

Application of Kohonen maps to kinetic analysis of human gait

Silvia Elizabeth Rodrigo*, Claudia Noemí Lescano, Rodolfo Horacio Rodrigo

Abstract In recent years the use of artificial neural networks for classification and analysis of kinematic and kinetic characteristics of human locomotion has greatly increased. This happens in an attempt to overcome the limitations of traditional dynamic analysis and to find new clinical indicators for interpreting quick and objectively the large amount of information obtained in a gait lab. One of the most widely used neural networks for human gait analysis is the self-organizing or Kohonen map, based on unsupervised learning without prior definition of the formed natural groups. Among the advantages of using this type of neural network is the data dimensionality reduction, with minimal loss of information content, and the grouping of them in function of their similarities. Taking into account this, in this work an application case of a Kohonen map for clustering of locomotion kinetic characteristics in normal and Parkinson's disease individuals is presented. The results indicate that the groups identified by the map are consistent with the classification carried out by experts in function of traditional gait dynamic analysis, showing the potential of this technique for distinguishing between a population of individuals with normal gait and with gait disorders of different etiology.

Keywords Human gait, Parkinson's disease, Artificial neural network, Clustering.

Aplicação de mapas de Kohonen à análise cinética da marcha humana

Resumo Nos últimos anos, tem aumentado significativamente o uso de redes neurais artificiais para a classificação e análise cinemática e cinética da marcha humana, em uma tentativa de superar as limitações da análise dinâmica tradicional e de encontrar novos indicadores clínicos para interpretar, de forma rápida e objetiva, a grande quantidade de informação obtida em laboratórios de marcha humana. Uma das redes neurais mais utilizadas para análise de marcha é o mapa de Kohonen ou mapa auto-organizado, baseado em aprendizado não supervisionado, sem uma definição prévia de grupos naturais que se formam. O uso deste tipo de rede neural tem mostrado benefícios significativos associados com a redução da dimensionalidade dos dados com mínima perda de conteúdo de informação e com o agrupamento de dados de acordo com suas semelhanças. Neste contexto, este trabalho apresenta um caso de aplicação de um mapa de Kohonen como classificador das características cinéticas da locomoção em indivíduos normais e com doença de Parkinson. Os resultados indicam que os grupos identificados no mapa são consistentes com a classificação feita por especialistas com base em análise dinâmica tradicional, que mostra o potencial desta técnica para diferenciar populações de indivíduos com marcha normal e de indivíduos com distúrbios da marcha de etiologia diferente.

Palavras-chave Marcha humana, Doença de Parkinson, Rede neural artificial, Classificação.

Introduction

Human gait is a complex spatiotemporal process involving structures and functions of the neuromusculoskeletal system of human body (Sánchez Lacuesta *et al.*, 1999; Winter, 2005). Although this process is of individual character, the similarities between individuals are such that a typical pattern of normal human gait can be found. In addition, this pattern experiences modifications due to the influence of diverse factors and particularly under certain pathological conditions, where the gait efficiency diminishes with the consequent increase of energy expenditure that this implies (Lehmann and De Lateur, 1999).

Among these pathological conditions, those from neuromuscular etiology such as Parkinson disease (PD), a neurodegenerative disorder of the basal ganglia, are included. Parkinsonian gait is characterized by small shuffling steps, muscle rigidity at limbs and neck, variability in stride length and walking speed during free ambulation and postural instability, with the risk of falls and loss of functional independence in the most severe cases (Davie, 2008; Lehmann and De Lateur, 1999).

Through computer analysis realized in gait laboratories it is possible to obtain kinematic, kinetic and electromyographic variables that characterize normal and pathological gait patterns (Medved, 2001). Proper processing and analysis of such locomotor information is a key factor to obtain an accurate diagnosis of the considered gait pathology and to plan its treatment (Battistella *et al.*, 2009; Rodrigo, 2008; Rodrigo *et al.*, 2008). However, these requirements imply spending many hours analyzing the collected raw data by trained personnel. Moreover, the difficulty to find new ways of linking information in complex locomotion disorders such as PD through traditionally used algorithms hinders the task of defining more precisely the underlying cause of this disorder and its effect on locomotor activity (Simon, 2004).

A possible solution to this problematic is to apply clustering techniques based on artificial neural network (ANN), which helps to quickly discriminate between normal and pathological gaits. In this regard, numerous studies show the high probability of success in achieving this classification from different types of supervised and unsupervised ANN (Barton and Lees, 1997; Barton *et al.*, 2006; Chau, 2001; Köhle and Merkl, 1996). Specifically, the aim of this paper is to show the feasibility of using an unsupervised ANN, as is the case of self-organizing maps or Kohonen maps, SOM (Kohonen, 2001), for distinction between normal and parkinsonian gait patterns.

For this purpose in the next section first, the ANN input variables or descriptors used to differentiate normal and Parkinson's disease inter-subject gait patterns are defined. Second, the structure of SOM map and the procedure used to achieve the classification is described. Then, the results obtained are shown and finally, a brief discussion precedes the conclusions of the work.

Materials and Methods

Human gait database

To demonstrate the suitability of the unsupervised ANN, a database corresponding to 60 normal individuals (mean 63.3 years, range 37 to 86 years, 47% men) and 60 patients with idiopathic Parkinson's disease (mean 68.8, range 45 to 84 years, 62% of men) was used as classifier of inter-subject gait variability. Among other variables, this database contains demographic information such as the weight of the analyzed subject and a commonly used system, such as the Hoehn & Yahr scale, for describing how the symptoms of Parkinson's disease progress (Hoehn and Yahr, 1967). According to the original scale, the natural progression of PD is developed through 5 stages of ever increasing severity. Since then, stage 0 has been added (corresponding to individuals with normal gait), and stages 1.5 and 2.5 have been proposed, resulting in a modified scale as is indicated in Table 1. In contrast to normal cases, pathological cases here considered correspond respectively to 40 and 20 cases of stages 2 and 2.5 based on the modified Hoehn and Yahr scale. These stages are related to patients that show an altered gait with bilateral symptoms but without loss of body balance (Hoehn and Yahr, 1967; PhysioBank, 2009).

As well, the database includes kinetic data for both feet that were acquired during a gait test of about

Table 1. Classification of severity level of PD according to modified Hoehn and Yahr scale (Hoehn and Yahr, 1967).

Stage	Signs y symptoms
0	No signs of disease.
1	Unilateral symptoms only.
1.5	Unilateral and axial involvement.
2	Bilateral symptoms. No impairment of balance.
2.5	Mild bilateral disease with recovery on pull test.
3	Balance impairment. Mild to moderate disease. Physically independent.
4	Severe disability, but still able to walk or stand unassisted.
5	Needing a wheelchair or bedridden unless assisted.

2 minutes on level ground at normal self-selected cadence for each test participant, which has been digitalized and recorded at 100 samples per second (PhysioBank, 2009). Underneath each foot there are 8 sensors that measure force (in Newtons) as a function of time and together represent the plantar pressures distribution for every foot. The database also contains two signals that reflect the sum of the 8 sensor outputs for each foot, corresponding to the vertical component of the ground reaction force (GRF) for both feet (Infotronic, 2005). Besides, to minimize the effects of start-up and to match the length of the vector data for all samples the first 5 seconds of data were discarded, opting finally for a vector length of 5600. Figure 1 displays the vertical component of GRF during a gait cycle (GC) for both feet, obtained for a subject with normal gait and a Parkinson's patient from the mentioned database.

Definition of variables

The choice of ANN input variables is based on typical inter-subject differences observed in the size and shape of the curve representing the temporal variation of the vertical component of GRF, depending on the condition of analyzed subject. Indeed, as displayed on Figure 1a for a case of normal gait, such curve has two peaks. The first one occurs during heel strike with the ground at the beginning of stance phase, while the second peak is caused by the upward force exerted by ground during toe off at the end of the same phase (Sánchez Lacuesta *et al.*, 1999). Besides, the stance and swing phases pass between 0 - 60 and 60 - 100% GC, respectively.

In contrast, the curve shown in Figure 1b for the case of Parkinson's disease patient exhibits some times, a reduction in the peaks height of GRF, in similar form to what occur in elderly individuals. Also of note is the variation of taken time for stance

and swing phases for PD patients with respect to normal individuals (in this case 70 and 30% of GC, respectively), so as to ensure as far as possible, the body stability during locomotion. Later stages of this disease are described by a gait with small shuffling steps and a single narrow peak of vertical component of GRF, thus reducing significantly the time required for swing phase (Koozekanani, 1987).

The observed differences in Figure 1 between kinetic patterns for normal and parkinsonian gaits are consistent with the general conclusion stated by Crowther (2008, p. 357)

[...] a stable movement pattern is a behavioral state that is reproducible and independent of others and equates to low variability.

Higher gait variability can be caused by injury, disease, or aging, indicating that the related neuromuscular system is less stable. Particularly in PD patients, different researches have shown increased gait variability, both intra- or within-subject and inter- or between-subject, which have been linked to an increased risk of falls in such patients (Hausdorff *et al.*, 1998).

According to these observations and in order to select the most appropriate input variables for fast differentiation of normal and parkinsonian gait patterns by ANN, distinct parameters were employed. Such parameters characterize the inter-subject gait variability throughout the considered temporal signal. Among them, were utilized: the mean standard deviation of GRF signal during consecutive stance phases, its maximum value (which defines the mean peak GRF intensity for successive stance periods), the mean coefficient of variation (CV) of GRF, and the mean sum of GRF over successive stance phases (as an approximation of the mean value of the area under the curve through these consecutive phases).

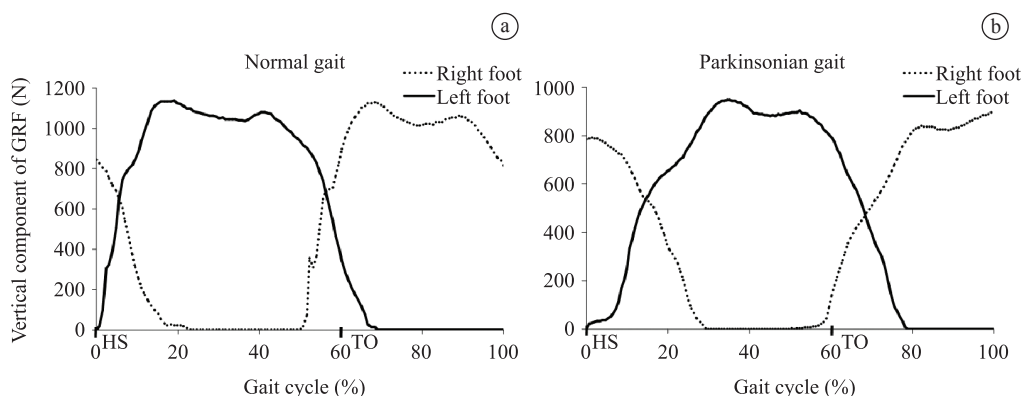


Figure 1. Vertical component of ground reaction force (GRF) for both feet during a gait cycle (GC). a) normal gait; b) parkinsonian gait. For normal gait stance phase (0-60% GC) begins at left heel strike (HS) and ends with toe off (TO). Swing phase takes between 60 and 100% GC. Notice how these percentages vary for parkinsonian gait.

As an example of inter-subject variability for GRF signal during successive stance phases, the variability exhibited by the mean coefficient of variation of GRF for the right and left legs is depicted in Figure 2. These curves correspond respectively to the 60 normal (on the left) and the 60 pathological (on the right) considered cases.

As well, in Figure 3 the inter-subject variability of mean sum of GRF through consecutive stance phases for the right and left legs is represented for the same normal (on the left) and pathological (on the right) cases.

As can be visualized both in Figures 2 and 3 on the left, the inter-subject variability values for normal gait exhibit not only a lower magnitude but also a minor temporal variability with respect to parkinsonian gait (Figures 2 and 3 on the right). This shows the feasibility of utilizing these indexes as descriptors of inter-subject gait patterns through ANN.

Taking into account these antecedents and after several trials, the following descriptors of the vertical component of GRF characteristics for both feet were finally selected:

- Mean coefficient of variation (CV) of GRF during consecutive stance phases to describe the inter-subject variability in GRF magnitude and waveform throughout analyzed gait cycles. It is determined in each sample and for left and right feet, as the ratio between the mean standard deviation and the mean value of GRF for each trial. After that, the average value of CV of GRF is determined for all successive stance phases of the analyzed gait cycles.
- Mean sum of GRF during successive stance phases, as an approximation of area under the GRF curve over these consecutive phases of analyzed gait cycles. It is calculated in each sample and for left and right feet, as the mean value of the sum of GRF for each trial, then averaging the achieved values for all consecutive stance phases of the temporal signal considered.

As well, such variables were normalized by their respective maximum values, so that they have the same rank, in this case between 0 and 1, and the same variance, equal to 1. Finally, the ANN input

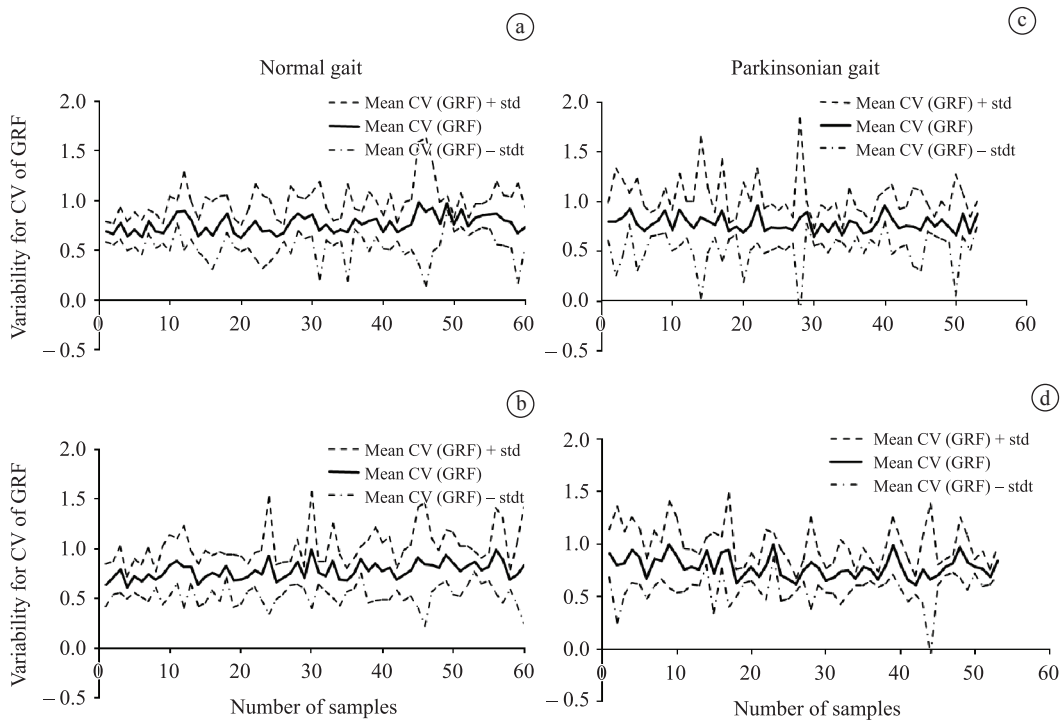


Figure 2. Inter-subject variability evaluated by the mean coefficient of variation (CV) of GRF during consecutive stance phases of gait cycle, corresponding to 60 normal subject and 60 parkinsonian patients analyzed. a) left foot for normal individuals; b) right foot for normal individuals; c) left foot for parkinsonian patients; d) right foot for parkinsonian. In each figure, the visualized values represent the mean coefficient of variation of ground reaction force during successive stance phases (mean CV(GRF)) and its corresponding standard deviation (std).

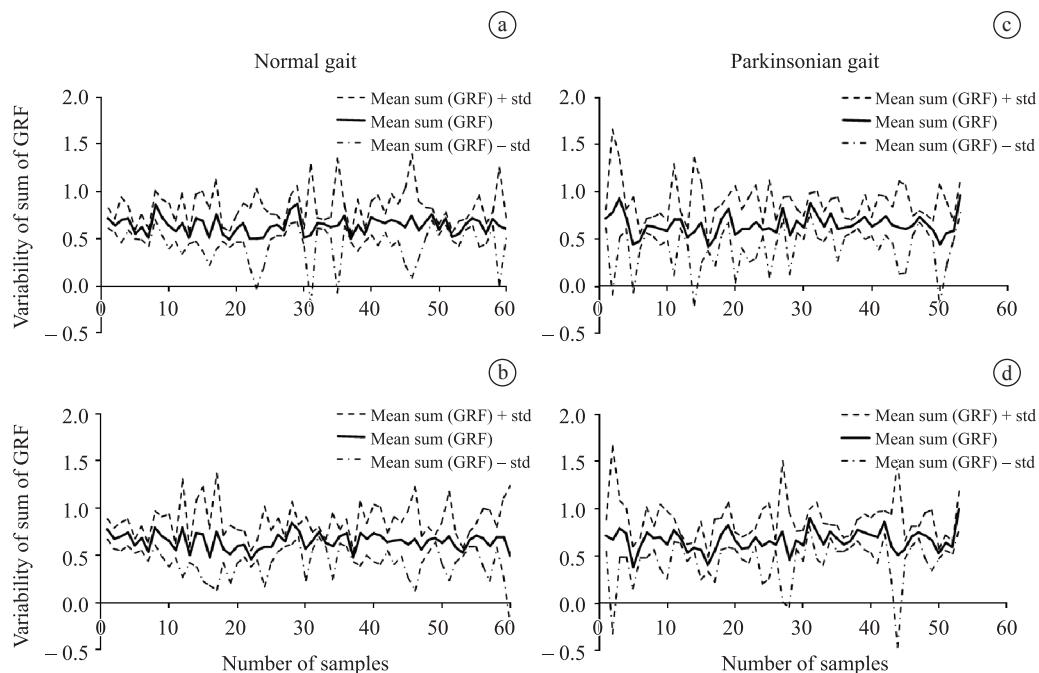


Figure 3. Inter-subject variability evaluated by the mean sum of GRF during consecutive stance phases of gait cycle, corresponding to 60 normal subject and 60 parkinsonian patients analyzed. a) left foot for normal individuals; b) right foot for normal individuals; c) left foot for parkinsonian patients; d) right foot for parkinsonian. In each figure, the visualized values represent the mean sum of ground reaction force during successive stance phases (mean sum (GRF)) and its corresponding standard deviation (std).

data are composed by normal and pathological cases randomly chosen.

Type and structure of the artificial neural network

The ANN chosen to classify the gait data between normal and pathological is the type called self-organizing map (SOM) or Kohonen map, a type of ANN algorithm whose aim is to discover and display the underlying structure of the data entered in it, usually of high dimensionality. A SOM map is defined by an array of neurons (connected by a neighborhood relationship that defines the structure of the network) of a single layer, which represents filters or processors selectively tuned by input data simultaneously presented to all network nodes. Consequently, the neurons transform the input signals in a probability distribution encoded by positions on a map structured according to its topology (Haykin, 2005; Kohonen, 2001; Merelo, 2004).

From a training process based on unsupervised competitive learning, the SOM group the input data (of n dimension) in network nodes (each of which also has dimension n) in terms of their similarity, projecting them on a smaller map output in where the input data characteristics are distributed in a gradual and order form (Kohonen, 2001). Thus, SOMs allow for two

tasks on the input data: to reduce their dimensionality and to visualize similarities. In this case and bearing in mind that the objective of this work only aims to assess the usefulness of ANN as differentiating normal and parkinsonian gaits, a 2×2 structure was chosen.

Generation, training and visualization of the artificial neural network

For SOM implementation a Matlab® Toolbox was utilized through *newsom* function, which generates an ANN defined in the following form (Vesanto *et al.*, 1999):

$$net = newsom(P, [d1, d2], tfcn, dfcn, steps, in) \quad (1)$$

being: P : matrix $R \times Q$, with R equal to input variable number and Q , sample number; $[d1, d2]$: neural network size (2×2); $tfcn$: neural network topology function, arranged in hexagonal, grid or randomly form (*hextop*, *gridtop*, or *randtop* respectively; '*hextop*' was chosen); $dfcn$: distance function from a particular neuron to its neighbors, Euclidian, by steps, Manhattan or box types (*dist*, *linkdist*, *mandist* y *boxdist*, respectively, '*linkdist*' was chosen); *steps*: steps from the neighborhood to reduce its size to 1 (default = 100); *in*: neighborhood initial size (=3).

After defining the ANN (with the values specified for the topology and distance function), the arrangements for the input data (an array of 120 samples of 4 elements each) were established. The number of samples for training, validation and testing was distributed in 90, 15 and 15, equivalent to 75, 12.5 and 12.5% of samples, respectively.

For training the Matlab® *train* function based on the function *NET.trainFcn* and training parameters *NET.trainParam* was used, whose syntax is as follows:

$$[net, tr] = train(NET, X, T) \quad (2)$$

where: *NET*: generated SOM map; *X*: input vector (in this case, 90 samples of 4 elements each), *T*: target vectors (in this case, do not apply), *net*: new ANN generated from training, *tr*: training register (number of epochs or iterations (=500) and ANN performance (quantization error of 0.01).

According to this function, the training algorithm move the SOM weight vectors (initially selected through Matlab® *rand*s function, which generates random values between -1 and 1) to extend then through the cloud of input data, so the map is organized in such a way that neighboring neurons in the ANN gain similar weight vectors. For this the Matlab® iterative training algorithm (by default *batch*) was used in two stages. In this process the total data set is presented to the network and the new weight vector is a weighted average of the input data vectors. In the first stage, which aims to organize the map, a high learning rate equal to 0.09 and a radius 3 neighborhood were employed, while in the second one, of fine tuning, these values were set respectively at 0.02 and 1, finishing the training (after 800 iterations in 12 seconds) with a stable arrangement of the input patterns. For a more detailed description about this issue the references cited can be consulted (Haykin, 2005; Kohonen, 2001; Merelo, 2004; Vesanto *et al.*, 1999).

Finally, after the training phase, the network produces an output suitable to all input samples through *sim* Matlab® function, whose syntax is:

$$Y = sim(NET, P) \quad (3)$$

where: *NET*: represents the trained ANN, *P*: set of input samples; *Y*: ANN output.

Results

Figure 4 displays the clusters obtained for both types of gait data the SOM with the set of selected input variables, that is, the mean coefficient of variation and mean sum of GRF during consecutive stance phases of the analyzed gait. In this figure, the small

hexagons represent the neurons, while the lines within the large hexagons connect neighboring neurons. In addition, the colors of the regions containing these lines indicate the distances between neurons, representing the dark and light colors, large and small distances respectively. As well, the dark segments dividing the two-dimensional map show that the Kohonen map grouped the input data into two distinct regions, corresponding to individuals with normal and parkinsonian gaits.

Furthermore, in Figure 5 the distribution of samples between normal and pathological cases is shown. From the analysis of these results it can be inferred that the SOM map operates as an appropriate classificatory taking into account that was able to recognize respectively, 72.09% (corresponding to 31 of 43 normal cases) and 70.21% (in this case, 33 of 47 parkinsonian cases) of normal and pathological gait patterns. Other evaluators of SOM performance

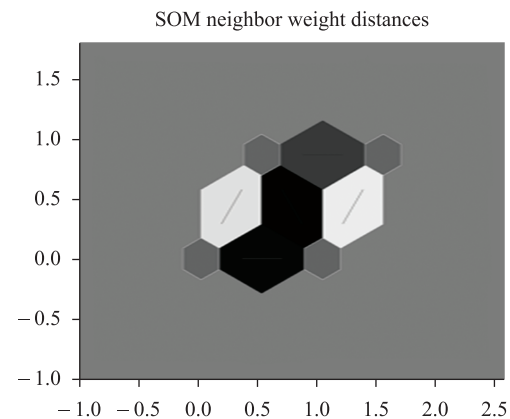


Figure 4. Results of SOM neural network corresponding to neighborhood defined by input vector samples.

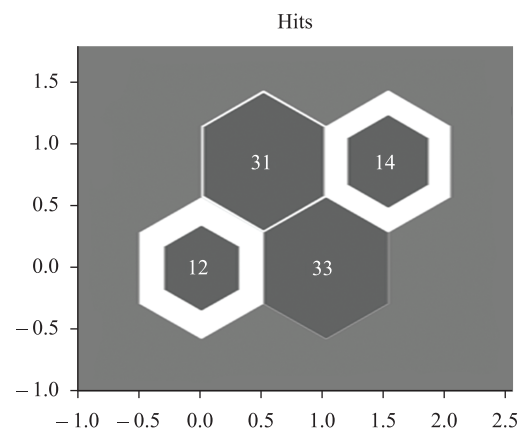


Figure 5. Classification of normal and parkinsonian gait pattern samples for the mean coefficient of variation (CV) of GRF and the mean sum of GRF as input variables to SOM.

are the sensibility and specificity indexes, being their values equal to 0.721 and 0.702, respectively. These indexes were calculated from a number of 31 true negatives and 12 false positive from a total of 43 normal samples, and 33 true positives and 14 false negative from 47 pathological samples.

In contrast with the selected variables for this SOM application, using as input variables the mean of maximum value and the mean standard deviation of GRF during consecutive stance phases, the results shown in Figure 6 are generated. In this case, the sensibility and specificity indexes were equal to 0.6744 and 0.6809, respectively, obtained from a number of 29 true negatives and 9 false positive from a total number of 43 normal samples, and 32 true positives and 20 false negative from 47 parkinsonian samples.

On the other hand, Figure 7 indicates the weight adjustment of input variables made by the SOM map in function of the spatial distribution of the samples presented during training (lighter small dots). In addition, the dark spots represent the neurons in the SOM map. Also, there exists a correlation between dark and light areas shown in Figure 4 and the distances between neurons in 7.

Finally, Figure 8 shows the weight map for each element of the input vector (in this case, two elements for each leg) that characterizes the influence degree of each input on each network neuron. Dark colors are indicative of greater weight and thus, greater influence in defining the clusters. Furthermore, if the connection patterns of two inputs are very similar, it can be assumed that the inputs are highly correlated. Such is the case of weights for inputs 1 and 2 and for the inputs 3 and 4, representing respectively, the mean CV of GRF and the mean sum of GRF for left and right feet.

Discussion

In the literature there are previous works oriented to classification of kinematic and kinetic gait patterns through ANN. About this, the work by Chau (2001) provides an excellent review on the subject. As application examples, an automatic classification of patients into groups with a SOM map was obtained using the vertical component of GRF under both feet, with results agreeing with the clinical classification of the patients (Köhle and Merkl, 1996). Other research group utilized a supervised ANN to differentiate gait patterns from hip and knee joint angles in the sagittal plane (Barton and Lee, 1997). The same group used then a SOM map to reduce the complexity of three-dimensional joint kinematic and kinetic

data, projecting them in a two-dimensional map (Barton *et al.*, 2006). Also, in a previous work a classification of normal and parkinsonian gait based on a multilayer perceptron ANN was achieved from the power spectral density of GRF for the left leg as input variables (Battistella *et al.*, 2009). These examples

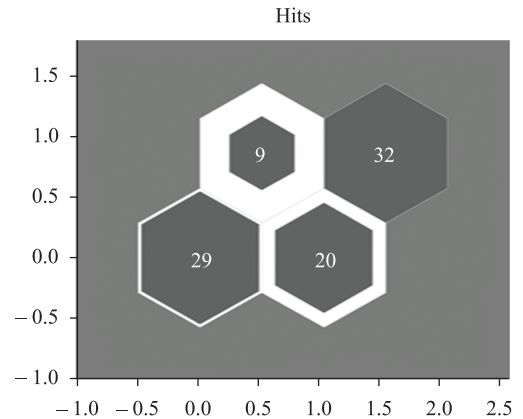


Figure 6. Classification of normal and parkinsonian gait pattern samples for the mean maximum value of GRF and the mean standard deviation of GRF as input variables to SOM.

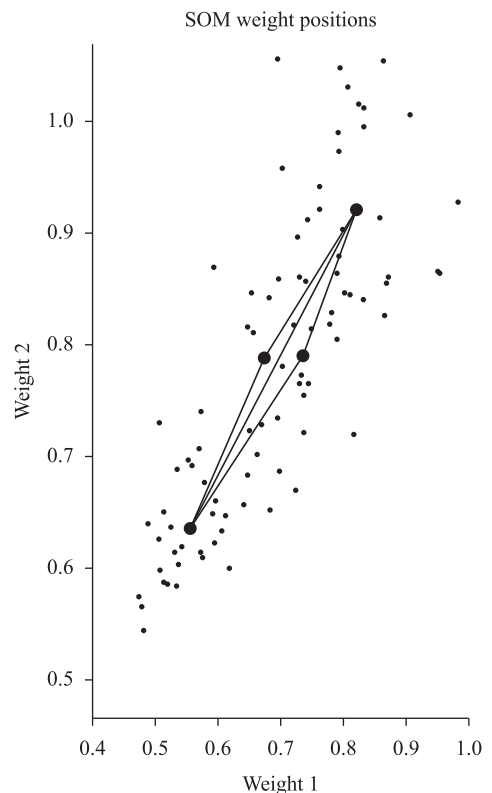


Figure 7. Results of SOM neural network corresponding to weight adjustment using the mean coefficient of variation (CV) of GRF and the mean sum of GRF as input variable.

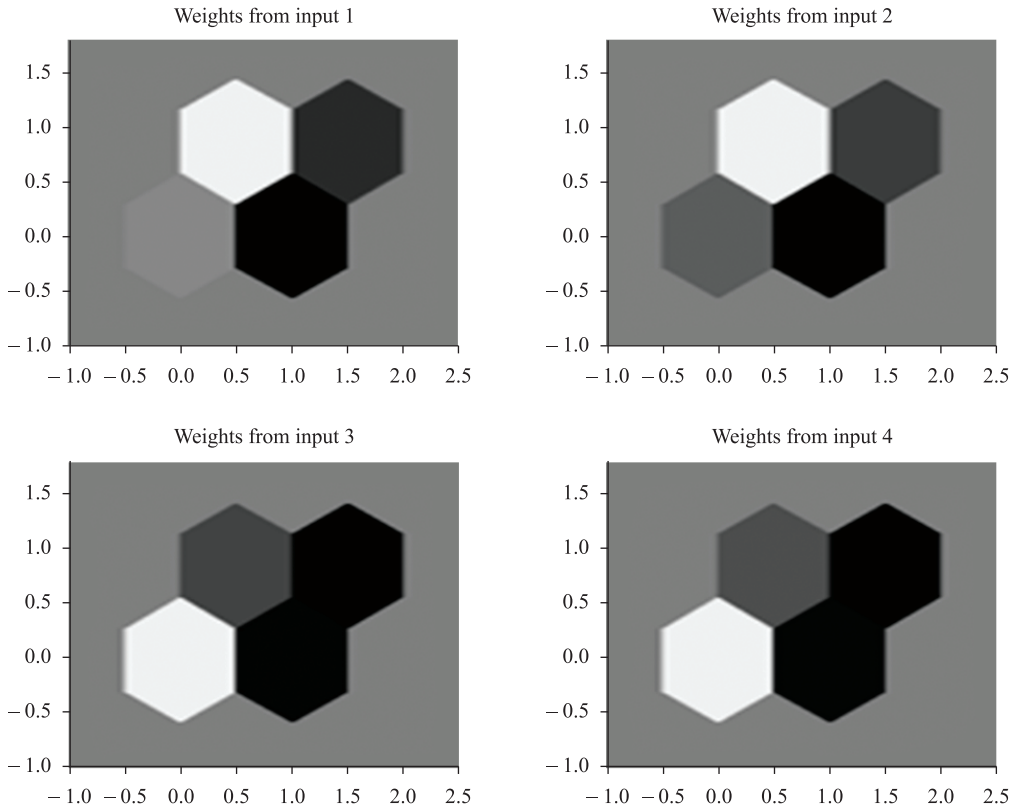


Figure 8. Influence of the input variables weights (mean coefficient of variation (CV) of GRF and the mean sum of GRF) on neural network neurons for clustering definition.

show the applicability of ANN as an objective tool that may enhance decision making.

In the present work, a series of test were performed with different input variables for the SOM. In contrast with the results obtained in Figure 6 for the mean maximum value of GRF and the mean standard deviation of GRF, the use of the mean CV of GRF and the mean sum of GRF over successive stance phases were the best option to obtain a relative good classification between normal and parkinsonian gaits (Figure 5). In the first case, the sensibility and specificity indexes were respectively equal to 0.6744 and 0.6809, meanwhile in the second one (that is, for the combination of SOM input variables used in this work), the values of sensibility and specificity indexes were 0.721 and 0.702, respectively. Such results demonstrate for this application, the generality of the obtained solution.

If the classification of gait patterns using a multilayer perceptron and a SOM ANNs are compared, better results are achieved in the first case. However, it is noteworthy that the use of a multilayer perceptron is based on supervised learning, unlike SOM, where

no target is included to contribute to the differentiation of input patterns.

From the point of view of the causative factors of gait variability, different hypothesis have been postulated. One of them proposes that it is due to structural complexity of the human neuromusculoskeletal system, what determines an even more complex functional system that is inherently variable, both within and among individuals (Bates *et al.*, 2004).

In addition, according to Bernstein's theory, effective organization of the multiple degrees of freedom of the neuromusculoskeletal system is a requisite for healthy functional and steady locomotion patterns (Bernstein, 1967), manifested by small fluctuations of gait parameters during free ambulation.

Nevertheless, in certain neuromuscular pathologies such as PD, such fluctuations are increased due to variability in stride length and walking speed during ambulation at self-selected cadence, determining an inability to maintain a steady walk with small stride-to-stride fluctuations (Hausdorff *et al.*, 2000, 2003; Hausdorff, 2005). Other research works also show the

importance of the integrity of cerebellum and basal ganglia structures, typically affected in PD patients, in adequately regulating force and timing control in motor activities such as locomotion (Davie, 2008; James, 2004; Lehmann and De Lateur, 1999).

Such considerations could explain the greater inter-subject variability observed in Figures 2 and 3 on the right for PD patients with respect to normal gait, on the left. Besides, consistent with the results of Figure 5, this differential gait attribute is reflected in the specificity index of 0.702, equivalent to a 70.21% of well classified pathological samples.

With respect to the clinical relevance of these issues, such changes in variability of gait parameters for PD patients could be a predictor of the level of deterioration of brain structures that regulates force and timing control for an efficient gait. In this sense, it is proposed here to estimate a reference range of normal gait that can be used then to compare the degree of gait impairment in the considered pathological condition. In this way, it can be quickly viewed in a first approximation, how the pathological condition deviates from normality.

Furthermore, greater inter-subject variability in PD patients would indicate an increased level of postural instability. As it is visualized in Table 1, the modified Hoehn and Yahr scale is based on the two-fold concept that the severity of overall parkinsonian dysfunction relates to bilateral motor involvement and the degree of gait balance.

In relation to the differential motor characteristics of these two categories of PD patients here considered (40 and 20 cases with Hoehn and Yahr index equal to 2 and 2.5, respectively) and according to Table 1, while the first has bilateral symptoms without impairment of balance, the second one exhibits mild bilateral disease with recovery on pull test. This last test evaluates postural instability, which is the relative ability or inability to recover from falling. In the pull test, the individual being tested is asked to maintain his balance while an examiner pulls him backward. As it is indicated in Table 1, PD patients with a Hoehn and Yahr index value of 2.5 are not affected by postural instability and for this reason, the observed differences can not be attributed to an increased risk of falls in such patients.

Summing up and according to these examples and observations, the results here obtained reveal the high probability of success in achieving differentiation between normal and parkinsonian gait patterns. In this sense, the specificity of the utilized descriptors played a key role not only in reducing the dimensionality of the input data, but also in finding similarities between both types of analyzed populations.

Conclusion

In this work a SOM map was used for clustering of normal and parkinsonian gait patterns. In the context of kinetic data analysis, the SOM can be regarded as a software tool that reduces the amount of data with minimal loss of information content. After the learning phase, input values can be classified as belonging to specific clusters, and in general hidden structures of the input data set can be detected.

From the biomedical point of view, the current application of SOM map indicates that the vertical component of GRF contains relevant information for differentiating quickly and objectively normal and pathological gait patterns. This shows that the utilized combination of input variables and the type of ANN is feasible for clustering of gait patterns. An additional advantage of using ANN is to obtain the classification automatically, without defining a biomechanical model of the human body to process the data. As well, the clinical relevance of this application of ANN is related with the fact that changes in gait variability with respect to normal gait, could be a predictor of the degree of ambulation impairment in the considered pathological condition.

Based on the obtained results, in future work it is expected to generate new descriptors to identify more accurately the mechanical and neurological behaviors of human body during parkinsonian gait, both intra- and inter-subject. Such descriptors could be parameters to identify the marked body imbalance or the increased variability between steps, typically seen in patients with Parkinson's disease.

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Authors

Silvia Elizabeth Rodrigo, Claudia Noemí Lescano, Rodolfo Horacio Rodrigo
Facultad de Ingeniería, Universidad Nacional de San Juan – UNSJ,
Av. Libertador General San Martín, 1109 (o), 5400, San Juan, Argentina