about secure grasp of objects.	<ul style="list-style-type: none"> SCI > 12 months SCI level C1-T1 Hand weakness: score of 2-4 out of 5 on manual muscle testing of finger extension, finger flexion, or finger abduction in either hand 	Recruiting	20	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Crossover Assignment, Masking: Single	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Time to achieve secure grasp upon initial contact Time to complete pick-up and placement of object Motion pathlength in moving object Error in placing object onto target <p>F/U immediately after the procedure</p>	May 2021	October 8, 2020	November 21, 2022	Bronx, NY, USA
NCT04578665	Shirley Ryan AbilityLab	Behavioral Interaction Conditions. Complete a tracking task (solo, collaboration task, competition task or cooperation task). Haptic Impedance Level (rigid, medium or soft), Skill Level of Partner (novice or expert). Participants will start experimentation paired as novice-novice, and at the end of the session may be invited to continue additional sessions to be paired as the expert in a novice-expert dyad.	<ul style="list-style-type: none"> Age 18-80 yrs AIS C-D SCI ≥ 6 months Ability to walk >10 m independently (w or w/o assistive devices or bracing) Normal hearing and vision 	Recruiting	544	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> lower limb motor control motor output from surface EMG of lower limb muscles 6 minute walking test 10 meter walking test Modified Ashworth Scale (Spasticity) BERG balance scale (BBS) functional gait assessment (FGA) <p>F/U 5 months (average)</p>	July 2021	October 8, 2020	July 15, 2022	Chicago, IL, USA
NCT04568928	Laval University	Overground locomotor training program using a powered exoskeleton combined with functional electrical stimulation (OLTP/PE+FES), or sham stimulation (OLTP/PE+shamFES), or OLTP without FES.	<ul style="list-style-type: none"> Age 18-70 yrs Incomplete SCI sufficient upper extremity strength and function to use a walker with wheels be able to stand >15 minutes NOT have osteoporosis NOT have epilepsy 	Enrolling by invitation	10	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (open label)	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> Modified 6 Minute Walk Test (6MWT) 10 Meter Walk Test (10mWT) Modified Timed Up and Go test (TUG) Walking Index for Spinal Cord Injury (WISCI-II) Brief pain inventory questionnaire Spinal Cord Injury Secondary Conditions scale (SCI-SCS) Modified Ashworth Scale (MAS) Electromyographic (EMG) activity of leg muscles during walking <p>F/U 3 months</p>	December 2020	September 29, 2020	August 24, 2022	Québec, QC, Canada
NCT04496609	Hopital Foch	Stimulation and automated rehabilitation for 40 working days, then washout during 30 days, then automated rehabilitation for 40 working days	<ul style="list-style-type: none"> Age 18-65yrs AIS B or C SCI above T10 (sensation preserved below level of lesion) Absence of significant motor deficit of the upper limbs or recovered motor deficit (muscular score ≥ 4/5) SCI > 2years 	Recruiting	14	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Assessment of number of patients able to move over a distance of 5 metres Percentage of patients capable of moving at the end of treatment Assessment of vesico-sphincter function Assessment of the genito-sexual function Assessment of the excitability of the spinal neuronal circuits Assessment of the kinetics of action of the induced effects Quality of life (EQ5D-3L) AE/SAE related to tolerance <p>F/U 6 months</p>	July 2021	August 3, 2020	August 3, 2022	Garches, France Suresnes, France
NCT04440709	University Hospital Tuebingen	The brain/neural hand exoskeleton restores hand motor function by translating user's intention into grasping motions	<ul style="list-style-type: none"> SCI C5-C6 SCI > 6M AIS B,C no severe spasticity (Ashworth grade ≥ 4) 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Supportive Care, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Change in Berlin Bimanual Task Assessment (BEBITA) score <p>F/U NA</p>	June 2020	July 22, 2020	June 22, 2020	Tübingen, Germany Berlin, Germany

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NCT04474106	AUVA Trauma Center Meidling	Extracorporeal shock wave therapy (shockwave generator orthogold 100)	<ul style="list-style-type: none"> traumatic SCI Age >= 18 yrs SCI <= 24 hours no serious traumatic brain injuries no pacemakers or implantable defibrillators 	Recruiting	246	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Quadruple	Technology	General Health	<ul style="list-style-type: none"> Total motor scores AIS grade Degree of spasticity Walking ability (WISCI II, TUG, 10MWT, 6MWT=) Urological function Plantar reflex SCIM II Adverse events (AEs) Dexterous control (Nine-Hole Peg Test, Grasp and Release Test, Pinch grip, Clenched grip, Pencil grip, Lumbrical grip) 	July 2020	July 16, 2020	February 24, 2022	Multicenter: Austria (Innsbruck, Bad Häring, Feldkirch, Graz, Tobelbad, Klosterneuburg, Linz, Salzburg, St. Pölten, Wien) Berlin, Germany
NCT04468919	Oregon Health and Science University	Brain-Computer Interface (BCI) Functional Implementation Toolkit (FIT) to customize the method for each individual end user	<ul style="list-style-type: none"> Age 18-89 yrs severe speech and physical impairment no implanted hydrocephalus shunt, cochlear implant or deep brain stimulator 	Recruiting	60	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Sequential Assignment, Masking: None (Open Label)	Technology	Mental Health	<ul style="list-style-type: none"> Typing Accuracy Typing Speed Information transfer rate User experience 	July 2022	July 13, 2020	September 13, 2022	Portland, OR, USA
NCT04460872	North Florida Foundation for Research and Education	Intramuscular testosterone injection and locomotor training (4 sessions/week for 2-3 months)	<ul style="list-style-type: none"> male Age >18 yrs incomplete SCI L1 and higher or complete SCI T2-L1 SCI >= 1y Locomotor dysfunction Diagnosis of first time SCI including etiology from trauma, vascular, or orthopedic pathology 	Recruiting	21	Phase: Phase 2, Primary Purpose: Supportive Care, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Drug	Standing/Walking/Mobility	<ul style="list-style-type: none"> muscle cross-sectional area 6MWT bone mineral density knee extensor peak torque 10mWT bone resorption/formation marker 	January 2021	July 8, 2020	September 28, 2022	Gainesville, FL, USA Jacksonville, FL, USA
NCT04458324	Spaulding Rehabilitation Hospital	Functional Electrical Stimulation Row Training (FESRT, 2-3 times/week for 6 months) combined with a medication against respiratory abnormalities (Bupropion Hydrochloride), and with non-invasive ventilation (NIV)	<ul style="list-style-type: none"> SCI outpatients 18-45 years SCI for 3 months - 6 years AIS A, B, or C SCI C1-T3 wheelchair users no diabetes, cancer, epilepsy 	Recruiting	70	Phase: Phase 2, Primary Purpose: Prevention, Intervention Model: Factorial Assignment, Masking: Double	Drug	General Health	<ul style="list-style-type: none"> aerobic exercise capacity ventilation during exercise glucoregulatory status (hemoglobin) serum lipids (cholesterol and triglycerides) dual x-ray absorptiometry (DXA) Change in baseline ventilatory function (spirometry) 	December 2020	July 7, 2020	January 4, 2023	Cambridge, MA, USA
NCT04453943	North Carolina State University	Exoskeleton Walking and Exoskeleton Sitting to Stading - with or without FES. Walking movements will be elicited by the hybrid walking platform that combines a powered exoskeleton and an FES system.	<ul style="list-style-type: none"> Age 18-60 yrs complete or incomplete SCI SCI >= 1yrs SCI T1-T10 no heart conditions and pacemakers 	Not yet recruiting	30	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Controls Algorithm Performance - Limb Angle Errors Muscle fatigue Index to Measure FES-Induced Muscle Fatigue Participant Verbal Feedback 	July 2020	July 1, 2020	July 1, 2020	Raleigh, NC, USA
NCT04393922	Shirley Ryan AbilityLab	Acoustic stimuli (Startle), Description: A startle stimulus (120 dB, 500 Hz, 50 ms) will be delivered through headphones.	<ul style="list-style-type: none"> Age 18-75 yrs Chronic SCI (>= 1 year) Incomplete SCI SCI T12 or above no pacemaker no history of seizures 	Recruiting	120	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Crossover Assignment, Masking: Single	Technology	Spasticity	<ul style="list-style-type: none"> MEP recruitment curves StartReact Modified Ashworth Scale (MAS) Pendulum Test 10-meter walk test GRASSP Toronto Rehabilitation Institute-Hand Function Test (TRI-HFT) 	May 2020	May 19, 2020	June 22, 2022	Chicago, IL, USA
NCT04369131	Quality Living, Inc.	Functional electrical stimulation of the calves, quads and/or abdominals	<ul style="list-style-type: none"> Age 19-70 yrs SCI C1-C5 AIS ns 	Enrolling by invitation	10	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Change in tilt table angle Maximum tilt table angle Blood pressure 	December 2021	April 30, 2020	September 8, 2022	Omaha, NE, USA
NCT04340063	VA Office of Research and Development	Gait training performed on a treadmill or in a Movement Amplification Environment	<ul style="list-style-type: none"> Age 18 - 80 years SCI C1-T10 AIS C or D >= 6 months post spinal injury no excessive spasticity in the lower limbs 	Recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> Daily Stepping LEMS WISCI II Functional Gait Assessment (FGA) 10MWT Activities Specific Balance Confidence (ABC) Scale Balance Evaluations Systems Test (BESTest) Berg Balance Scale (BBS) The World Health Organization Quality of Life Scale Urinary Incontinence Questionnaire (ICIQ-UI SF) 	October 2020	April 9, 2020	January 23, 2023	Hines, IL, USA
NCT04323449	VA Office of Research and Development	Control methods for a wheelchair-mounted robotic manipulator	<ul style="list-style-type: none"> Age 18yrs using a power wheelchair as primary means of mobility no impaired vision 	Recruiting	16	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Task completion time Mode Switching Frequency Task success rate NASA Task Load Index (TLX) System Usability Scale (SUS) 	April 2022	March 26, 2020	May 31, 2022	Pittsburgh, PA, USA
NCT04307303	Salisbury NHS Foundation Trust	functional electrical stimulation of the abdominal muscles	<ul style="list-style-type: none"> Age >=18 yrs SCI C1-T12 AIS NS >=1 year post spinal injury no implanted electrical devices (cardiac pacemaker in situ or other) 	Not yet recruiting	36	Phase: Not Applicable Primary Purpose: Treatment Intervention Model: Parallel Assignment Masking: Triple	Technology	Bowel Health	<ul style="list-style-type: none"> Time required for defecation Neurogenic bowel dysfunction sexual function index 	May 2020	March 13, 2020	March 13, 2020	Salisbury, UK
NCT04302259	Rhode Island Hospital	Intelligent Spine Interface - Commercial (ISI-C) is a Epidural Electrical Stimulation (EES). The aim is to restore volitional control of the lower limb.	<ul style="list-style-type: none"> Age 18-65yrs SCI C7/T1- T10 AIS A-B traumatic SCI SCI > 1 year able to ambulate with a wheelchair or crutches no intrathecal baclofen or morphine pump no implanted device 	Not yet recruiting	3	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Safety: adverse events ISNCSCI 6MWT 10MWT TUG Spinal cord Injury Functional Ambulation Inventory SCI-QOL Berg Balance Scale 	September 2021	March 10, 2020	October 26, 2022	Providence, RI, USA

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NCT04292717	University of Zurich	Gait training three times a week for 6 weeks (18 training sessions). Each exercise session will last for 1 hour and will be conducted and supervised by an experienced physical therapist.	<ul style="list-style-type: none"> Age 18-80 yrs SCI ≥ T12 SCI >6 months AIS C-D walk without assistance or devices on the treadmill and 10m over ground 	Recruiting	56	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> 6MWT ISNCSCI WISCI II 10MWT TUG Kinematics, EMG SCIM III MR/MRS 	January 2021	March 3, 2020	August 16, 2022	Zurich, Switzerland
NCT04288245	Baylor Research Institute	Active Vagus Nerve Stimulation paired with upper extremity rehabilitation.	<ul style="list-style-type: none"> Age 18-64yrs cervical SCI ASIA B, C, D traumatic SCI ≥ 12 months no SCI due to sharp objects, firearms 	Recruiting	20	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Crossover Assignment, Masking: Triple	Technology	General Health	<ul style="list-style-type: none"> Functional Independence Score Description Spinal Cord Independence Measure (SCIM) III AIS Walking Index for Spinal Cord Injury II (WISCI II) GRASSP 	February 2021	February 28, 2020	January 5, 2023	Dallas, TX, USA
NCT04286191	Medical University of South Carolina	Operant Conditioning training intervention in which the brain-spinal cord-muscle pathways are strengthened. Transcranial magnetic stimulation (TMS).	<ul style="list-style-type: none"> SCI ≥1 year AIS NS SCI level NS able to ambulate at least 10 m with or without an assistive device (except for parallel bars) Signs of weak ankle dorsiflexion at least unilaterally 	Recruiting	44	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Triple	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> MEP/CMEP Short Interval Intra-cortical Inhibition (SICI) H-reflex amplitude F-wave amplitude and occurrence tibialis anterior EMG amplitude Ankle function 10-meter walk test 6-minute walk test 	February 2021	February 26, 2020	December 5, 2022	Charleston, SC, USA
NCT04250688	Shirley Ryan AbilityLab	Ekso Training with FES	<ul style="list-style-type: none"> Age 18-70 yrs SCI C7-T11 AIS A, B, C, D SCI <6 months at completion of study able to tolerate upright standing for a minimum of 30 minutes 	Enrolling by invitation	10	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 6MWT 10MWT 	January 2016	January 31, 2020	May 4, 2022	Chicago, IL, USA
NCT04241250	VA Office of Research and Development	EAW+SCES (exoskeleton and spinal cord epidural stimulation): 3 months of EAW training followed by 6 months of SCES.	<ul style="list-style-type: none"> Age 18-70 yrs SCI C1-T10 traumatic SCI AIS A, B no implanted pacemakers and/or implanted defibrillator devices 	Recruiting	10	Phase: Phase 2, Primary Purpose: Health Services Research, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 10-meter walking speed (m/sec) Muscles electromyography (EMG) during locomotion 	July 2020	January 27, 2020	April 13, 2022	Richmond, VA, USA
NCT04221373	Icahn School of Medicine at Mount Sinai	Ekso™ powered exoskeleton-assisted walking (EAW) for early training compared to standard of care.	<ul style="list-style-type: none"> Age ≥18 yrs SCI level NS AIS NS Height between 5'2" and 6'2" (1.6 meters to 1.9 meters) 	Enrolling by invitation	40	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Technology	Pain	<ul style="list-style-type: none"> ISNCSCI FIM SCIM Pain: ISCBPDS 2.0 and SCIPI 	September 2019	January 9, 2020	April 20, 2022	New York, NY, USA
NCT04194099	Taipei Medical University Hospital	Paired Nerve Stimulation (PNS) with various stimulation parameters of repetitive transcranial magnetic stimulation (rTMS) combined with trans-spinal electrical stimulation (tsES) vs. Sham PNS, followed by cycling exercise.	<ul style="list-style-type: none"> Age 20-65 yrs Vertebral level above T10 AIS B, C, D Chronic SCI SCI>1yr 	Recruiting	12	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single (Outcomes Assessor)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> MEP Resting Motor Threshold Modified Ashworth Scale LEMS fMRI EMG 	December 2019	December 11, 2019	May 5, 2022	Taipei City, Taiwan
NCT04193709	University of Louisville	Arm 1: Cross-sectional and observational autonomic function data collection during bladder filling and bowel stimulation Arm 2: use of scES in the lab and at home configured for maintenance of normative blood pressure and heart rate during bladder filling and bowel evacuation	<ul style="list-style-type: none"> Age 18-70 yrs SCI Level NS AIS A, B, C, D 	Recruiting	70	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Bladder Health	<ul style="list-style-type: none"> 24hr Blood Pressure monitor Bladder Function 	January 2021	December 10, 2019	May 9, 2022	Louisville, KY, USA
NCT04105114	University of Louisville	Three arm study of different combinations of non-invasive electrical spinal cord stimulation, oral buspirone vs. placebo, BWSTT, gravity neutral device, Ekso Bionics Exoskeleton, rolling walker. Group 1 complete SCI-gravity neutral stepping; Group 2 complete SCI exoskeleton assisted stepping; Group 3 incomplete SCI overground stepping.	<ul style="list-style-type: none"> Age 18-65 yrs SCI Level T2-T7 AIS A, B, C Chronic SCI> 1yr 	Recruiting	15	Phase: Early Phase 1 Primary Purpose: Basic Science, Intervention Model: Crossover Assignment Masking: Triple (Participant, Investigator, Outcomes Assessor)	Technology	General Health	<ul style="list-style-type: none"> EMG Kinematics Blood Pressure Heart Rate DEXA Bone/Soft Tissue Urodynamics 	September 2019	September 26, 2019	July 12, 2022	Louisville, KY, USA
NCT04101916	Helsinki University Central Hospital	Paired Associative Stimulation (PAS) using transcranial magnetic stim paired with Lower Extremity (LE) peripheral nerve electrical stimulation vs. Sham PAS. Several sessions/wk for 12 weeks.	<ul style="list-style-type: none"> Age 18-75 yrs Cervical SCI AIS NS Voluntary finger muscle or MEP obtained from hand muscles Subacute SCI 1m<SCI≤4m 	Enrolling by invitation	24	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Double	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Muscle Strength Testing SCIM 	October 2019	September 24, 2019	March 16, 2022	Helsinki, Finland
NCT04102826	Robert Jones and Agnes Hunt Orthopaedic and District NHS Trust	Functional Electrical Stimulation (FES) to nerves supplying weakened UE muscles combined with the use of mobile arm supports to compensate for diminished strength where necessary	<ul style="list-style-type: none"> Age NS Cervical SCI Subacute-Chronic SCI SCI≤6wks 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Canadian Occupational Performance Measure (COPM) 	September 2021	September 15, 2019	November 11, 2021	Oswestry, UK
NCT04077346	University of Louisville	BioStim-5 Transcutaneous Spinal Stimulation (TcStim) alone, Activity-Based Locomotor Training (AB-LT) alone, and in combination (AB-LT+TcStim). 120 weekday sessions (5x/week within 8 months)	<ul style="list-style-type: none"> Age 4-12 yrs SCI T12 & above Unable to stand, walk, initiate steps Chronic SCI> 1yr 	Enrolling by invitation	19	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Ability to take independent step EMG Angular excursions of LE Blood Pressure Heart Rate Compliance with Sessions 	April 2021	September 4, 2019	May 25, 2022	Louisville, KY, USA

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NCT04052776	Centre Hospitalier Universitaire Vaudois	Orally administered buspirone and levodopa/carbidopa taken individually and in combination plus placebo, by subjects who have completed the STIMO Study NCT02936453 (see above) and are enrolled in the STIMO ESS extension study. 4 arms-all subjects will receive all 4 (crossover).	<ul style="list-style-type: none"> Age 18-65 yrs Enrolled in STIMO Ext. Level NS AIS A, B, C, D Chronic SCI Enrolled in STIMO Ext. (see above) 	Not yet recruiting	8	Phase: Phase 1. Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Quadruple	Technology	General Health	<ul style="list-style-type: none"> Safety: AE, SAE, Side Effects Blood Pressure Heart Rate Pendulum Test ISNCSCI Kinematics EMG 10MWT 	May 2020	August 12, 2019	May 26, 2022	Lausanne, Switzerland
NCT04050696	BrainQ Technologies Ltd.	Physical Therapy run-in/baseline with Machine Learning analysis of EEG/MEG (BCI) and EMG patterns during functional motor tasks to create low-intensity, non-invasive patterned electromagnetic field CNS stimulation.	<ul style="list-style-type: none"> Age 18-75 yrs SCI C1-C8 AIS B, C, D GRASSP motor 5-30/50 at least one side Chronic SCI 12msSCIs30m 	Recruiting	8	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> GRASSP CUE-T ISNCSCI Ashworth SCIM III SCI-QoL MRI 	August 2019	August 8, 2019	September 28, 2021	Miami, FL, USA West Orange, NJ, USA Ramat Gan, Israel
NCT04043715	University of Washington	Transcutaneous Spinal Stimulation (TSS), Epidural Spinal Stimulation (ESS), Physical Therapy (PT) in Phased intervention: baseline testing→PT only→PT+TSS→Washout→PT+ESS→follow-up testing	<ul style="list-style-type: none"> Age 21-70 yrs LE impairment Level NS AIS NS Candidate for ESS implant (pain) Chronic SCI≥1yr 	Recruiting	6	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Neuromuscular Recovery Scale ISNCSCI WISCI SSEP, MEP Berg Balance Gait Kinematics 6MWT 10MWT NBSS 	August 2019	August 2, 2019	October 8, 2020	Seattle, WA, USA
NCT03998527	University of Louisville	6 Groups: 1) Transcutaneous electrical spinal cord stimulation (TcESCS), 2) Respiratory training (RT), 3) TcESCS+RT, 4) TcESCS+Arm training, 5) TcESCS+Trunktraining, 6) TcESCS to Non-Disabled control group	<ul style="list-style-type: none"> Age 18-99 yrs SCI T5 or above ≥15% deficit in FVC or FEV1 Chronic SCI≥1yrs 	Enrolling by invitation	36	Phase: Early Phase 1, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single (Outcomes assessor)	Technology	General Health	<ul style="list-style-type: none"> Respiratory EMG Pulmonary Function Spirometry Max Airway Pressure FNPA MMR 	May 2023	July 26, 2019	June 3, 2022	Louisville, KY, USA
NCT04032990	University of Louisville	Transcutaneous spinal stimulation with activity-based upper extremity training (40 sessions, 1.5 hours/day, 5 days/week); stimulation will be applied intermittently (with Biostim-5 transcutaneous spinal stimulator) for no more than 10 minutes at a time	<ul style="list-style-type: none"> Age 4-18 yrs SCI T1 or above Moderate-severe UE deficit Chronic SCI>1yr 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Device Feasibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Safety/Feasibility Skin Irritation Pain Blood Pressure Arm Excursion, Grip Strength 	November 2019	July 25, 2019	March 28, 2022	Louisville, KY, USA
NCT03949660	University of Louisville	Must be already be enrolled in epidural spinal stimulation and training study. Various epidural stimulation protocols will be performed with assessments of bowel function and QoL.	<ul style="list-style-type: none"> Age 18-75 yrs SCI Level NS AIS A, B Cardiovasc & bowel dysfunction Chronic SCI≥2yrs 	Recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Bowel Health	<ul style="list-style-type: none"> Wireless Bowel Motility Capsule Blood Pressure Heart Rate Anorectal Pressure Bowel Diary SCI-QoL 	September 2019	May 14, 2019	May 12, 2022	Louisville, KY, USA
NCT03930056	Shirley Ryan AbilityLab	C-Brace II use vs. traditional KAFO use. Following evaluation and brace fitting, participants will receive 10-20 one hour training sessions with assigned brace, then transition to home use for 3 month period.	<ul style="list-style-type: none"> Age 18-80 yrs SCI Level NS AIS NS Requires LE orthotic bracing including knee for instability Chronic SCI 3mos≥SCI≥24mos 	Recruiting	30	Phase: Not Applicable, Primary Purpose: Supportive Care, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 6MWT 10MWT MMT LE ROM Ashworth WISCI Gaitrite Data Capture 	April 2019	April 29, 2019	October 25, 2022	Chicago, IL, USA
NCT03922802	Shirley Ryan AbilityLab	Acute intermittent hypoxia (AIH) + non-invasive spinal cord stimulation (TSCS) + walking rehabilitation.	<ul style="list-style-type: none"> Age ≥ 18yrs SCI > 6 months Level below C2 AIS A, B, C, D No epilepsy or a concussion within 6 months No metal implants in the head or face No implanted cardiac pacemaker or drug pump 	Recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Single (Participant)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Gait kinematics using Gait Rite 10 Meter Walk Test 6 Minute Walk Test using V02 analysis TUG 	December 2020	April 22, 2019	October 26, 2022	Chicago, IL, USA
NCT03892746	Cleveland Clinic US Department of Defense	Non-invasive brain stimulation tDCS (Transcranial Direct Current Stimulation) vs. sham tDCS to the area in the brain controlling the weaker muscle of the weakest upper limb while receiving task-oriented training for 15 Session (5d/wk X 3wks)	<ul style="list-style-type: none"> incomplete SCI Age 18-75yrs SCI level C1-C8 AIS NS Chronic SCI≥1yr Triceps strength of weaker UE is 1 grade- stronger UE 	Recruiting	49	Phase: Phase 1. Phase 2, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> ISNCSCI UEMS GRASSP Ashworth COPM SCIM TMS H-reflex 	July 2019	March 27, 2019	October 4, 2022	West Orange, NJ, USA Cleveland, Ohio, USA
NCT03714997	Indiana University	Comparison of two different intensities of walking training (30 one-hour sessions on a treadmill, overground, and on stairs). High intensity training will target achievement of heart rates close to 80% of heart rate reserve; lower intensity to heart rates from 30% to 40% of heart rate reserve.	<ul style="list-style-type: none"> Age 18-75 yrs SCI C1-T10 AIS C, D Must tolerate 10m of standing without hypotension Chronic SCI≥1yr 	Not yet recruiting	80	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> 10MWT 	July 2019	October 22, 2018	April 22, 2019	Indianapolis, IN, USA
NCT03698149	University of California, San Francisco	Brain implantation of cortical electrodes to enable electrocorticography (ECoG) recording of brain activity. Study subjects will undergo training and assessment of their ability to control a complex robotic system and/or produce speech	<ul style="list-style-type: none"> Age≥21yrs Limited UE use due to SCI or other neuro disability. Lives close to UCSF Chronic SCI≥1yr 	Recruiting	3	Phase: Early Phase 1 Primary Purpose: Device Feasibility Intervention Model: Single Group Assignment Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Adverse Events associated with ECoG-based interface 	August 2018	October 5, 2018	August 2, 2022	San Francisco, CA, USA

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
NCT03690700	Spinal Cord Injury Centre of Western Denmark	Active vs. Sham low-intensity blood-flow restricted exercise (BFRE); low-intensity strength training (20-30 % of max) while using circumferential cuffs during exercise adjusted to maintain arterial inflow to the muscles while preventing venous return.	<ul style="list-style-type: none"> Age >18 yrs Elbow flex & Wrist extend strength 2-4/5 AIS A, B, C, D Subacute/Chronic SCI>1yrs 	Recruiting	28	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Rehabilitation	Standing/Walking/Mobility	<ul style="list-style-type: none"> SCAR 10MWT 6MWT TUG H-Reflex Pain Level QoL BDS COPM 	May 2020	October 1, 2018	September 27, 2022	Viborg, Denmark
NCT03680872	Chad Bouton	Bidirectional Neural Bypass System: Implantation of microelectrode arrays into the primary motor cortex to record neural activity associated with desired movements and into the primary somatosensory cortex to deliver stimulation in order to provide sensory perception. Participation in 3 study session/wk X 12mos	<ul style="list-style-type: none"> Age 22-65 yrs Stable cervical SCI Finger strength 0-2 Plamar finger sensation 0-2 Chronic SCI≥1yr 	Recruiting	3	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> GRASSP-restoration of hand movement Restoration of tactile sensation of the hand 	September 2019	September 21, 2018	December 29, 2022	Manhasset, NY, USA
NCT03504826	Brooks Rehabilitation	Locomotor training with Cyberdyne Hybrid Assistive Limb (HAL) and locomotor training overground with or without the HAL device. 60 sessions (5 days/week for 12 weeks).	<ul style="list-style-type: none"> Age 18-80 yrs Level NS AIS B, C, D Can walk 10ft Chronic SCI>1yr 	Recruiting	24	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 10MWT 6MWT EMG 	November 2018	April 20, 2018	December 20, 2022	Jacksonville, FL, USA
NCT03447509	VA Office of Research and Development	Transcranial non-invasive Magnetic Stimulation targeting late indirect descending volleys (TMS) (vs. sham TMS) with acoustic startle during performance of UE movement tasks/training	<ul style="list-style-type: none"> Age 18-85 yrs SCI above C8 AIS A, B, C, D Visible grip/UE movement ability Chronic SCI>1yr 	Recruiting	300	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: Single	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Motor Evoked Potential Amplitude Grip Strength 9-Hole Peg Test 	January 2020	February 27, 2018	January 27, 2023	Miami, FL, USA Hines, IL, USA
NCT03443700	Montecatone Rehabilitation Institute S.p.A.	EKSO-GT locomotor training plus 8 weeks standard locomotor training vs. 8 weeks of standard locomotor training alone	<ul style="list-style-type: none"> Age 18-65 yrs SCI T1-L1 AIS C, D Functional Gait (incl with braces) Chronic SCI 1yr<SCI<5yrs 	Recruiting	40	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 10MWT 6MWT WISCI II Ashworth LEMS EMG SSEP fMRI 	December 2020	February 23, 2018	December 16, 2021	Imola BO, Italy
NCT03364660	University of Louisville	Spinal Cord Epidural Stimulation utilizing stim parameters for voluntary movement, standing, or cardiovascular responses combined with leg movement training or stand training while sitting or supine.	<ul style="list-style-type: none"> Age≥18 yrs SCI level NS Unable to move legs/stand Chronic SCI≥2yrs 	Recruiting	36	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Cardiovascular Assessments Functional Movement Assessments Standing Assessments 	November 2017	December 6, 2017	November 18, 2022	Louisville, KY, USA
NCT03161067	Johns Hopkins University	Investigation on the Bidirectional Cortical Neuroprosthetic System	<ul style="list-style-type: none"> Age 22-65 yrs AIS A-C SCI level C4-C6 >1 year after SCI >5 years life expectancy NO memory impairment 	Recruiting	5	Phase: Not Applicable, Primary Purpose: Device Feasibility Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Sensory function	ARAT (Successful control of an assistive device)	August 2017	May 19, 2017	September 27, 2022	Baltimore, MA, USA
NCT03057652	The University of Texas Health Science Center, Houston	algorithmic-based evaluation and treatment approach for wearable robotic exoskeleton (WRE) gait training using ReWalk, Ekso, and REX systems; randomly assigned order of device use. Up to 15 training sessions per device.	<ul style="list-style-type: none"> Age≥18 yrs SCI level NS AIS NS Ashworth <3 Chronic SCI>6m 	Recruiting	75	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 10 Meter Walk 6 Minute Walk Surface EMG Oxygen Consumption Gait Kinematics Bone Mineral Density 	March 2016	February 20, 2017	May 23, 2022	Houston, TX, USA
NCT03053791	University of Zurich	Unilateral implantation of a Medtronic Activa SC deep brain stimulation system in the mesencephalic locomotor region	<ul style="list-style-type: none"> Age 18-75 yrs SCI T10 & above AIS C, D walk 10 meters Chronic SCI≥6m no implanted devices (e.g., pacemaker) 	Recruiting	5	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> 6 Minute Walk 10 Meter Walk Test TUG Ashworth AIS GRASSP SF-36 Lower urinary tract function Activity monitoring 	February 2017	February 17, 2017	May 20, 2022	Zürich, Switzerland
NCT03026816	University of Minnesota	Implanted epidural spinal cord stimulator for improving volitional motor activity/autonomic function in persons with chronic motor complete SCI; comparing outcomes with stimulator on vs. off (sham stimulation).	<ul style="list-style-type: none"> Age≥22 yrs SCI C6-T10 AIS A, B Chronic SCI>1yr 	Recruiting	100	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Brain Motor Control Assessment Volitional Movement Systolic blood pressure during epidural stimulation Cerebral Blood Flow (tilt table) Stroop test 	August 2017	January 20, 2017	May 2, 2022	Minneapolis, MN, USA
NCT02991248	Shirley Ryan AbilityLab	Three arm study comparing robotic/pelvic force-perturbation treadmill training with 1) active vs. 2) sham transcutaneous spinal direct current stimulation (tsDCS), and 3) standard treadmill training only. Three treatment sessions per week for 6 weeks.	<ul style="list-style-type: none"> Age 18-65 yrs SCI C4-T10 AIS C, D Able to walk 10 meter with no more than AFO Time after SCI NS 	Recruiting	54	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Parallel Assignment, Masking: Single	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Gait Speed (overground) 6MWT Dynamic Gait Index Berg Balance Scale 	November 2018	December 13, 2016	November 3, 2022	Chicago, IL, USA
NCT02978638	Palo Alto Veterans Institute for Research	Implantation of Finetech Vocare Bladder System—a sacral nerve root stimulator. The study tests the use of the system to inhibit bladder contractions by electrically stimulating sensory nerves (as an alternative to cutting sensory nerves).	<ul style="list-style-type: none"> Age≥22 yrs SCI below C4 AIS A Dyssynergia/Detrusor Hyper-reflexia Chronic SCI≥2yr 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Bladder Health	<ul style="list-style-type: none"> Bladder Capacity (Cystometry) Frequency of Incontinence 	September 2014	December 1, 2016	April 25, 2022	Palo Alto, San Jose, CA, Albuquerque, NM, USA

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
NCT02451683	Shirley Ryan AbilityLab	Study of motor task training with real or sham stimulation assessing electrophysiological parameters of time domain and location	<ul style="list-style-type: none"> Age 18-85 yrs SCI C8 & above Some grasp and reach ability Chronic SCI≥6months 	Recruiting	300	Phase: Not Applicable, Primary Purpose: Diagnostic, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Functional tests of arm/hand function Cortical Neurophysiology Upper limb movements scale F/U 5 months	May 2020	May 22, 2015	May 25, 2021	Chicago, IL, USA
NCT02446210	Shirley Ryan AbilityLab	Magstim 200 stimulator for Transcranial Magnetic Stimulation and electrical Peripheral Nerve Stimulation	<ul style="list-style-type: none"> Age 18-85 yrs Injury above L2 Can grip bilat Can ambulate a few steps Sub acute/Chronic SCI≥1 month 	Recruiting	514	Phase: Not Applicable, Primary Purpose: Diagnostic, Intervention Model: Crossover Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Motor Cortical Excitability EEG/EMG Enhanced motor UE Enhanced motor LE F/U 5 months	January 2021	May 18, 2015	May 25, 2021	Chicago, IL, USA
NCT0329652	Kevin Kilgore	Implantation and use of networked neuroprosthesis system (NNS) for arm, hand and trunk function.	<ul style="list-style-type: none"> Age≥17 yrs SCI C5-C7 AIS A, B, C, D Elbow flex≥2/5 Chronic SCI≥6m 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Treatment, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> ADL Abilities Test Grasp-Release Test F/U 3 months	December 2014	December 31, 2014	July 21, 2022	Cleveland, OH, USA
NCT01964261	University of Southern California Rancho Los Amigos	Implantation of 3 Neuroport electrode arrays to enable learned control of an end effector (for reach and grasp tasks) by thought augmented with sensory feedback via intracortical brain stimulation	<ul style="list-style-type: none"> Age 22-65 yrs High cervical SCI AIS NS Time after SCI NS 	Recruiting	2	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Patient control of end effector (virtual or physical) Infection or Irritation F/U 1 year	November 2013	October 17, 2013	June 1, 2022	Downey, CA, USA Los Angeles, CA, USA Pasadena, CA, USA
NCT01958086	Richard A. Andersen, PhD	Implantation of two Neuroport electrode arrays in posterior parietal cortex allowing direct brain-control of a computer interface. Ultimate objective is to allow the patient autonomous control over the Google Android tablet operating system.	<ul style="list-style-type: none"> Age 22-65 yrs high cervical SCI Lives<60 miles from study center; not on ventilator Time post SCI NS 	Recruiting	2	Phase: Not Applicable, Primary Purpose: Basic Science, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Subject control of tablet computer Absence of infection or irritation Adverse Events F/U 6 years	October 2013	October 8, 2013	April 13, 2022	Los Angeles, CA, USA Pomona, CA, USA
NCT01923662	VA Office of Research and Development	Device: IST-16 (16-Channel implanted stimulator-telemeter) for standing in persons with paralysis resulting from neurological disorder such as low cervical/thoracic spinal cord injuries (C6-T12)	<ul style="list-style-type: none"> Age≥21 yrs C6-T12 AIS NS Chronic SCI≥6m 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Device easibility, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Elapsed Standing Time Subject Impression of Stability Body Weight Distribution Standing Stability Measures F/U 12 months	April 2013	August 15, 2013	December 13, 2022	Cleveland, OH, USA
NCT01894802	Michael Boninger	Implantation of microelectrode Cortical Recording and Stimulating (CRS) arrays in the motor cortex and sensory cortex of the brain for neural activity recording and use in control of external devices	<ul style="list-style-type: none"> Age 22-70 yrs Limited or no ability to use hands due to cervical SCI or other condition Chronic SCI≥1yr 	Recruiting	5	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	Arm/Hand Function	<ul style="list-style-type: none"> Safety Efficacy: long-term recording of neural activity and successful control of external devices F/U 12 months	December 2013	July 10, 2013	September 26, 2022	Pittsburgh, PA, USA
NCT01474148	VA Office of R&D	Device: IRS-8 (8-Channel implanted stimulator-telemeter) to facilitate stability of the trunk and hips; Study the effect of stabilizing and stiffening the trunk with FES to change the way persons with SCI sit, breathe, reach, push a wheelchair, roll in bed	<ul style="list-style-type: none"> Age≥21 yrs SCI C4-T12 AIS A, B, C Chronic SCI≥6m 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Treatment Intervention Model: Single Group Assignment Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> seated posture respiration reach ability seated stability personal mobility F/U up to 36 months	July 2011	November 18, 2011	January 6, 2023	Cleveland, OH, USA
NCT00912041	NIDCD US Dept of Veterans Affairs NINDS	Implantation of the one or two BrainGate2 sensor electrode arrays into the motor cortex; training implanted subjects to control a computer cursor and other assistive devices with their thoughts	<ul style="list-style-type: none"> Age 18-80 yrs Cervical SCI AIS A, B, C, D Lives3hr drive Time post SCI NS 	Recruiting	15	Phase: Not Applicable, Primary Purpose: Other, Intervention Model: Single Group Assignment, Masking: None (Open Label)	Technology	General Health	<ul style="list-style-type: none"> Safety Feasibility of BrainGate2 F/U 1 year	March 2009	July 3, 2009	January 10, 2023	Stanford, CA, USA Sacramento, CA, USA Boston, MA, USA Cleveland, OH, USA Providence, RI, USA Atlanta, GE, USA
NCT00623389	Case Western Reserve University NIH	Device: IST-16 (16-channel implanted stimulator-telemeter) with pre- and post- surgical training to facilitate exercise, standing, stepping and/or balance in people with various degrees of paralysis	<ul style="list-style-type: none"> Age 21-75 yrs C6-T12 or other paralysis AIS A, B, C Normal ROM Chronic SCI≥6m 	Recruiting	10	Phase: Not Applicable, Primary Purpose: Device Feasibility Intervention Model: Single Group Assugnment Masking: None (Open Label)	Technology	Standing/Walking/Mobility	<ul style="list-style-type: none"> Device reliability and technical performance Device operability F/U 12 months	June 2018	February 26, 2008	October 6, 2022	Cleveland, OH, USA

This table is abstracted from the clinical trial registration website www.clinicaltrials.gov using the search term "Spinal Cord Injuries" and is updated periodically. The most recent update occurred January 31, 2023 at which time the www.clinicaltrials.gov search found a total of 1,564 SCI trials. Of these, there were 358 interventional trials that are enrolling or not-yet-enrolling. Review of these 358 studies for those that are targeting improvement in neurological or related functional outcomes yielded the current list. The table includes 170 SCI trials from the search that: 1) are currently actively recruiting or soon-to-be recruiting subjects; 2) are interventional (testing an intervention/treatment) using rehabilitation, neural stimulation and/or assistive technology strategies; and 3) targeted sensorimotor neurological or related functional improvement of the spinal cord as outcome measures. Trials meeting these criteria are included if sufficient information is available on the [clinicaltrials.gov](http://www.clinicaltrials.gov) webpages to adequately determine basic protocol design, the nature of the intervention, its delivery method, and relevant outcome measures. Interventional clinical trials are routinely registered on www.clinicaltrials.gov based on legal requirements* and because scientific journals may require registration for publication of the trial results. The [clinicaltrials.gov](http://www.clinicaltrials.gov) website is the largest repository of current and past clinical trials for all diseases and disorders-as of January 31, 2023 the registry contained information on 440,875 trials including research conducted in all 50 states in the USA and 221 countries. Investigators may choose not to register some early phase trials and those testing behavioral interventions, even though they may be important and scientifically rigorous studies. *U.S. Public Law 110-85 requires the registration and reporting of results of "certain applicable clinical trials," i.e., controlled interventional clinical trials that are subject to FDA regulation and that involve a Drug or Biologic (other than Phase I investigations), or Device (other than small feasibility studies); <http://prsrinfo.clinicaltrials.gov/lda.html>. More detailed information on individual trials may be accessed by using the NCT number found in the first column of the table. All trials registered with www.clinicaltrials.gov are assigned a registration number that begins with NCT (e.g. NCT01321333).

When an electronic version of the tables is used (e.g. when downloaded as a pdf file from www.scope-sci.org), the webpages describing a specific trial can be directly accessed by using the hyperlink (left Click to follow the link) of the NCT number in the table. Listing of a clinical trial on the [clinicaltrials.gov](http://www.clinicaltrials.gov) website does not reflect an endorsement by SCOPE or the National Institutes of Health. Information appearing on [clinicaltrials.gov](http://www.clinicaltrials.gov) is provided by study sponsors/investigators and is not verified by SCOPE or [clinicaltrials.gov](http://www.clinicaltrials.gov) for scientific validity or relevance. Before volunteering to participate in a clinical trial, patients are urged to discuss all options with their healthcare provider and other trusted advisors.

NCT ID	Sponsor	Intervention	Criteria	Status	Target Enrollment	Study Phase & Design	Primary Intervention Type	Primary Potential Benefit	Outcome Measures and Follow Up	Study Start Date	First Posted	Last Updated	Location(s)
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Terms/Abbreviations

- AIH: Acute Intermittent Hypoxia. Short duration (<2 min) exposure to breathing reduced oxygen concentration levels (~10% inspired oxygen), with alternating exposures to breathing air with normal oxygen concentration (~21% inspired oxygen). This intervention is commonly delivered with a breathing mask device as a series of multiple brief hypoxic exposures alternating longer breathing exposure to "room air" with normal oxygen content.
- AIS: the ASIA (American Spinal Injury Association) Impairment Scale is a component of the ISNCSCI that classifies the degree of motor/sensory sparing below the level of injury. The AIS scale ranges from A (most severe, complete injury without sensory function in the lowest sacral segments) to E (normal). AIS B describes sensory only sparing; AIS C describes sensory and very weak motor sparing; AIS D describes sensory and stronger but not normal motor sparing.
- Ashworth/Modified Ashworth: a scale used to measure spasticity severity
- Barthel Index: a measure of performance in Activities of Daily Living (ADL) and Mobility
- Box and Block Test: a test of manual dexterity-how many blocks a person can grasp and transfer in one minute.
- Central Cord Syndrome/Cervical Central Cord Syndrome: motor incomplete cervical SCI in which the upper extremities are significantly more impaired than the lower extremities
- COPM: Canadian Occupational Performance Measure
- DASH: Disability of Arm, Shoulder, Hand scale is a measure of the upper extremity function
- EMG: the electromyogram refers to a physiological test of muscle and nerve function.
- ESWT: extracorporeal shock wave therapy. Delivery of sound wave energy to the spinal cord using a transducer applied to the skin (extracorporeal i.e. outside of the body).
- FIM: Functional Independence Measure was developed to measure the burden of care in persons who were not independent in ADL, hygiene/self-care, and mobility. The FIM and its subscales have been used as an outcome measure of a subject's independence.
- Frankel Scale: an older scale for classifying severity of injury that was modified in 1992 to create the AIS.
- FU: follow-up
- GRASSP: Graded Redefined Assessment of Strength, Sensibility, and Prehension is a clinical measurement of upper limb function for use in person with tetraplegia (quadriplegia).
- HDE: Humanitarian Device Exemption is a U.S. Food & Drug Administration (FDA) application that, if successful, authorizes the applicant to market a Humanitarian Use Device (HUD) subject to certain profit and use restrictions.
- HUD: Humanitarian Use Device is a designation of the U.S. FDA for medical devices intended to benefit patients in the treatment or diagnosis of a disease or condition that affects or is manifested in fewer than 4,000 persons in the United States per year. HUDs cannot be sold for profit, except in narrow circumstances, and they can only be used in a facility after an Institutional Review Board has approved their use in that facility except in certain emergencies.
- IANR-SCIFRS: the International Association of Neurorestoratology-Spinal Cord Injury Functional Rating Scale. Changes in motor and sensory scores assessed by IANR-SCIFRS scale (total score range from 0 to 51, higher values represent a better outcome)
- ICSSH: International Classification for Surgery of the Hand in Tetraplegia is a clinical measure of hand function used by surgeons performing reconstructive surgery to improve function in persons with tetraplegia.
- IRB: Institutional Review Board is a multidisciplinary group that has been formally designated by an institution such as a hospital to review, approve and monitor biomedical research involving human subjects.
- ISAFSCI: International Standards to document remaining Autonomic Function after Spinal Cord Injury.
- ISNCSCI: International Standards for Neurological Classification of Spinal Cord Injury-sometimes referred to as the ASIA (American Spinal Injury Association) standards. This refers to the accepted international standards for performing motor/sensory physical examination of persons with SCI and the classification scheme for documenting the neurological level and the severity (completeness) of injury.
- IT: intrathecal, within the subarachnoid space surrounding the spinal cord-e.g. administration of a drug into the subarachnoid space which contains the cerebrospinal fluid (CSF)
- IV: intravenous-administration of a drug by vein
- Kinematics: analysis of movement
- Kunming Locomotor Scale: a 10-grade Roman numeral locomotion scoring system describing ability to stand, ability to walk, and required support/devices.
- N/A: not applicable
- NS: not specified
- Penn Spasm Frequency Scale: a measure of spasticity based on frequency of spasm occurrence
- Phase of Study: Clinical trials usually progress in phases from 1 to 4. (Note: trials that are not on the path to FDA/regulatory approval (e.g. trials of surgical techniques or rehabilitation therapies) may not have a phase designation.)
 - 1 Phase 1 trials are usually first-in-human or first-in-disease/condition experiments that are intended to demonstrate feasibility (can it be done), safety (is it reasonably safe), and tolerability (are the side effects tolerable). These trials usually do not include a comparison control group, and do not provide direct evidence of the interventions efficacy. Phase 1 trials usually enroll a small number of subjects and are commonly done at a single study center but may use a small number of collaborating centers.
 - 2 Phase 2 trials follow successful completion of Phase 1. Phase 2 trials are used to develop information on intervention administration (how to give), dose (how much to give), timing (when and how long to give), effect of the intervention on the body (what does it do, beneficial or harmful). These trials commonly utilize multiple study centers, many subjects, and a randomized control group to provide direct information about efficacy and safety of the intervention. Phase 2 trials enable refinement of how to administer the intervention and how to measure its beneficial effects.
 - 3 Phase 3 trials are conducted using the refined protocols developed from Phase 2 trials. They are often termed "pivotal" studies because they are sufficiently well-designed and rigorously conducted that their results, if positive, can be used for regulatory approval (e.g. FDA approval). These trials almost always enroll large numbers of subjects (hundreds or more), use multiple study centers, and a randomized control group design (placebo control and double blinding). The FDA generally requires two successful confirmatory Phase 3 trials of an intervention for approve
 - 4 Phase 4 trials are conducted after regulatory (e.g. FDA) approval to gather additional safety and efficacy data.
- Pharmacokinetics: study of the absorption, distribution, metabolism, and excretion of drugs; commonly involving periodic sampling of bodily fluids (e.g. blood, serum, CSF, urine) to determine drug concentration changes over time.
- Open Label: a trial in which there is no attempt to conceal the identity of the intervention from the subjects; i.e. there is no "blinding" or "masking" of the intervention-the subjects know that they are receiving either an "active ingredient" or a placebo.
- RCT: Randomized Controlled Trial. A clinical trial in which subjects are randomly assigned to either receive the active treatment or an alternative (control). Well-designed RCT's minimize the influence of variables other than the intervention that might have an effect on the desired outcome. For this reason, they provide the best evidence of efficacy and safety. The most rigorous RCT's utilize a placebo (inactive) control group and blinding (concealing active vs. control assignment) to minimize bias in the interpretation of study results.
- Residual Urine: Changes in residual urine measured after voiding by ultrasound test (volume of urine in mL, lower values represent a better outcome)
- ROM: Range of Motion
- SCIM/SCIM II/SCIM III: the Spinal Cord Independence Measure is a measure of a person's ability to perform certain activities independently
- SQ: subcutaneous-administration of a drug by injection beneath the skin
- SF-36: the Short Form-36 is a patient-reported survey of health status. The SF-36 is commonly used as a measure of Health-Related Quality of Life
- TMS/UEMS/LEMS: Total Motor Score/Upper Extremity Motor Score/Lower Extremity Motor Score are components of the ISNCSCI that include the ASIA Motor Index Score (the TMS) and the sub-components of the UEMS and the LEMS which are commonly analyzed and reported separately.
- Tardieu Scale: test of spasticity by assessing muscle resistance to passive movement at both slow and fast speed.
- VAS: Visual Analogue Scale-a scale commonly used to assess the severity of pain
- 9 Hole Peg Test: a test of manual dexterity
- 6MWT: 6 minute walk test. An assessment of the distance that the subject can walk in 6 minutes.
- 10MWT: 10 meter walk test. An assessment of the time required to walk 10 meters.