REPUBLIC OF LEBANON MINISTRY OF PUBLIC HEALTH Lebanon Clinical Trials Registry

The Effect of Rhythmic Auditory Stimulation on Gait in Chronic Stroke Patients

11/09/2025 04:13:21

Main Information	
Primary registry identifying number	Protocol number
LBCTR2021024744	CEUA 056
MOH registration number	
Study registered at the country of origin	Study registered at the country of origin: Specify
Yes	
Type of registration	Type of registration: Justify
Prospective	N/A
Date of registration in national regulatory agency 20/01/2021	
Primary sponsor	Primary sponsor: Country of origin
Farah Ayoubi	Lebanon
Date of registration in primary registry	Date of registration in national regulatory agency
14/02/2021	20/01/2021
Public title	Acronym
The Effect of Rhythmic Auditory Stimulation on Gait in Chronic Stroke Patients	
Scientific title	Acronym
The Effect of Rhythmic Auditory Stimulation Tempo Shifts Application on Gait Parameters in Chronic Stroke Patients: A Pilot Study	
Brief summary of the study: English	
To examine the effect of rhythmic auditory stimulation tempo shifts on gait in stroke patients. Ten to twenty chronic stroke patients with an onset of six months and above will be recruited based on the proposed criteria and after a thorough assessment and obtaining their consent. Participants will be instructed to walk under a self selected tempo based on their average walking cadence followed by three randomly applied conditions (-10%, +10%, and +20%) calculated based on the self-selected tempo, then finally each participant will walk without stimulation to monitor immediate effect. Kinovea (motion analysis software) and 10 meter walk test will be used to evaluate gait parameters at session 1 and session 12 respectively through all applied walking conditions to monitor short term effect.	
Brief summary of the study: Arabic	
دى مرضى السكنة الدماغية. سيتم توظيف عشرة إلى عشرين مريضا بالسكنة المرّمنة (سنة أشهر وما يهم دقيق والحصول على موافقة المرضى سيتم توجيه المشاركين للسير وفقا للايقاع المحدد ذاتيا متبوعا المشي. ثم أخيرا سيمشى كل مشارك دون تحفيز ٢٠ + , ١٠. , ١٠ بنالاثة شروط وظيفية عشوائية ٪ في في Kinovea أمتار و برنامج ١٠ المراقبة تأثير العلاج الفوري. سيستخدم اختبار المشي على مسافة	فوق). بناءً على المعابير المُقترحة وبعد تقو + محددة استنادا إلى متوسط معدل سرعة

Health conditions/problem studied: Specify

 \sim

Health Condition: Chronic Stroke Patients (Hemiplegia)



Problems Studied: Gait (spatio-temporal parameters)/ Walking Issues/ Motor Issues/ Impairment/ Function

Interventions: Specify

Each participant will walk along a wide unobstructed 10-m walkway at preferred speed with the possibility to use a uniform walking aid when needed, based on this each participant's cadence will be calculated using a pedometer and then averaged over 3 trials to obtain the mean cadence. The acceleration and deceleration periods will be neglected, and this step is to be repeated each session. The self selected tempo will be set each session according to the mean cadence (number of steps/min= number of beats/min). Each participant will then walked under three randomly applied conditions which will be administered based on block randomization (-10%, +10%, and +20%) calculated from self-selected tempo (0%). Each participant will walk 3 times under baseline tempo (0%), followed by 3 times under each condition. Finally, each participant will be asked to walk for a distance of 10m without any stimulation to monitor short term effects. The obtained data will be analyzed using Kinovea software and 10-m walk test.

Conventional approach (ethical considerations): The conventional treatment will consist of customized functional tasks targeting transfers such as; weight shifting while in sitting and standing, transfers from sitting to standing, and from one chair to another (with and without arm support) as well as enhancing static balance; while sitting and standing in addition to management of the upper extremity; through stretching, range of motion exercises of high repetition, and functional reaching exercises.

Key inclusion and exclusion criteria: Inclusion criteria

The inclusion criteria were as follows: 1) able to walk for a 10-m distance. 2) Able to follow instructions. 3) Doesn't suffer from cognitive impairment with a Modified Mini Mental State Examination (3MS) score of 79 or higher. 4) Brunnstrom recovery scale of stages (4-6).

Key inclusion and exclusion criteria: Gender	Key inclusion and exclusion criteria: Specify gender
Both	
Key inclusion and exclusion criteria: Age minimum	Key inclusion and exclusion criteria: Age maximum
35	65

Key inclusion and exclusion criteria: Exclusion criteria

Individuals were excluded if any demonstrated one or more of the following conditions: 1) Auditory or visual problems. 2) Chronic pain and/or an orthopedic condition which affects gait. 3) Symptomatic cardiac failure. 4) Neurologic condition other than initial stroke. 5) Major depression or psychological issues.

Type of study Interventional	
Type of intervention	Type of intervention: Specify type
Rehabilitation strategies	N/A
Trial scope	Trial scope: Specify scope
Therapy	N/A
Study design: Allocation	Study design: Masking
Single Arm Study	Blinded (masking used)
Study design: Control	Study phase
Active	N/A
Study design: Purpose	Study design: Specify purpose
Treatment	N/A
Study design: Assignment Single	Study design: Specify assignment N/A
IMP has market authorization	IMP has market authorization: Specify
Name of IMP	Year of authorization Month of authorization
Type of IMP	



REPUBLIC OF LEBANON Lebanon Clinical Trials Registry

Pharmaceutical class

N/A

Therapeutic indication

Such approaches are essential and crucial for stroke individuals which are hemiplegic or have a functional impairment in their lower extremities or have a motor control issues in addition to hemiparesis. Additionally such an approach is very beneficial for other neurologic populations as a primary treatment method.

Therapeutic benefit

- Optimizing Gait and Gait parameters (deviations, symmetry, cadence, velocity, step length, etc...)
- Improving Balance (Static+Dynamic)
- Decreasing postural issues and optimizing postural control
- Motor Relearning
- Inducing and Optimizing Brain plasticity
- Optimizing quality of life
- Optimizing independence
- Optimizing functional capacities
- Decreasing morbidity rates
- Decreasing mortality rates
- Prevention of recurrence of health issues
- Improving Mental state and cognition

Study model

N/A

Study model: Explain model N/A

Time perspective: Explain time perspective

Study model: Specify model

N/A

Time perspective

N/A

Time perspective: Specify perspective

N/A

Target follow-up duration

Number of groups/cohorts

Biospecimen retention

None retained

Biospecimen description N/A

Target follow-up duration: Unit

N/A

Target sample size

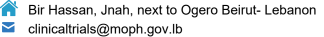
30

Date of first enrollment: Type
Actual

Date of study closure: Type Actual Actual enrollment target size 10

Date of first enrollment: Date 04/01/2021

Date of study closure: Date 04/03/2021





Recruitment status	Recruitment status: Specify
Recruiting	
Date of completion	
14/02/2021	
IPD sharing statement plan	IPD sharing statement description
Νο	At this current point we would prefer to keep our data disclosed, as soon as we are done with the study as eligible we will be submitting the trial to a journal for a prospect publication.

Additional data URL

Admin comments

Trial status

Approved

Secondary Identifying Numbers		
Full name of issuing authority	Secondary identifying number	
Ethics Committee at the Antonine University (CEUA)	CEUA 056	

Sources of Monetary or Material Support

No Sources

Secondary Sponsors
Name
NA



Contac	ct for Public/Scientific Queries					
Contact type	Contact full name	Address	Country	Telephone	Email	Affiliation
Public	Farah AYOUBI	Tripoli	Lebanon	71765367	farah.ayoubi@ua .edu.lb	Research Advisor/ Supervisor / Research Instructor/ MPT/ PhD/ PT at Antonine University
Scientific	Marie Catherine Baradhii	Baabda	Lebanon	70686738	mariecatherine2 @hot mail.com	Researche r/ Research Supervisor / Clinical Instructor/ DPT/ MPT/ PT

Centers/Hospitals Involved in the Study			
enter/Hospital name		Principles investigator speciality	Ethical approval
Arc en ciel (Jesr I Wate)	ate) Dr. Jeanine Matar (yet, researcher conducted trial)		Approved
Mousawat Foundation	N/A (researcher conducted trial)	PT	Approved

Ethics Review				
Ethics approval obtained Approval date Contact name C		Contact email	Contact phone	
Other Ethics committe of the Antonine University (CEUA)	26/11/2020	Nidaa Abou Mrad	nidaa.aboumrad@ua.edu.lb	05927000

Countries of Recruitment

Name

Lebanon

Health Conditions or Problems Studied		
Condition	Code	Keyword
Stroke/ Hemiplegia	Hemiplegia (G81)	stroke/ hemi/ gait/ function/ RAS





Interventions		
Intervention	Description	Keyword
Gait Re-education under Rhythmic Auditory Stimulation	Rhythmic Auditory Stimulation is a neurologic technique used to facilitate the rehabilitation of movements that are intrinsically biologically rhythmical, most importantly gait. RAS uses the physiological effects of auditory rhythm on the motor system to improve the control of movement in rehabilitation of functional, stable, and adaptive gait patterns in patients with significant gait deficits due to neurologic impairment.	RAS

Primary Outcomes

Name	Time Points	Measure
Angle measurements	At session 1 and session 12	Angles in degrees via Kinovea software
Angle symmetry	At session 1 and session 12	SI (symmetry index)
Velocity	At session 1 and session 12	10m walk test
Cadence	At session 1 and session 12	Number of steps/min using pedometer
Step Length	At session 1 and session 12	Steps in cm using kinovea software

Key Secondary Outcomes		
Name	Time Points	Measure
Cognition (pre-requisite to enrollment)	At session 1 (pre- experiment)	3MS (Modified Mini Mental State examination)
Muscular strength/ Motor ability (global)	At session 1 (pre- experiment)	Brunnstrom scale





Trial Results Summary results Study results globally Date of posting of results summaries Date of posting of results summaries Date of first journal publication of results Results URL link Baseline characteristics Participant flow Adverse events Outcome measures URL to protocol files