NOVEL SEMG PARAMETERS FOR EARLY DIAGNOSTICS OF NEUROLOGICAL DISEASES AND AGING

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Abstract. The early-stage diagnostics is critical for early treatment of neurodegenerative diseases. Although Parkinson’s disease (PD) as one of the most studied neurological diseases has been intensively studied, the pre-clinical markers of PD is yet to be established. Here we report on the non-linear characteristics of surface electromyogram (sEMG) and tremor acceleration as a possible diagnostic tool, and, in prospective, as a predictor for PD. We found that such nonlinear parameters as percent of determinism, sample entropy and correlation dimension allowed discriminating some 85% of healthy controls from PD patients. We intend to expand this approach to multiply sclerosis, schizophrenia and essential tremor.

Key words: Parkinson’s disease, electromyography, nonlinear parameters, early diagnostics, ageing.
1. Why to search for novel biosignals and early diagnostics in neurology?

1.1. Economical and personal burden.

Neurodegenerative diseases, such as Alzheimer’s disease (AD), Parkinson’s disease (PD), multiple sclerosis (MS), along with schizophrenia (SZ), essential tremor (ET) and Huntington’s disease (HD) imply strong economical burden of several kinds on the patients, their relatives and society. In the USA at present, AD afflicts 13% of people aged over 65 and 43% of those 85 and older (in total, an estimated 5.4 million) (Ambrose, 2013). PD is the second most common, after AD, neurodegenerative disease. The incidence of PD in developed countries is estimated at about 1% in people older than 60 years and 4% in people over 80 years (de Lau, 2006). SZ affects approximately 1% of the population but SZ patients occupy 25% of all hospital beds in the USA (Davies, 1994). The number of people with MS is 2–2.5 million globally, with rates varying from 0.5 per 100,000 in Africa to 200 per 100,000 in certain populations of Northern Europe (Adelman, 2013).

The personal burden means dramatic decrease of the quality of the patient’s life due to motor (tremor, rigidity, akynesia, atonia, postural instability, fatigue, and muscle weakness) and non-motor symptoms (impaired autonomic functions, depression, sleep, emotional and cognitive disorders). These motor and non-motor symptoms may decrease person’s ability to work and thus may imply early retirement, restrictions on profession choice and, hence, the salary (Keränen, 2003). Cost burden of neurological patients also includes medication, care costs, insurance etc. For example, annual economical burden on one PD patient in the developed countries exceeds 6000-25000 USD (Noyes, 2006; Tolosa, 2006), and on one MS patient 8000-50000 USD (Adelman, 2013). The overall U.S. 2002 cost of schizophrenia was estimated to be 62.7 billion USD (van den Heuvel, 2010).

These diseases, in a sense, are “non-lethal” pathologies because they do not cause immediate fatal outcome. They can last, like PD under medication, for decades. The average life expectancy of men with SZ is only slightly lower than would be expected for the population as a whole (Davies, 1994).

1.2. Early diagnosis.

In some neurological patients, diagnosis is highly uncertain, especially in the early stages. For example, only 70% of patients are correctly diagnosed with PD (Tolosa, 2006). PD, ET, MS, AD and HD share many motor symptoms among each other. There is a large body of papers on comparative studies of PD and ET (Becker, 2002). However, a diagnostic tool to differentiate these two pathologies is yet to be elaborated.

Neurodegenerative diseases are characterized by a lengthy prodromal due to their chronic course, which is known as “preclinical” phase (Berendse, 2006). For example, PD becomes clinically recognizable only when 60% of dopaminergic neurons in the compact part of substantia nigra are lost. This phase is believed to start long before emergence of clinical PD and then may slowly progress over 4.5 years (Postuma, 2013). The prodromal period presents unique opportunity in disease prevention or delay in PD onset. Some 10% of subjects over 60 years are in the “prediagnostic” phase of PD according to neuropathological reports (Stern, 2004). Therefore, predating the diagnosis of PD and identifying subjects at risk is an important goal for research aimed to postpone the onset of PD by neuroprotective therapy (Siderowf, 2006). Thus, the ultimate goal of the early diagnosis of PD would be switch from medical treatment to the disease management.
A predictive study would help to identify the subjects at-risk of future disease, to start earlier treatment, and to develop effective neuroprotective treatment strategies (Siderowf, 2006). Here, we report on current approaches in early diagnostics and differential diagnosis of PD with special stress on the nonlinear parameters of interference surface electromyography (sEMG) signal.

2. Current approaches to early diagnosis of neurological disease.

The preclinical (early clinical) predictors become a hallmark of the current neurology, which would present a window of therapeutic opportunity for many neurological diseases. PD is one of the best studied diseases in respect with early diagnosis. Visual, smell, sleep, vocal, handwriting, autonomic, motor and mental dysfunction due to dopaminergic deficit affect 60-80% of PD patients (Tolosa, 2006) and they are usually synchronized. These are used as prodromal markers for PD. However, definitive PD-sensitive diagnostic tool is lacking. So, the decisive diagnosis of PD is still post-mortem. All in all, current methods help to diagnose only 70-80% of the PD cases (Becker, 2002; Berendse, 2006; Müller, 2001; Tissingh, 2001), which is not satisfactory. It corresponds with misdiagnosis of PD estimated as 20-30% (Jankovic, 2008). Novel biomarkers for PD must be presented. Over the last years, single photon emission computed tomography (SPECT), positron emission tomography (PET), fMRI and transcranial sonography are widely used to assess dopaminergic function in PD patients, their relatives and healthy controls (Becker, 2002; Müller, 2001). However, their diagnostic yield is still far from obvious (Morrish, 1997). For example, up to 15% of subjects with normal imaging findings, have clinically evident PD, and vise versa, decreased dopamine content in basal ganglia seen on MRI is often associated with neurodegenerative diseases other than PD (Bajaj, 2013). Also, methods based on nuclear medicine and ultrasound are not appropriate for population-based studies due to their high costs, and insufficient availability (Becker, 2002; Müller, 2001). The concept of a preclinical phase of disease is not foreign also for AD (Nitrini, 2010; Sperling, 2011) and MS (Andersen, 2010).

3. EMG as a potential early marker for PD.

To our mind, the method for early diagnostics of neurological disease must be sensitive, cost-efficient, non-invasive, high throughput and non-hazardous. We believe that surface electromyography (sEMG) is such a method. EMG is a kind of “mirror” from skeletal muscles for central nervous system (CNS) because it reflects the activity of the spinal motoneurons due to motor units (MUs). No consensus exists about applicability of spectral-based analysis of sEMG to PD diagnosis because conventional linear parameters are still lacking to provide definitive difference between the PD and healthy controls.

The morphology-based analysis has shown promising results in discriminating PD and healthy controls. The method is based on the histogram and crossing rate analysis of sEMG signals (Rissinanen, 2007). sEMG kurtosis, a parameter based on higher order statistics, is reportedly increased in PD patients, that might reflect increased number of spikes due to increased synchronicity of MU firing (Meigal, 2009). Also, previous works have reported that sEMG waveform can better be modeled as an output of a nonlinear dynamic system, rather than a stochastic output of a white-noise driven linear system (Rissanen, 2007). Nonlinearity is a hallmark of complex dynamic systems (Elbert, 1994; Stam, 2007). As a nonlinear signal, sEMG displays chaotic behavior, i.e. its time series 1) evolves over the time, 2) depends on the initial state, and 3) is fractal in the terms of dimensionality (Nieminen, 1996). Thus, as a nonlinear signal, sEMG can be characterized by the state of deterministic chaos (Webber, 1995). Therefore, sEMG might give clues to describe dynamics of the neuronal circuits in the terms of regularity, predictability, and complexity (Goldberger, 2002). Indeed, it has been recently found that such nonlinear parameters as
approximate entropy, percent of determinism based on recurrence quantification analysis (RQA), and dimensionality based on fractal analysis are highly sensitive for hidden rhythms on sEMG in subjects under fatigue and condition of increased MU synchronization (Del Santo, 2007).

sEMG in the PD patients is known to be rich in regular clusters (grouping) at the characteristic tremor frequencies (4-6 Hz) due to increased synchronization of MU (Vaillancourt, 2000). Acceleration signal has also been studied in PD patients using both linear and nonlinear parameters (Sturman, 2005; Vaillancourt, 2000; Vaillancourt, 2003). These findings led us ask whether either readily visible or “hidden” rhythms in sEMG contribute to its nonlinear features and thus yield difference between the PD patients and healthy controls? If this hypothesis holds, the PD patients might present a more regular time-dependent structure of sEMG and acceleration time series, while healthy subjects – a less regular and more complex signal. Also, PD patients with less expressed sEMG clustering might present less regular signal. To test this hypothesis, we compared a variety of novel nonlinear parameters with the classic linear parameters between PD patients with various UPDRS scores, healthy young and older controls.

We found that nonlinear parameters of sEMG signal in the PD group significantly differed from the ones in the healthy control groups (Meigal, 2009). In particular, percent of determinism (%DET) values of sEMG were significantly higher in the PD group, while sample entropy (SampEn) and correlation dimension (CD) were lower in comparison to old and young controls. Instead, such traditional parameters as RMS and MDF did not differ between groups. Decreased CD of sEMG in the PD group may indicate increased self-similarity of the myoelectrical signal over time (Siderowf, 2006) and hence, lowered complexity of the underlying neural network. Decreased SampEn of sEMG signal in the PD group may reflect higher regularity of sEMG. High %DET reflects abundant wave features in sEMG, either readily visible by eye or quasi-waves during, e.g., increased motor unit synchronization (Rissanen, 2007; Rissanen, 2009). Clustering of MU action potentials was the characteristic of raw sEMG of many our PD patients (Meigal, 2009). Interestingly, %DET also was increased in some patients with visually stationary sEMG. We have also found that such sEMG parameters, as %REC, and %DET were correlated with Unified Parkinson’s Disease Rating Scale (UPDRS) score. Most of novel sEMG characteristics correlated also with finger tapping scores (Meigal, 2009). Correlation values were the most significant in the state without additional loading, and they decreased when loading increased. This probably indicates emergence of "regular normal" postural muscle tonus, which erased the difference between the groups. In a sense, under loading sEMG of PD patients became more “normal”. Thus, sEMG signal in PD is less complex, more predictable and regular. It means that rhythmic activity takes place in the spinal cord, resulting in more or less obvious clustering of sEMG. From the physiological point of view, these data reflect increased synchronization of MU activity or increased clustering coefficient of signal generator (the spinal neuron circuitry) (Stam, 2007). It could also well be that the spinal cord rather relays this periodicity from the upper levels of CNS, than generates it itself.

4. sEMG parameters in SZ patients.

Investigations of nonlinearity of biosignals in SZ are usually focused on brain and EEG. For example, application of graph theory revealed that brain network is substantially disintegrated in SZ (Wu, 2005). Muscles may reflect aberrant descending motor flow thus forming a hypothetical “muscle channel” for sEMG diagnostics of SZ. Dopamine hypothesis of SZ argues that its symptoms can be explained by elevated level of dopamine in the brain (Carlsson, 1963). As such, one can suggest that sEMG parameters must be opposite to those found in PD, at which dopamine production is decreased (Hornykiewicz, 1975). Namely, we expect that sEMG in SZ patients might have been characterized by high entropy and correlation dimension, and low percent of determinism. Further measurements would support or refute that assumption.
5. Non-linear parameters of sEMG under ageing.

Because of overwhelming prevalence of movement disorders among elderly individuals, it is important to distinguish healthy aging from neurological pathologies (Sturman, 2005). Studies, in which the linear parameters of sEMG were used, showed that EMG signal changes over lifetime (Gerasimova, 2004). There is also a strong conviction that biosignal in human organism become simpler over the lifetime (Goldberger, 2002) due to either loss of CNS oscillators over lifetime or losing capability to balance these oscillators (Vaillancourt, 2003). The outcome of such condition would be dominance of one (few) oscillators which are able to express their own rhythm on sEMG. However, we did not observe reduced complexity of sEMG signal in older controls (Meigal, 2009; Meigal, 2012). Moreover, SampEn and CD were even slightly elevated in aged group, though non-significantly (Figure). It means that in old controls sEMG signal was even more complex in comparison to PD patients and young controls. Thus, nonlinear parameters might help to distinguish between pathologically affected and “normal” sEMG samples. By now, our methods allow to discriminate almost 87% of PD patients from healthy controls (Rissanen, 2009). It could well be that trajectory of complexity over lifetime bifurcates at definite age forming two new trajectories - one for “pathology” and another – for “normal” ageing. Such trajectory may have diagnostic value to predict pathological changes of the motor system.

**Figure.** Trajectories of the nonlinear sEMG parameters over lifetime. Bifurcation of trajectories is denoted as a light grey circle. Difference is statistically significant between PD patients and young controls, but it was not significant between young and older controls.

CD – correlation dimension, DET% - percent of determinism, En – sample entropy.

6. Conclusion

The novel nonlinear sEMG parameters have potential in the preclinical diagnosis of PD due to their relatively high discrimination power, cost efficiency and high throughput. Additionally, nonlinear parameters of sEMG may help to distinguish between healthy ageing and pathology.
during elderly age. Finally, we believe that sEMG could be helpful in early detection of other neurodegenerative diseases, such as AD, MS, and also SZ.

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