


# FNIRS application in Parkinson's Disease

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## Definition

The management of people affected by neurological disorders, such as Parkinson's disease, requires the adoption of targeted and cost-effective interventions to cope with chronicity. Although therapy adaptation and rehabilitation represent major targets, affordable and reliable neurophysiological correlates of cerebral activity to be used throughout treatment stages are often lacking. The functional Near-Infrared Spectroscopy (fNIRS) represents a versatile optical neuroimaging technology for investigating cortical hemodynamic activity in the most common chronic neurological conditions, including Parkinson's disease, with the advantages of non-invasiveness and portability which make fNIRS suitable for carrying out multiple measurements in rehabilitation settings.

## 1. Introduction

The management of people affected by age-related neurological disorders requires the adoption of targeted and cost-effective interventions to cope with chronicity <sup>[1][2]</sup>. Therapy adaptation and rehabilitation represent major targets requiring long-term follow-up of neurodegeneration or, conversely, the promotion of neuroplasticity mechanisms <sup>[3]</sup>. However, affordable and reliable neurophysiological correlates of cerebral activity to be used throughout treatment stages are often lacking <sup>[4][5]</sup>. The application of functional neuroimaging methods, including functional Magnetic Resonance Imaging widely accepted as gold standard modality <sup>[6]</sup>, is often limited either by high costs, long scan times, poor temporal resolution and sensitivity to motion that reduce patient compliance and the possibility to assess responses evoked by complex stimuli. Functional Near-infrared Spectroscopy (fNIRS) <sup>[7]</sup> is gaining an increasing relevance as a more versatile solution to investigate cerebral activity based on the principle of neurovascular coupling. It is a non-invasive optical imaging technique that employs near-infrared (NIR) light to measure cortical oxygenation and consequently cortical activation. This technique employs pairs of NIR sources and detectors placed over predefined scalp locations to estimate light intensity attenuations, next converted to oxygenated (HbO<sub>2</sub>) and deoxygenated hemoglobin (HbR) concentration changes according to the modified Beer-Lambert law <sup>[8]</sup>. Other chromophores are essentially stationary and allow sufficient transmittance in the NIR region. fNIRS is better suited to perform low-cost and multiple acquisitions in the rehabilitation context during the execution of a wide variety of tasks: motor, cognitive and somatosensory <sup>[9]</sup>, as well as postural control and free-walking conditions <sup>[10][11]</sup>.

## 2. fNIRS in Parkinson's Disease

Parkinson's Disease (PD) is a slowly progressive neurodegenerative disorder and represents one of the most common pathologies related to ageing, involving about 6.1 million individuals worldwide and presenting considerable implications on national healthcare systems <sup>[12][13]</sup>. This pathology is characterized by a prominent impairment of motor functions (bradykinesia, muscular rigidity, rest tremor, postural and gait impairment) together with non-motor features (olfactory and autonomic dysfunction, cognitive impairment, psychiatric symptoms, sleep disorders, pain and fatigue) <sup>[14][15]</sup>. As a consequence, complex clinical pictures lead to numerous PD subtypes which can show different clinical profiles and disease progression <sup>[16]</sup>. This paragraph provides a review of actual fNIRS applications either to assess the effects of interventional procedures referred to Deep Brain Stimulation (DBS) <sup>[17][18][19][20]</sup> or to investigate clinical phenotypes associated to PD during usual and dual task (DT) walking <sup>[21][22][23][24][25][26][27][28]</sup>. Most of available articles investigated the cortical activity of the prefrontal cortex (PFC), while only a single study considered the primary motor cortex (PMC) as the region of interest [18]. Finally, one article employed a high-density probe configuration to simultaneously investigate brain activity associated to temporal and occipital cortices <sup>[20]</sup>.

### 2.1. Deep Brain Stimulation

The use of fNIRS and NIRS in the context of DBS is based on the underlying hypothesis that the stimulation delivered at

basal ganglia level can modulate the global cortical activity, which in turn is reflected by the neurovascular activity [18]. Sakatani et al. [17] were the first to employ NIRS for assessing changes in PFC oxygenation induced by frequency- and intensity-varying stimulation of the thalamic nucleus ventralis intermedius and globus pallidus internus. They observed HbO<sub>2</sub> and HbR changes that were comparable to those induced by cognitive tasks, thus suggesting a possible interaction between the frontal lobes and the stimulated deep brain structures. More recently, Morishita et al. [18] investigated PMC cortical activity pre-operatively and at 1-month follow-up, while motor scores were assessed by means of the Unified Parkinson's Disease Rating Scale (UPRS). Results show pre- to post-operative improvement of the UPRS motor score and the group analysis of fNIRS revealed a post-operative cortical activity comparable to the pre-operative one though more confined to the motor cortex for HbO<sub>2</sub>. Mayer et al. [19] studied the effects of bilateral subthalamic nucleus stimulation on working memory functions. Early PD patients showed a reduced frontal activation with respect to the control group. Overall worsening of working memory performance was accompanied by an increased frontal activation under DBS and on-medication, while no modifications were observed with respect to medication states, thus suggesting a DBS-induced compensatory mechanism operating within the basal ganglia-prefrontal network. Finally, Eggebrecht et al. [20] utilized a new High-Density Diffuse Optical Tomography (HD-DOT) probe array to map distributed brain functions and resting-state networks in 3 patients undergoing subthalamic nucleus DBS. This study demonstrated that HD-DOT showed a reliable overlap with fMRI, thus suggesting that it can be used to provide individualized functional images when other traditional functional imaging modalities are unavailable.

## 2.2. Walking and Dual Walking Task

Most of reviewed studies in PD were focused on walking and DT conditions to investigate the extent of PFC motor vs. executive and cognitive dysfunction. The underlying hypothesis is that PFC compensates for the motor impairment, hence cortical activation can be considered as an overall index of cognitive load. Within the context of rehabilitation, the promotion of more localized cortical activation associated with executive-attentional functions could be a viable way to activate this compensatory mechanism [24]. Mahoney et al. [21] reported that PD patients showed greater PFC activation in order to successfully achieve the same level of postural stability with respect to age-matched HCs and other mild PD patients. Nieuwhof et al. [22] observed overall increases of HbO<sub>2</sub> levels in the PFC during three dual walking tasks (walking while counting forward, serially subtracting, reciting digit spans) compared to a rest condition (standing still). Another study by Cornejo et al. [23] found that both gait stability and PFC activation were enhanced when walking on a treadmill at a self-selected pace with respect to usual walking, suggesting that an external rhythmic pacing may reduce the cognitive mediation on gait. Stuart et al. [24] also found that ageing and pathology affect the PFC compensatory mechanism due to the cognitive control required to perform turning-in-place and walking tasks, while this effect is reduced once the action has begun.

Maidan et al. [25][26][27][28] carried out an extensive work to investigate the relationship between this compensation mechanism and PFC activation and promote strategies to reduce cognitive load. In 2015 the authors studied the interplay of PFC activation during freezing of gait (FOG), a common disturbance among PD patients, associated with anticipated and unanticipated turns, in order to distinguish opposite cognitive requests such as motor planning and reflexive responses [26]. Results revealed that frontal activation levels can be associated to different typologies of FOG. Successively, they studied PD patients without FOG and noticed increased activation during usual walking and a decrease during turning in the absence of cognitive load [25]. Increased activation was also found in a sub-group of patients with impaired ambulation, which further supports the role of PFC in motor-cognitive compensation. Yet another study suggested that a combination of motor and cognitive tasks - namely obstacle negotiation, usual, and DT walking—determined different involvements of PFC activity during gait in HCs and PD patients [28]. Finally, the same research group carried out a longitudinal randomized controlled trial to assess the effects of treadmill training (TT) alone or combined with virtual reality (VR) [27]. PFC activation during obstacle negotiation and DT walking was reduced in the combined TT-VR program condition. However, both experimental conditions induced an overall reduction in the rate of post-intervention falls and an improvement in gait performances, suggesting that simultaneous motor and cognitive training promotes the recruitment of more specific PFC areas.

## 3. Conclusion

In conclusion, the available literature supports the idea that fNIRS can be a viable tool to detect functional differences

between normal ageing and people affected by the most common chronic neurological disorders, including PD. We found that this technology is mainly employed for the characterization of the patients' clinical phenotype, whereas a systematic adoption of intervention-based monitoring still remains to be seen. Overall results open the scenario of fNIRS as low-cost and portable tool to monitor cerebral plasticity during disease progression in order to promote subject-specific intervention strategies.

## References

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1. Eli Carmeli; Physical Therapy for Neurological Conditions in Geriatric Populations. *Frontiers in Public Health* **2017**, *5*, null, 10.3389/fpubh.2017.00333.
2. Christopher J. Poulos; Antony Bayer; Lauren Beaupre; Linda Clare; Roslyn G. Poulos; Rosalie H. Wang; Sytse Zuidema; Katherine S. McGilton; A comprehensive approach to reablement in dementia. *Alzheimer's & Dementia: Translational Research & Clinical Interventions* **2017**, *3*, 450-458, 10.1016/j.trci.2017.06.005.
3. Fary Khan; Bhasker Amatya; Mary Galea; Roman Gonzenbach; Jürg Kesselring; Neurorehabilitation: applied neuroplasticity. *Journal of Neurology* **2016**, *264*, 603-615, 10.1007/s00415-016-8307-9.
4. Ayse A Küçükdeveci; A Tennant; G Grimby; Franco Franchignoni; Strategies for assessment and outcome measurement in Physical and Rehabilitation Medicine: An educational review. *Journal of Rehabilitation Medicine* **2011**, *43*, 661-672, 10.2340/16501977-0844.
5. Shannon L. Risacher; Andrew J. Saykin; Neuroimaging Biomarkers of Neurodegenerative Diseases and Dementia. *Seminars in Neurology* **2013**, *33*, 386-416, 10.1055/s-0033-1359312.
6. Vanessa Scarapicchia; Cassandra Brown; Chantel Mayo; Jodie R. Gawryluk; Functional Magnetic Resonance Imaging and Functional Near-Infrared Spectroscopy: Insights from Combined Recording Studies. *Frontiers in Human Neuroscience* **2017**, *11*, 419, 10.3389/fnhum.2017.00419.
7. Meryem A. Yücel; Juliette J. Selb; Theodore J. Huppert; Maria Angela Franceschini; David A. Boas; Functional Near Infrared Spectroscopy: Enabling routine functional brain imaging. *Current Opinion in Biomedical Engineering* **2017**, *4*, 78-86, 10.1016/j.cobme.2017.09.011.
8. D T Delpy; M Cope; P Van Der Zee; S Arridge; S Wray; J Wyatt; Estimation of optical pathlength through tissue from direct time of flight measurement. *Physics in Medicine & Biology* **1988**, *33*, 1433-1442, 10.1088/0031-9155/33/12/008.
9. Simone Cutini; Sara Basso Moro; Silvia Bisconti; Functional near Infrared Optical Imaging in Cognitive Neuroscience: An Introductory Review. *Journal of Near Infrared Spectroscopy* **2012**, *20*, 75-92, 10.1255/jnirs.969.
10. Gilles Allali; Helena M. Blumen; Hervé Devanne; Elvira Pirondini; Arnaud Delval; Dimitri Van De Ville; Brain imaging of locomotion in neurological conditions. *Neurophysiologie Clinique* **2018**, *48*, 337-359, 10.1016/j.neucli.2018.10.004.
11. Vera Gramigna; Giovanni Pellegrino; Antonio Cerasa; Simone Cutini; Roberta Vasta; Giuseppe Olivadese; Iolanda Martino; Aldo Quattrone; Near-Infrared Spectroscopy in Gait Disorders: Is It Time to Begin?. *Neurorehabilitation and Neural Repair* **2017**, *31*, 402-412, 10.1177/1545968317693304.
12. E. Ray Dorsey; Alexis Elbaz; Emma Nichols; Foad Abd-Allah; Ahmed Abdelalim; Jose C. Adsuar; Mustafa Geleto Ansha; Carol Brayne; Jee-Young J Choi; Daniel Collado-Mateo; et al.Nabila DahodwalaHuyen Phuc DoDumessa EdessaMatthias EndresSeyed-Mohammad FereshtehnejadKyle J ForemanFortune Gbetoho GankpeRahul GuptaGraeme J. HankeySimon I. HayMohamed I HegazyDesalegn T. HibstuAmir KasaeianYousef KhaderIbrahim KhalilYoung-Ho KhangYun Jin KimYoshihiro KokuboGiancarlo LogroscinoJoão MassanoNorlinah Mohamed IbrahimMohammed A. MohammedAlireza MohammadiMaziar Moradi-LakehMohsen NaghaviBinh Thanh NguyenYirga Legesse NirayoFelix Akpojene OgboMayowa Ojo OwolabiDavid M. PereiraMaarten J PostmaMostafa QorbaniMuhammad Aziz RahmanKedir T. RobaHosein SafariSaeid SafiriMaheswar SatpathyMonika SawhneyAzadeh ShafieesabetMekonnen Sisay ShiferawMari SmithCassandra E I SzoekiRafael Tabarés-SeisdedosNu Thi TruongKingsley Nnanna UkwajaNarayanawamy VenketasubramanianSantos VillafainaKidu Gidey WeldegewergsRonny WestermanTissa WijeratneAndrea S. WinklerBach Tran XuanNaohiro YonemotoValery L FeiginTheo VosChristopher J L Murray Global, regional, and national burden of Parkinson's disease, 1990–2016: a systematic analysis for the Global Burden of Disease Study 2016. *The Lancet Neurology* **2018**, *17*, 939-953, 10.1016/s1474-4422(18)30295-3.
13. Alves, G.; Pedersen, K.F.; Pedersen, K.F. Epidemiology of Parkinson's disease. *J. Neurol.* 2008, *5*, 525–535.
14. Lorraine V. Kalia; Anthony E. Lang; Evolving basic, pathological and clinical concepts in PD. *Nature Reviews Neurology* **2016**, *12*, 65-66, 10.1038/nrneurol.2015.249.
15. Lorraine V Kalia; Anthony E Lang; Parkinson's disease. *The Lancet* **2015**, *386*, 896-912, 10.1016/s0140-6736(14)61393-3.
16. Eduardo De Pablo-Fernández; Andrew J. Lees; Janice L. Holton; Thomas T. Warner; Prognosis and Neuropathologic Correlation of Clinical Subtypes of Parkinson Disease. *JAMA Neurology* **2019**, *76*, 470-479, 10.1001/jamaneurol.2018.4377.
17. Kaoru Sakatani; Yoichi Katayama; Takamitsu Yamamoto; Susumu Suzuki; Changes in cerebral blood oxygenation of the frontal lobe induced by direct electrical stimulation of thalamus and globus pallidus: a near infrared spectroscopy study. *Journal of Neurology, Neurosurgery & Psychiatry* **1999**, *67*, 769-773, 10.1136/jnnp.67.6.769.
18. Takashi Morishita; Masa-Aki Higuchi; Kazuya Saita; Yoshio Tsuboi; Hiroshi Abe; Tooru Inoue; Changes in Motor-Related Cortical Activity Following Deep Brain Stimulation for Parkinson's Disease Detected by Functional Near Infrared Spectroscopy: A Pilot Study.

*Frontiers in Human Neuroscience* **2016**, *10*, 1-9, 10.3389/fnhum.2016.00629.

19. Jutta S. Mayer; Joseph Neimat; Bradley S. Folley; Sarah K. Bourne; Peter E. Konrad; David Charles; Sohee Park; Deep brain stimulation of the subthalamic nucleus alters frontal activity during spatial working memory maintenance of patients with Parkinson's disease. *Neurocase* **2016**, *22*, 369-378, 10.1080/13554794.2016.1197951.
20. Adam T. Eggebrecht; Silvina L. Ferradal; Amy Robichaux-Viehoever; Mahlega S. Hassanpour; Hamid Dehghani; Abraham Z. Snyder; Tamara Hershey; Joseph P. Culver; Mapping distributed brain function and networks with diffuse optical tomography. *Nature Photonics* **2014**, *8*, 448-454, 10.1038/nphoton.2014.107.
21. Jeannette R. Mahoney; Roe Holtzer; Meltem Izzetoglu; Vance Zemon; Joe Verghese; Gilles Allali; The role of prefrontal cortex during postural control in Parkinsonian syndromes a functional near-infrared spectroscopy study. *Brain Research* **2016**, *1633*, 126-138, 10.1016/j.brainres.2015.10.053.
22. Freek Nieuwhof; Miriam F. Reelick; Inbal Maidan; Anat Mirelman; Jeffrey M. Hausdorff; Marcel G. M. Olde Rikkert; B. R. Bloem; Makii Muthalib; Jurgen A. H. R. Claassen; Measuring prefrontal cortical activity during dual task walking in patients with Parkinson's disease: feasibility of using a new portable fNIRS device.. *Pilot and Feasibility Studies* **2016**, *2*, 59, 10.1186/s40814-016-0099-2.
23. Pablo Cornejo Thumm; Inbal Maidan; Marina Brozgol; Shiran Shustak; Eran Gazit; Shirley Shema Shiratzki; Hagar Bernad-Elazari; Yoav Beck; Nir Giladi; Jeffrey M. Hausdorff; et al. Anat Mirelman Treadmill walking reduces pre-frontal activation in patients with Parkinson's disease. *Gait & Posture* **2018**, *62*, 384-387, 10.1016/j.gaitpost.2018.03.041.
24. Samuel Stuart; Valeria Belluscio; Joseph F. Quinn; Martina Mancini; Pre-frontal Cortical Activity During Walking and Turning Is Reliable and Differentiates Across Young, Older Adults and People With Parkinson's Disease. *Frontiers in Neurology* **2019**, *10*, 536, 10.3389/fneur.2019.00536.
25. Inbal Maidan; Hagar Bernad-Elazari; Nir Giladi; Anat Mirelman; Jeffrey M. Hausdorff; When is Higher Level Cognitive Control Needed for Locomotor Tasks Among Patients with Parkinson's Disease?. *Brain Topography* **2017**, *30*, 531-538, 10.1007/s10548-017-0564-0.
26. Inbal Maidan; Hagar Bernad-Elazari; Eran Gazit; Nir Giladi; Jeffrey M. Hausdorff; Anat Mirelman; Changes in oxygenated hemoglobin link freezing of gait to frontal activation in patients with Parkinson disease: an fNIRS study of transient motor-cognitive failures. *Journal of Neurology* **2015**, *262*, 899-908, 10.1007/s00415-015-7650-6.
27. Inbal Maidan; Freek Nieuwhof; Hagar Bernad-Elazari; Bastiaan R. Bloem; Nir Giladi; Jeffrey M. Hausdorff; Jurgen A. H. R. Claassen; Anat Mirelman; Evidence for Differential Effects of 2 Forms of Exercise on Prefrontal Plasticity During Walking in Parkinson's Disease. *Neurorehabilitation and Neural Repair* **2018**, *32*, 200-208, 10.1177/1545968318763750.
28. Inbal Maidan; Freek Nieuwhof; Hagar Bernad-Elazari; Miriam F. Reelick; Bas R. Bloem; Nir Giladi; Judith E. Deutsch; Jeffrey M. Hausdorff; Jurgen A. H. Claassen; Anat Mirelman; et al. The Role of the Frontal Lobe in Complex Walking Among Patients With Parkinson's Disease and Healthy Older Adults. *Neurorehabilitation and Neural Repair* **2016**, *30*, 963-971, 10.1177/1545968316650426.

## Keywords

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neurovascular coupling;fNIRS;neurological disease;Parkinson's Disease

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