Neuromotor Strategy of Gait Rehabilitation for Lower-Limb Spasticity

Jinan Charafeddine, Sylvain Chevallier, Mohamad Khalil, Didier Pradon, Samer Alfayad

To cite this version:
Jinan Charafeddine, Sylvain Chevallier, Mohamad Khalil, Didier Pradon, Samer Alfayad. Neuromotor Strategy of Gait Rehabilitation for Lower-Limb Spasticity. International Conference on Advances in Biomedical Engineering (ICABME), Oct 2019, Tripoli, Lebanon. 10.1109/ICABME47164.2019.8940263. hal-02541681

HAL Id: hal-02541681
https://hal.uvsq.fr/hal-02541681
Submitted on 14 Apr 2020

HAL is a multi-disciplinary open access archive for the deposit and dissemination of scientific research documents, whether they are published or not. The documents may come from teaching and research institutions in France or abroad, or from public or private research centers.

L’archive ouverte pluridisciplinaire HAL, est destinée au dépôt et à la diffusion de documents scientifiques de niveau recherche, publiés ou non, émanant des établissements d’enseignement et de recherche français ou étrangers, des laboratoires publics ou privés.
Neuromotor Strategy of Gait Rehabilitation for Lower-Limb Spasticity

Jinan Charafeddine¹, Sylvain Chevallier², Mohamad Khali³, Didier Pradon¹, Samer Alfayad²

1- EndiCAP U1179 - APHP – UVSQ
Hôpital Raymond Poincare, Garches, France

2- Laboratoire d’Ingenierie des Systemes de Versailles Universite de Versailles Saint-Quentin, France

3- Faculty of engineering, lebanese university, Tripoli, Lebanon

Abstract— Assisting users and restoring human locomotion for patients with lower limb spasticity is a challenging task. Studies focusing on patients with abnormal walking behavior are scarce because there is an important variability from one patient to another. Those patients could benefit the most from rehabilitation and assistive mechatronic devices, there is no generic controlling scheme or any dynamical gain indicator. This contribution introduces a bio-kinematic index which is called Neuro-motor index (NMI), based on electromyographic (EMG) and joint angles measurements. NMI is derived from the nonlinear regression with a combination of two co-contraction indices (CCI), which allows addressing the variability of walking situations. This new index is evaluated on patients with cerebral palsy and a stroke. Then, the estimation error was calculated in comparison with the other co-contraction indices. This estimation shows that this index has the highest signification of joint angles prediction. Thus it can be suitable in adaptive rehabilitation control for spasticity cases.

Keywords— Rehabilitation Exoskeleton, Spasticity, Bio-kinematic, co-contraction, Neuro-motor.

I. INTRODUCTION

The neurological diseases, such as Cerebral Palsy (CP) and strokes (Cerebral Accident, CA), may lead to lower-limb spasticity which conducts to gait disorders. CP is a disorder that affects muscle tone and motor skills, inducing uncoordinated movements [1]. For CA, this disease causes death or spasticity due to low blood flow to brain cells [2]. Spasticity is an exaggerated contraction of the antagonist muscle, which generates an opposition couple or even an inversion of the desired movement [3]. This could affect all the walk phases which are first double support (DS1), single support (SS), second double support (DS2) and swing phase (SWP) that are detailed on Fig. 1. Mechatronic devices such as exoskeletons are one of the solutions for the rehabilitation process. However, they are difficult to control as each patient has specific motor capabilities and physical integrity should be ensured. Control strategies rely on the fundamental assumption that the system should intervene if the patient is moving along a trajectory considered as “correct”. When deviating from this reference trajectory, the exoskeleton should generate a compensation force, through an adjustment of the mechanical impedance in the controller [4]. This kind of correction includes adjusting the walking speed or step length. There are two known control strategies for these devices (i) Interaction force controllers using impedance or admittance, which are generally predetermined and do not consider the user's physical condition [5,6] (ii) Musculoskeletal model based on EMG signals which rely on the detection of muscle activation. But EMG signals are not effective while applying on patients having a muscular disorder[5,6]. Consequently, the aforementioned strategies are inappropriate for patients with cerebral palsy (CP) or after stroke. The EMG signals not only contain data about patient intention but also give information about the muscle co-contraction around the joint [7]. Co-contraction is an operation between a pair of muscles (agonist/antagonist). The quadriceps (contains rectus femoral muscle (RF)) and hamstring (contains biceps femoral muscle (BF)) are bi-articular muscles for knee and hip joints. The angles of these joints result while movement due to the exchange of muscles roles between agonist and antagonist. Co-contraction is related to joint stability [8] and is an important factor which participates to the inefficiency of pathological movement [9].

Fig. 1: Illustration of the walk phase: First double support (DS1), single support (SS), second double support (DS2) and swing phase (SWP).

The most three important approaches to estimate this co-contraction are proposed by Falconer and Winter [9], Hessee et al. [10] and Unihan et al. [11]. These approaches are efficient for offline analysis and diagnosis but they are not appropriate for designing a control strategy. Consequently, a new index is needed which depends on physiological signals (Electromyography EMG) to estimate joint angles of walking. The strategy is to use the EMG as a proxy for determining the patient’s intention, initiating assistance when the muscular contraction is characteristic of a predefined move and detected with sufficient intensity. This approach stimulates movements initiated by the patient, which is an essential part of motor rehabilitation. The main objective of this work is to propose an approach suitable for people with
cerebral palsy or strokes and can be used in the future for exoskeleton control. Where the patient is considered as an expert while moving. The contributions are the following:

- A complete and uniform description for the most important co-contraction indices existing in the literature.
- A novel index proposition to characterize the relation between electromyographic activity and kinematic parameters.
- An evaluation on bio-kinematic recordings from 20 subjects (9 healthy, 6 strokes and 5 with cerebral palsy).

Section II presents the materials and methods. Section III details the results by evaluating the proposed index. Section IV concludes this paper.

II. MATERIALS AND METHODS

There were twenty subjects (9 healthy adults: 4 females and 5 males, aged 50 years±7; five kids with cerebral palsy: 3 females and 2 males, aged 10 years±2; 6 adults with stroke: 3 females and 3 males, aged 50 years±7), were recorded in Gait Laboratory (Raymond Poincare Hospital, arches, France).

A. Experimental protocol

The normal gait for each subject was tested on the ground by performing 11 trials. The activity of quadriceps and hamstring muscles were recorded using a surface EMG system (MA311). Using a 3D optoelectronic system, the gait analysis was recorded at 100 Hz following the Helen Hayes model which is commonly used by the biomechanical community for gait analysis, a group of markers were placed on the lower part of the patient [12]. The marker trajectories were then filtered using a Butterworth filter (4th order, 6 Hz cut-off frequency) [13]. The peak of flexion and extension was the main kinematic parameters as appropriate for the hips and knees. For both lower limbs: a calculation was made for spatiotemporal parameters (velocity, cadence, step and stride length, step width) and for kinematic parameters on each sub-phase of the gait cycle.

B. Signal Processing

The processing of the raw EMG signals was divided into several steps: first were band-pass filtered (10–400 Hz), second were rectified, third were low-pass filtered using Butterworth filter (4th order) from 4 till 6 Hz cut-off frequency depends on the subject cadence [14], and finally normalized by the detected maximal voluntary contraction value (MVC) of each muscle on gait trials [4]. For the kinematic data, the processing was begin by filtering using a Butterworth filter (4th order, 6 Hz cut-off frequency) which is connected to the acquisition system, then these data is segmented in 1001 values for knee and hip joints to obtain matrices with fixed dimensions which are equal to those of recorded EMG matrices. After that, the estimation of their mean and variance was done.

C. Co-contractions Indices

In the case of a normal walk, there is a very large literature on the CCI, which depends on numerous parameters such as the walking speed, inertia, considered muscle groups, age and sex [11]. The CCIs are always computed as a ratio between agonist and antagonist muscles, and as a function of the phase in the walking cycle. Among the various methodologies to compute CCI, three are more robust and accurate and will be detailed hereafter.

The CCI(t) is introduced by [9] and is computed as:

$$CCI(t)=\frac{\int_{t_1}^{t_2} ENV_{Ag}^{emg}(t) \cap ENV_{Ag}^{emg}(t) dt}{\int_{t_1}^{t_2} ENV_{Ag}^{emg}(t) \cup ENV_{Ag}^{emg}(t) dt} \times 100 \quad (1)$$

Where ENV(t) is the normalized envelop of EMG signals, as shown on Fig. 3. The period [t1, t2] denotes the time span where the envelop of Rectus Femoral (RF) is lower than the envelop of Biceps Femoral (BF), whereas [t3, t4] denote the period where the BF envelop is lower than the RF envelop. CCI(t) is computed by finding the area of the total antagonistic activity (area under RF + BF) divided by the integral of the sum of the EMG envelop (area under RF + BF). This method is commonly used in the literature, for quantifying the co-activation of muscle groups during multitjoint movements, such as gait. However, it exhibits discontinuities in two points of the cycle, between the single support (SS) and the second double support (DS2) on the one hand and, on the other hand, between the

The CCI2(t) is proposed by [10] and is defined as:

$$CCI_2(t)=\frac{2 \int_{t_1}^{t_2} ENV_{Ag}^{emg}(t) \cap ENV_{Ag}^{emg}(t) dt}{\int_{t_1}^{t_2} ENV_{Ag}^{emg}(t) \cup ENV_{Ag}^{emg}(t) dt} \times 100 \quad (2)$$

This method derived by the equation proposed by [15] and [9]. The CCI2(t) is evaluated by finding the overlap between the agonist and antagonist muscle (area under RF ∩ BF) curves and divided by the integral of the sum of the EMG envelop (area under RF + BF). On could note that CCI2(t) is continuous throughout a complete gait cycle.

$$CCI_3(t)=\frac{2 \int_{t_1}^{t_2} ENV_{Ag}^{emg}(t) dt + \int_{t_1}^{t_2} ENV_{Ag}^{emg}(t) dt}{\int_{t_1}^{t_2} ENV_{Ag}^{emg}(t) \cup ENV_{Ag}^{emg}(t) dt} \times 100 \quad (3)$$

The last co-contraction index is a reformulation of the index proposed by [15] and [11]. CCI3(t) is evaluated by finding the overlap between the agonist and antagonist (area under RF∩BF) curves and normalized by the area under RF∪BF.

![Fig. 2 Examples of muscles study, biceps femoral (BF) and rectus femoral (RF), conducted on one subject. a) normalized EMGs of the agonist and antagonist muscles. b) normalized EMGs of the antagonist and the total](image-url)
As CCI$_2(t)$, CCI$_3(t)$ is also continuous during the gait cycle. Nonetheless, CCI$_2(t)$ and CCI$_3(t)$ relies on the arbitrary definition of the peak co-contractions, that is the precise value of $t_1$, $t_2$ and $t_3$ which could be set priori for normal walk but is not suitable for abnormal walk without manual annotations of the data. They are thus not appropriate for the control of rehabilitation or assistive exoskeleton.

D. Proposed Index

Even with this ensured continuity for CCI$_2(t)$ and CCI$_3(t)$, the gait angles and the CCI varies independently along the walk cycle. It is difficult to find a mapping from the joint angle variation to the CCI, thus to propose a mapping function for the control process. We address this problem by defining a regression from the CCI to the angle variation of the considered articulation, that preserve the necessary information and act as a smoothing function. We use a Hermite polynomial to combine the CCI that are estimated from the EMGs recorded from the quadriceps and the hamstrings muscles. The proposed neuro-motor index is built on the carefull detection of user-specific peak co-contraction activities which are then injected in non-linear combination of two cocation indices [10],[16]. The NMI allows to characterize the envelop peak of a muscle pair during each flexion/extension of a walk cycle, and to estimate the joint angular trajectory. The first step is to detect the peak of $f(t)$ defined as:

$$\text{NMI}(t) = \frac{1}{2} \left\{ \text{CCI}_1(t) + R_x(t) \text{CCI}_2(t) \right\}$$  \hspace{1cm} (4)

Where, $R_x(t)$ is a nonlinear regression based on Hermitian function

$$R_x(t) = h_1(t)f_0 + h_2(t)p_0 + h_3(t)f_1 + h_4(t)p_1$$

The peaks are identified with: $\text{argmin } |f'(t)|$

$$f(t) = EMG_{anago}(t) \cap EMG_{ago}(t)$$

$h_1, h_2, h_3, h_4 \in He(t)$, $p_0$ and $p_1$ are tangent to $f_0$ and $f_1$.

The NMI is determined based on the flexion/extension of a joint and does not depend on the definition of a standard gait cycle, a requirement for working with cerebral palsy and stroke patients. The non-linear regression ensures that this method could be applied for an abnormal walk, either for adult or children. Starting from a minimal calibration to reduce the patient fatigue, in order to provide comfortable, stable and secure rehabilitation for the patient, the NMI allows a direct prediction of the patient intention because it depends on his EMG, and since it is derived from the co-contraction index, its results rely on the residual capabilities of the patient.

III. RESULTATS

The evaluation is conducted on all 20 subjects.CCI$_1(t)$, CCI$_2(t)$, CCI$_3(t)$ and NMI(t) are calibrated for one walking cycle of each subject and evaluated on the 10 other cycles.

Each index allows to estimate the angular value of the knee, and an estimation error is computed using kinematics data as ground truth. To proceed this evaluation, we manually annotate the dataset to indicate the $t_1$, $t_2$ and $t_3$ time values corresponding to each phase of the walking cycle (first double support, single support, second double support, and swing phase). Healthy subjects that have a normal walking cycle do not present any change. However this provides a valuable information for assisting patients with an abnormal walk, as the CCI are not appropriate for the latter case. Moreover, it should be mentioned that the $\Theta(t)$ angular value of the knee, which is to be predicted, is used only during training for NMI (calibration phase on 1 cycle) and is not provided to the algorithm during the rest of the test (evaluation phase on 11 cycles). Fig.3 shows the relation between the joint angles and the CCI during the gait cycle. It shows that the CCI do not provide a reliable or proper function to predict the joint angle for the control of an actuator. However, the determination of these angles is possible using NMI, which is a surjective function that is valid for control. It could be seen that each input corresponds to a single output, that is crucial requirement for designing a control function. As it could be seen on Fig.4, the NMI consistently achieved the best prediction for the knee angle, based on the ground truth and only after a short calibration. The CCI performed well for subjects with a normal walk but the main source of error was with patients suffering from cerebral palsy or stroke.

Fig.3 Knee joint angle variation $\Theta(t)$ computed for CCI1, CCI2, CCI3 and NMI as a function of walk cycle.

Fig.4 Comparison of error estimation for CCI1, CCI2, CCI3 and NMI of stroke patient. Walking phases are DS1 (first double support), SS (single support), DS2 (second double support) and SWP (swing phase).
The estimation error, in Table 1, shows a big variability for CCIs in function of Knee angles for 11 gait cycle of a stroke subject, where the estimation error of CCI1 reaches (48.1 ± 2.2 ) in DS1, for each CCI between (26.8±1.6 to 57±2.9) in SS, (30.9 ± 1.7 to 66.9 ± 3.1) in DS2, and (13.2±3 to 40.9 ±2) in SWP, however, this estimation reaches a maximum value (2.9±1.1 ) for NMI in all the phases of the gait cycle.

IV. CONCLUSION

The NMI is a new bio-kinematic index, focused on producing a reliable and generic estimation of the articulation flexion/extension using EMG to determine muscle co-contraction. The main contribution of this new index is to provide a model-free estimation of the walk cycle, without imposing a pre-defined trajectory on the patient. The patient is thus considered as an expert, providing a personal optimal walk cycle for calibration, and the NMI yields a robust index for designing an exoskeleton controller. This index evaluated on a dataset of 20 subjects, some with normal walk cycle and others with abnormal ones, using kinematic data as ground truth. The NMI largely outperforms existing CCIs. Following this work, the NMI will be integrated in the control scheme for a lower limb exoskeleton used as walking assistance for people suffering of spasticity.

REFERENCES