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Feasibility of using Lokomat combined with FES for the rehabilitation of foot drop

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Abstract—This study investigated the clinical feasibility of combining the electromechanical gait trainer Lokomat with functional electrical therapy (LokoFET), stimulating the common peroneal nerve during the swing phase of the gait cycle to correct foot drop as an integrated part of gait therapy.

Five patients with different acquired brain injuries trained with LokoFET 2-3 times a week for 3-4 weeks. Pre- and post-intervention evaluations were performed to quantify neurophysiological changes related to the patients' foot drop impairment during the swing phase of the gait cycle. A semi-structured interview was used to investigate the therapists' acceptance of LokoFET in clinical practice. The patients showed a significant increase in the level of activation of the tibialis anterior muscle and the maximal dorsiflexion during the swing phase, when comparing the pre- and post-intervention evaluations. This showed an improvement of function related to the foot drop impairment. The interview revealed that the therapists perceived the combined system as a useful tool in the rehabilitation of gait. However, lack of muscle selectivity relating to the FES element of LokoFET was assessed to be critical for acceptance in clinical practice.

I. INTRODUCTION

Over a third of the patients suffering from acquired brain injury (ABI) have gait impairments [1], [2]. An inadequate control of dorsiflexion during the swing phase of gait (called foot drop) is often seen [3]. Foot drop is often caused by weakness of the tibialis anterior (TA) muscle [4], which can result in compensatory movement patterns [3], slowed gait velocity [5], limited functional mobility, and increased risk of falls [4].

Electromechanical gait trainers (EMGT) such as Lokomat (Hocoma AG, Volketwill, Switzerland) are often utilized to initiate early intensive gait rehabilitation [6]. Lokomat enables the patient to train gait movements with many repetitions, more independently of the therapist, potentially with their body weight supported, and with an automated movement of the lower extremities [6].

Stroke patients subjected to EMGT therapy (using Lokomat) combined with conventional therapy develop larger muscle mass and lower fat percentage compared to controls only receiving conventional therapy [7]. Despite the beneficial effects of this EMGT therapy, Lokomat has shown to inhibit dorsiflexion of the ankle joint during the swing phase of the gait cycle, rather than facilitating it [8]. This conflicts with the guidelines for rehabilitation of pathological gait, which advocate that rehabilitation should facilitate a physiological gait pattern, hereby supporting restorative neurorehabilitation [3].

Functional electrical stimulation (FES) administered at the common peroneal nerve has shown to facilitate dorsiflexion of the ankle joint during the swing phase of the gait cycle [9], and there is clinical evidence supporting FES having a therapeutic effect on the foot drop impairment [10].

The combination of EMGT and FES for rehabilitation of gait impairments in sub-acute stroke patients has resulted in increased gait speed compared to conventional therapy [11]. However, only two studies have combined FES with Lokomat, showing that the combined system was technically feasible [12] and that when tested by one stroke patient, the patient was able to achieve better dorsiflexion and TA muscle activation during the swing phase of the gait cycle, in comparison to the normal gait pattern [13].

The goal of this feasibility study was therefore to test a clinical protocol combining Lokomat with Functional Electrical Therapy (FET), i.e. LokoFET, as preparation for larger clinical trials. The aims of the study were to:

- quantify the neurophysiological changes related to the patient’s foot drop impairment after LokoFET treatment
- investigate the therapists’ acceptance of LokoFET in clinical practice

II. METHODS

A. Test of LokoFET in clinical practice

Subjects

Five patients were recruited [Table I] based on the following inclusion criteria: suffering from ABI, age between 18-80 years, weight 50-100kg, femur length 48-21cm, able to fully extend the knees, able to communicate and understand instructions, tolerate electric stimulation, tolerant to be supported passively in the body weight support (BWS) harness for 5 min, decreased dorsiflexion during the swing phase, and able to walk a minimum of 30 steps with the ankle strap from Lokomat in neutral position. Patients with the following criteria were excluded: height over 2 m, leg length difference > 2 cm, bone instability, infection and/or orthopedic problems in the placement area of the electrodes, heart or lung disease, pregnancy, pacemaker, prior incidences of neurological or musculoskeletal diseases, suffering from mental diseases, and lack of cooperation.
corresponded to heel-strike of the contralateral leg and the output signals from the Lokomat. The start signal duration of the stimulation period was controlled by the FES was triggered using custom made software [13]. The done in each session, based on the evoked motor response.

At each session, the Lokomat exoskeleton, gait speed, and BWS were adjusted to fit the patient. The ankle straps were adjusted to have the patient’s ankle joint in neutral position. When the gait training was initiated, Lokomat supplied timing information, defining the start- and end of the stance phases.

**Functional electrical stimulation**

To stimulate the common peroneal nerve, a one-channel, computer-controlled stimulator (Noxitest, Danmark) and two surface electrodes were used. The cathode (Pals Platinum Round 3.2 cm, Axelgaard Ltd., USA) was placed above the common peroneal nerve, close to the fibular head on the patient’s most affected leg. The anode (Pals Platinum Oval 4.0x6.4 cm, Axelgaard Ltd., USA) was placed on the proximal aspect of the TA muscle on the same leg. The rectangular monophasic stimulation had a pulse duration of 300 µs and a frequency of 30 Hz. Adjustments regarding the location of the electrodes and the stimulation intensity were done in each session, based on the evoked motor response. FES was triggered using custom made software [13]. The duration of the stimulation period was controlled by the output signals from the Lokomat. The start signal corresponded to heel-strike of the contralateral leg and the end signal corresponded to the beginning of the stance phase of the most affected leg [13]. Consequently, the patient’s most affected side was stimulated during the push-off and swing phases of the gait cycle.

**Outcome measurements**

Electromyography (EMG) was recorded using a bipolar configuration. Two electrodes (Medicotest, Oelstykke, Denmark) were placed on the muscle belly of TA on the patient’s most affected leg and a third electrode of the same kind was placed on the tibial bone serving as reference. The recordings were amplified, based on the patient’s individual maximum voluntary contraction, band-pass filtered (10-500 Hz, second order), sampled at 2 kHz, and saved.

Kinematics of the ankle joint was recorded in the sagittal plane using an electronic goniometer (SG110/A, Biometrics Ltd., Gwent, UK). The goniometer was placed on the lateral side of the ankle of the most affected leg. Data was sampled at 2 kHz and saved.

The protocol for the study was approved by the local ethical committee (ESDH 1-10-72-135-12) and the experiments were conducted in accordance with the declaration of Helsinki.

**Lokomat**

Lokomat was used to provide gait training. Prior to the first experimental session, the patients had at least one training session in Lokomat to familiarize them to the EMGT.

When the gait training was initiated, Lokomat supplied timing information, defining the start- and end of the stance phases.

**Pre- and post-intervention evaluation**

Pre- and post-intervention evaluations were performed to assess the effect of training. The goniometer and EMG electrodes were mounted while the patients were seated. The patients then walked in Lokomat for 2-5 minutes before recording the ankle kinematics and TA EMG during 30 steps.

**LokoFET training**

Patients had one hour set aside per session. First, the FES stimulation intensity was determined starting at 10mA and adding steps of 2mA until adequate dorsiflexion was observed (≥10°) with the patient in sitting position. The Lokomat’s BWS system, exoskeleton, and treadmill were then adjusted. Immediately afterwards, the patients walked assisted by Lokomat and, if necessary, the stimulation intensity was adjusted until appropriate dorsiflexion was visually confirmed, or the stimulation intensity became intolerable.

The patients received LokoFET training 2-3 times a week for 3-4 weeks, giving a total of 6-8 treatments.

**Data analysis**

The time spent on preparing for FES (e.g. placement of electrodes and adjusting stimulation intensity) before the actual training started was noted. The time spent on preparing Lokomat’s BWS and exoskeleton, and setting the speed of the treadmill was likewise noted.

**Acceptance of LokoFET in clinical practice**

To investigate the acceptance of LokoFET in clinical practice a semi-structured interview was performed and data was saved as transcripts [14]. The respondents for the semi-structured interview were two of the physiotherapists who administered LokoFET.
Data analysis
The analysis of the qualitative data was done using a conventional qualitative content analysis [15]. After reading the transcripts, the material was coded allowing for emerging themes to appear as sub-themes, resulting in a meaning condensation, and interpretation of the qualitative data [15].

III. RESULTS AND FINDINGS

A. Test of LokoFET in clinical practice
Out of the five patients originally included in the study, four patients completed the LokoFET training course. Patient 5 was excluded due to low motor ability and low cognitive stamina making him unable to complete the LokoFET training within the one-hour limit per session.

The mean stimulation intensity across patients was 34.98±8.25 mA. The mean duration of the sessions was 17.12±6.34 minutes. 10.35±3.83 and 35.91±6.00 minutes were spent for FES and Lokomat preparation, respectively.

Effect of LokoFET training
The TA EMG activity was significantly increased (73.34%) at the post-intervention evaluation, compared to the pre-intervention evaluation (paired t-test, p<0.05) (Fig. 1).

The maximal dorsiflexion during the swing phase was significantly increased (66.78%) at the post-intervention evaluation, compared to the pre-intervention evaluation (paired t-test, p<0.02) (Fig. 2).

The maximal plantarflexion during push-off was not changed at the post-intervention evaluation, compared to the pre-intervention evaluation (paired t-test, p=0.67) (Fig. 3).

B. Acceptance of LokoFET in clinical practice
The therapists mentioned the FES element as a positive and active add-on treatment technique to the conventional EMGT therapy. However, it was also observed that in some LokoFET training sessions, FES resulted in dorsiflexion coupled with eversion of the ankle joint.

Selective dorsiflexion was assessed by the therapists to be crucial for the acceptance of LokoFET in clinical practice. This is because the coupled dorsiflexion and eversion is not a part of the non-pathological gait pattern, which the therapists strive towards avoiding during gait training, especially during the sub-acute phase. Training in the application and use of FES for foot drop correction were therefore assessed to be crucial by the therapists.

Time spent on FES preparation prolonged the usual time spent on preparation before training with Lokomat. The therapists argued that this could potentially harm the usefulness of LokoFET because many patients suffer from fatigue, hereby requiring a short preparation time in order to train before fatigue sets in.

The therapists also argued that the mere use of Lokomat in the standard EMGT therapy (without FES) was highly complex on its own, especially when training heavily motor impaired patients. The introduction of FES could therefore add a level of complexity making them unable to use both Lokomat and FES.

IV. DISCUSSION

A. Test of LokoFET in clinical practice
It was feasible for patients with different ABI to complete the LokoFET training in a clinical environment. The British Medical Research Council highlights the importance of feasibility studies evaluating the practical feasibility when introducing new interventions in everyday practice before testing them in larger clinical trials [16]. This study did not aim to show the effectiveness of LokoFET compared to conventional EMGT therapy or spontaneous remission. However, when comparing the pre- and post-intervention evaluation, both the TA EMG activity and the ankle joint kinematics (dorsiflexion) showed significant gains related to the foot drop impairment.

In this study, the stimulation intensity was adjusted until appropriate dorsiflexion was visually confirmed, or the stimulation intensity became intolerable. Alternatively, the adjustment of the stimulation intensity could have been based on more precise kinematic information using electronic goniometers or other sensors external to the Lokomat. However, one of the goals of this FES add-on
The undesired eversion could be caused by the higher stimulation intensities, which affect larger areas, depolarizing a larger amount of fibers of the superficial peroneal nerve, leading to increased activation of the peroneal muscle, which everts the ankle joint. Patients needing higher stimulation intensities (e.g. patients with severe foot drop, primarily caused by spasticity of the calf muscles) might therefore not be candidates for training with LokoFET or FES in general. Unfortunately, this study did not assess the spasticity or the passive mechanical resistance of the ankle joint of the patients before inclusion and therefore, no relation between the degree of foot-drop, the level of spasticity, and the stimulation amplitude could be made.

The therapists also mentioned that time spent on preparation instead of training could result problematic for some patients who can easily get fatigued. Post-stroke fatigue is a common stroke symptom [22]. Using the abovementioned electrode arrays, might help reducing the time needed for preparation. However, the preparation time for Lokomat was far greater than that for FES, whereby the patient was placed in sitting position for the latter preparation procedure. A possible reduction of the Lokomat preparation time might therefore be more beneficial.

The introduction of the FES element could result problematic for the therapists, especially when working with heavily motor impaired patients, due to the added complexity of operating both Lokomat and the FES system. In this study, FES was administered by a FES specialist and not by the therapists administering the EMGT therapy. Whether or not the added complexity of FES is a real problem is therefore unknown.

Training of personnel in both the application of FES and EMGT therapy is therefore important in order to secure optimal muscle selectivity and time efficiency. Furthermore, care should be taken to include eligible patients based on their level of spasticity and fatigue.

V. CONCLUSION

This feasibility study showed that LokoFET was able to provide ABI patients with active training of their ankle joint and resulted in improvements related to their foot drop impairment. However, the effectiveness of LokoFET compared to conventional EMGT therapy is still unknown.

Therapists perceived LokoFET as a useful tool in the rehabilitation of gait, pointing towards possible acceptance of the technology. However, muscle selectivity and the added complexity related to the FES element of the treatment were assessed to be critical for LokoFET to be useful in clinical practice.

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