Epidemiology of Parkinsonism and Parkinson’s disease in Sub-Saharan Africa: Nigerian profile

Parkinsonism and Parkinson’s disease (PD) are the most common movement disorders (neurological syndromes characterized by paucity of voluntary and automatic movements or excessive movements unrelated to muscle weakness or spasticity), and they are generally perceived to have lower prevalence and incidence in Sub-Saharan Africa (SSA) because of the relative youthfulness of the SSA population (<5% are over 65 years of age), under-recognition, paucity of published reports, and cultural perception of neurologic disorders generally as being part of normal ageing.[1]

In Nigeria, Africa’s most populous nation, the earliest detailed report on Parkinsonism and PD dates back to about 40 years ago when Osuntokun[2] described extra pyramidal disorders in 1.9% of 9600 patients with neurological conditions seen over a 12 year period (1957–69) at the University College Hospital Ibadan, Southwestern Nigeria. Most subsequent reports on this syndrome from Nigeria have emanated similarly from the same region, the current report by Owolabi et al[3] being the very first from the expansive northern region of the country.

In 1978, Osuntokun and Bademosi described 217 Nigerians with Parkinsonism, of whom 38% were classified as PD.[1,3,4] However, in a recent 10 year review of Parkinsonism as seen in the major tertiary hospital in Lagos, the Nigerian commercial nerve center in the Southwest, 79% of the 124 patients reviewed had PD[4] a figure very close to the 83% documented from the current review from Kano, Northern Nigeria.[1] The increased proportion of PD is thought to be due to recent refinements in the diagnostic criteria for PD and Parkinsonism syndromes.[1] The spectrum of secondary Parkinsonism and Parkinson-plus syndromes documented from these hospital series and other anecdotal reports in Nigeria includes Parkinsonism secondary to Wilson’s disease, cerebral vascular disease, drugs, trauma, tumor excision, typhoid septicemia, and human immunodeficiency virus (HIV) infection. Others are multiple system atrophy, dementia with Lewy bodies, progressive supranuclear palsy, hemiparkinsonism-hemiatrophy, juvenile parkinsonism with dystonia and hemiatrophy and primary amyloidosis with parkinsonism.[1,3,4]

A pioneering epidemiologic survey of PD from a rural community in Southwestern Nigeria yielded an age-adjusted prevalence rate of 67 per 100,000 (above 39 years),[5] while a case-control study on risk factors found blacksmithing (OR 8.0 95% CI 1.3- 50.7)[6] to be significantly associated with PD. There are no data on exposure to pesticides, herbicides, rural living, and well water use. A preliminary analysis of the genetic contributions to PD in Nigerians explored the role of mutations in LRRK2, PRKN, and ATXN3 genes and found some novel but non-pathogenic mutations.[7]

Neuropathological studies involving brain tissue samples of neurologically normal Nigerians showed number of melanized neurons[8] and Lewy body pathology burden[9] similar to findings in Caucasian populations implying that the frequency of PD-related pre-symptomatic neuropathology (and indirectly the risk of PD) among Nigerian and Caucasian populations may be similar, but the effect of yet unidentified genetic factors, environmental exposures, differences in population structure and reduced life expectancy of Nigerians (about 50 years) may explain the differences in the prevalence and incidence rates observed between the two population groups.

Findings from this seminal work of Owolabi et al[3] reveal some shared features of PD in both Northern and Southern Nigeria.[2-4] These include onset of disease in the sixth decade, male preponderance, under-diagnosis, late presentation, higher mortality, higher proportion of tremor dominant PD, and relatively low proportion of young onset and familial disease compared to Western
populations. However, the relatively older age of onset in the current study\(^1\) compared to older studies\(^{1,2,3,4}\) may be a subtle evidence of ongoing epidemiologic transition.

Non-motor manifestations including cognitive dysfunction, autonomic dysfunction, psychiatric symptoms (especially apathy, depression, and anxiety), gastrointestinal and sudomotor disturbances all of which significantly impact patients’ quality of life have also been described among Nigerian PD patients.\(^{10,11}\)

The management of PD in Nigeria is fraught with challenges which include high cost and narrow spectrum of available treatment options and medications, paucity of movement disorders specialists, late presentation of available treatment options and medications, and high caregiver burden.\(^{11,2,4}\)

Overall, though the current report\(^3\) represents an important milestone in the annals of movement disorders in Nigeria and SSA, there still exists knowledge gaps in several aspects including current prevalence and incidence, natural history, quality of life, cost of care, treatment outcomes and clinico-genetic-pathological correlations of PD and other movement disorders. Intra- and international collaborations are needed to enhance research and capacity building while advocacy through public awareness campaigns and support groups will improve the recognition and adequate holistic treatment of Parkinsonism and PD in rural and urban SSA.

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