Parkinson's disease and degenerative parkinsonism: relationship between clinical and neurobiological aspects

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<table>
<thead>
<tr>
<th>Section</th>
<th>Page</th>
</tr>
</thead>
<tbody>
<tr>
<td>Introduction</td>
<td>3</td>
</tr>
<tr>
<td>1. Dementia in Parkinson's disease</td>
<td>5</td>
</tr>
<tr>
<td>1.1 Clinical Features</td>
<td>5</td>
</tr>
<tr>
<td>1.2 Neuropathology of cognitive disorders in Parkinson's disease</td>
<td>8</td>
</tr>
<tr>
<td>1.3 Alzheimer-like pathological changes</td>
<td>9</td>
</tr>
<tr>
<td>1.4 Vascular white matter lesion load</td>
<td>10</td>
</tr>
<tr>
<td>2. Monitoring cerebral blood flow: cerebral autoregulation and neurovascular coupling</td>
<td>12</td>
</tr>
<tr>
<td>2.1 Cerebral Autoregulation</td>
<td>12</td>
</tr>
<tr>
<td>2.2 Neurovascular Coupling</td>
<td>13</td>
</tr>
<tr>
<td>3. Cerebral Vasoreactivity: methods of evaluation</td>
<td>17</td>
</tr>
<tr>
<td>3.1 Vasoreactivity to CO2</td>
<td>18</td>
</tr>
<tr>
<td>4. Cerebral hemodynamics and cognitive functions</td>
<td>21</td>
</tr>
<tr>
<td>5. Objectives of the study</td>
<td>27</td>
</tr>
<tr>
<td>6. Materials and Methods</td>
<td>27</td>
</tr>
<tr>
<td>6.1 MCI in Parkinson's disease</td>
<td>28</td>
</tr>
<tr>
<td>6.2 Eco-color doppler of the neck vessels, transcranial Doppler ultrasound at baseline and Breath-Holding unit test</td>
<td>31</td>
</tr>
<tr>
<td>7. Statistical Analysis</td>
<td>32</td>
</tr>
<tr>
<td>8. Results</td>
<td>32</td>
</tr>
<tr>
<td>9. Discussion</td>
<td>36</td>
</tr>
<tr>
<td>10. Conclusions</td>
<td>40</td>
</tr>
<tr>
<td>References</td>
<td>41</td>
</tr>
</tbody>
</table>
Introduction

Parkinson's disease (PD) has been classically considered as discrete disease entity, characterized clinically by rigidity, tremor and bradykinesia, and from the neuropathological point of view, the neuronal degeneration of the substantia nigra of the midbrain and the presence in the neuronal population surviving in this same region of inclusions containing α-synuclein (α-syn) in the cell bodies and synaptic processes (respectively known as Lewy bodies and Lewy neurites).

However remain valid this core descriptive, the progressive acquisitions on different aspects of Parkinson's disease tend to recognize now the disease as a more complex clinical-pathologic entity: a "protean" multiorgan disease caused by the synergistic convergence of different pathophysiological moments.

Consistent with this vision "extended" the degenerative process, the cognitive impairment in Parkinson's disease is considered to be the result of complex interactions between neurobiological and pathophysiological variables, rather than the expression of a linear degenerative process.

Clinically, some nuanced cognitive dysfunction are typically found already in the early stages of Parkinson's disease; such dysfunctions tend to evolve in a typical cognitive impairment only in a subset of patients to progress, in some cases, within a framework of dementia.

The cognitive alterations most frequently found at the onset of the disease are due to impairment of the fronto-striatal systems, manifested by alterations of executive functions, planning, goal-directed behaviors and working-memory\(^1\).

An initial deterioration of visuo-spatial expression of an involvement by the pathological process of the posterior brain structures, is another aspect of cognitive potentially involved already in the early stages of the disease, although they are not necessarily the expression of a full-blown dementia.
Extremely heterogeneous even the progression of cognitive impairment in patients with Parkinson's disease.

Based on the functions involved, we observe at least two patterns of progression: a relatively slow decline in cognitive function of "nature" fronto-striatal and a more rapid decline when to be involved are the functions classically attributed to the posterior structures\(^{(2)}\).

However it is plausible that such a division would be an oversimplification; in fact the cognitive decline in patients with Parkinson's disease has a more considerable variability, probably reflecting the complex interaction of different substrates pathophysiological, neuropathological and genetic studies.
1. Dementia in Parkinson's disease

Cognitive impairment in Parkinson's disease is a primary determinant associated with a significant reduction in quality of life, with a greater risk of developing psychiatric disorders, with a greater risk of institutionalization and reduced life expectancy.

Studies in patients with Parkinson's disease to accept the frequency showed considerable variability.

Two studies have investigated the incidence of dementia in patients with Parkinson's disease in the period pre-levodopa: Lewy (1923) observed that 54 of 70 patients investigated (77%) developed dementia and Monroe reported that the painting in approximately one third of patients Parkinson's disease.

A recent review estimated the prevalence of Parkinson-dementia (PDD) in 40% of patients in a sample of 4336 individuals included in 27 studies. However, with the use of different methodological criteria, other authors have reported a given prevalence of 31.3% . (3)

How common is dementia in Parkinson's disease is still a matter of discussion.

Two longitudinal studies are now available: a study of 149 patients on PD de-novo Sydney where it was reported a prevalence of dementia of 28% after 5 years, 48% after 15 years and 83% after 20 years (4) and the study Stavanger where the prevalence of PDD increased up to 78% after 8 years of observation (5).

1.1 Clinical Features

A certain degree of cognitive impairment in PD is potentially present in the early stages, can also be observed in patients not on dopaminergic therapy, although it is generally not very noticeable to the patient and the neurologist.

Muslimovic at al. (6) assessed the neuropsychological profile in a cohort of 129 patients De-novo suffering from Parkinson's disease, subjecting them to an
extensive neuropsychological battery. The authors found cognitive deficits in 23.5% of patients with Parkinson's disease; performance on tests of psychomotor speed, language functions, features attentional/executive, memory functions and visuo-spatial abilities are aspects results in deficit. While in the domain of language disorders were relatively rare (22%), alterations in attention functions/execution were found in nearly all of the subgroup of patients whose cognitive deficits emerged.

The same authors emphasized as the age of onset of the disease is an independent risk factor in the development of cognitive impairment in patients with Parkinson's disease. The figure is in according with previous studies indicate that it is predictive of the development of cognitive impairment in patients with Parkinson's disease, age of onset of the disease rather than the duration.

In a study by P. Marini et al.\textsuperscript{(7)} on a small group of patients with Parkinson's disease de-novo has been observed that some executive dysfunction tend to decline after the administration of levodopa.

A recent study\textsuperscript{(8)} investigated the association between cognitive and motor symptoms in a cohort of patients, identifying a strong correlation in patients suffering from non-demented Parkinson's disease between cognitive deficits and non-motor symptoms levodopa-responsive (impaired gait, postural abnormalities and dysarthria); the data therefore stresses the important role of extra nigro-striatal systems in the development of cognitive impairment.

The initial cognitive impairment in patients with Parkinson's disease, non-demented, include a wide range of ailments, mostly in relation to deficit of frontal-executive functions; they may be detected through the use of specific psychometric instruments:

- Verbal fluency and phonemic alternating
- Verbal fluency "phenomenon tip-of-the-tongue": proof of fluency phonetic
- Sustained attention: Stroop test and Wisconsin card sorting test
- Working memory: digit span backward
- Speed of cognitive processing: Digit Symbol test.
- Verbal and visual memory: words and figure of Rey
- Test of Watch
- Visuo-spatial ability: mental rotation of shapes, orientation lines.

With the progression of the disease, in addition to a further deterioration of the domains mentioned above, may arise deficit regarding the following areas:
- Verbal fluency semantics: semantic fluency by category
- Name
- Recognition
- Constructional apraxia and a copy of the figure: Pentagons.

However the cognitive impairment in Parkinson's disease remains extremely heterogeneous in terms of both temporal progression of that dysfunctional profile. Besides the already mentioned clinical phenotype of Parkinson's disease, also the presence of psychiatric disorders is a risk factor for the development of cognitive impairment; in particular visual hallucinations have been associated with alterations in cognitive profile in the domains of verbal learning and attention\(^9\). Comparing groups of patients suffering from non-demented Parkinson's disease who presented with visual hallucinations (VH) with a control group of patients suffering from Parkinson's disease, but without VH, some authors have observed the correlation of the psychiatric disorder with different non-motor symptoms, such as depression, anxiety and sleep disorders.

Cognitive deficits, as well as neuropsychiatric disorders, are both believed to be extra nigro-striatal symptoms. This aspect supports the hypothesis that patients
with PD and visual hallucinations, are a subgroup of patients at high risk for developing dementia, even when cognitively intact. The symptom is in fact due to a complex interaction of various neurotransmitter systems. Similarly, the potential role of an imbalance between different neurotransmitter pathways is also suggested in the genesis of cognitive impairment, as supported by recent scientific evidence. Monastero et al. \(^\text{[10]}\) assessed the neuropsychiatric profile of 410 patients in a large cohort of patients with non-demented Parkinson's disease: the authors observed that neuropsychiatric disorders were more frequent in patients with PD and mild cognitive impairment, in particular in patients in whom emerged dysfunction borne of multiple domains.

### 1.2 Neuropathology of cognitive disorders in Parkinson's disease:

**a-Synucleinopathia**

The presence of intracytoplasmic neuronal inclusions (inclusions known as Lewy Bodies) and confined to neuronal processes (known as Lewy neurites, LN) is the pathognomonic neuropathological feature of Parkinson's disease. The involvement of the substantia nigra with the destruction of dopaminergic neurons in the pars compacta is unanimously considered to be the hallmark of Parkinson's disease. A study carried out to examine the neuropathological substrates of cognitive dysfunction in Parkinson's disease\(^\text{[11]}\) indicates that the extension of cortical neuropathological inclusions typical of so-called "synucleinopathies" is the element that most correlates with cognitive impairment; the data would therefore be in line with the hypothesis of Braak on the progression of alpha-synuclein.
1.3 Alzheimer-like pathological changes

Although several studies indicate that cortical neuropathological alterations similar to those seen in Alzheimer's disease are considered an important factor in the development of dementia in Parkinson's disease, the real meaning of plaques of beta-amyloid (Aβ) and neurofibrillary tangles in dementia Parkinson's disease is still the subject of research (12), since there is no consistent correlation between cortical Aβ load and assessments of cognitive deficits in patients with clinically determined PDD.

However in the study by Irwin (2012) previously cited, the authors observed through a sub-analysis of the PDD group, the presence of beta-amyloid plaques and neurofibrillary tangles may influence cognitive impairment and disease progression in a subset of patients; specifically the presence of Alzheimer-like changes seem associated with cognitive decline in the group of patients at onset of illness in old
age and with a reduced free interval between onset of motor symptoms and onset of cognitive impairment.

Figure from Kalaitzakis ME (Acta Neuropathol 2009). Immunohistological analysis for the deposition of b-amyloid in the caudate nucleus. A) Numerous deposition of Ab in the caudate nucleus in a case PD with dementia. B) caudate nucleus, without Ab deposits in the event a non-demented PD. Magnification 920 Parkinson's disease PD, Ab amyloid b peptide.

**1.4 Vascular white matter lesion load**

The white matter lesion load has been classically associated with motor disability and cognitive dysfunction in elderly subjects not suffering from dementia or Parkinson's disease; therefore vascular disease may have an additive effect on clinic sintomotology in Parkinson's disease, including cognitive deficits. A review of Bohnen NI\(^{(13)}\) highlights this concept.

Figure by Bohnen NI, Nat Rev Neurol, 2011

However the impact of vascular disease on cognitive dysfunction in Parkinson's disease has not been widely accepted.
A recent study by Gonzalez-Redondo R.\(^{14}\) analyzed the vascular lesion load by means of magnetic resonance brain imaging (MRI) in a large cohort of patients with PD; to identify a possible relationship between silent infarcts and dementia or MCI in this population has been structured assessment transverse and longitudinal. The authors found no association between white matter hyperintensities and global cognitive profile in patients with Parkinson's disease; also the presence of vascular lesion load was not a negative prognostic index after a mean follow-up of 30 months.
2. Monitoring cerebral blood flow: cerebral autoregulation and neurovascular coupling

Cerebral blood flow (CBF) is regulated by two major homeostatic physiological systems, cerebral autoregulation that maintains a relatively constant CBF to changes in systemic blood pressure and cerebral perfusion pressure and neurovascular coupling, consisting of the neurovascular unit, which adapts the cerebral blood flow to neuronal activity in the brain. These two systems work in part with the same physiological mechanisms.

2.1 Cerebral Autoregulation

The cerebral circulation has a fundamental homeostatic mechanism called self-regulation that is the ability to maintain cerebral blood flow (CBF) relatively constant in terms of variability acute or chronic systemic blood pressure and cerebral perfusion pressure, ensuring, within certain limits, a adequate metabolic supply to the brain tissue; therefore unlike other tissues exempt from autoregulation, in the brain there is a nonlinear relationship between systemic arterial pressure and flow. The self-regulating mechanism is able to maintain the cerebral flow relatively constant within the limits of mean systemic arterial pressure between 60 and 150 mmHg; beyond these limits the flow becomes passively dependent on perfusion pressure, causing ischemia if it is reduced or if edema increases. The principal mechanism employed to keep the cerebral blood flow constant when the perfusion pressure is represented mainly from changes in the local vascular resistance according to the formula:

Regional blood flow (rCBF) = cerebral perfusion pressure (CPP) / resistance cerebrovascular regional (rCVR).
2.2 Neurovascular Coupling

The Neurovascular Coupling is the process in which the neurovascular unit, represented by complex-vessel wall endothelium of small blood vessels, neurons, astrocytes and glial cells interact with each other by modulating the complex relationship between neuronal activity, hemodynamic factors and intracellular signals. These interactions allow you to synchronize the neuronal brain perfusion with the request, allowing an adequate metabolic supply to the brain areas in relation to the functional demand; in addition to the trophic support the neurovascular unit is involved in immune surveillance in trade at the level of the blood-brain barrier and hemostatic balance.

Currently the physiological mechanisms implicated in the regulation of regional cerebral blood flow are represented by:

. myogenic mechanism
. neurogenic mechanism
. metabolic mechanism

These mechanisms may act alone or in synergy with each other.

In mechanism myogenic vascular tone is modified in response to the mechanical tension on the vessel wall induced by the pressure and the amount of flow. The small vessels and arterioles (vessels more accountable vascular resistance) constrict or dilate in response respectively to an increase or reduction in the transmural pressure gradient with every heartbeat. Increased pressure within the vessel mechanically generate a greater circumferential wall tension and relax the smooth muscle fibers of the muscle layer, thus resulting in depolarization of the cell membrane and contraction of the muscle cell. An increase in flow within the vessel generates a voltage increase grazing (shear stress) on endothelial cells, resulting in
the relax of smooth muscle cell with a mechanism Nitric oxide (NO) -mediate. The flow-induced muscle relaxation allows to answer the need for increased local flow. The metabolic regulation provides that changes in molecules and metabolic substrates such as O2, CO2, ATP, glucose, glutamate, at the level of the tissue microenvironment generate vessel regulators regional adaptations, physiological condition exists as a direct relationship between cerebral blood flow and cerebral metabolism. The increase of neuronal activity in physiological conditions produces an increase in production and release of glutamate, adenosine triphosphate (ATP) and an increase in the activity of cyclooxygenase-2. The activation of NMDA receptors for glutamate increases the production of NO (vasodilator) to increase the activity of Nitric Oxide Synthase. Also through the glutamate binding to its receptors and metabotropic at the membrane causes an increase in astrocytic Phospholipas C, with an increase in consensual inositol triphosphate, release of calcium from the sarcoplasmic reticulum and activation of calcium-dependent potassium channels. The resulting in hyperpolarization of the cell membrane, caused by the leakage of potassium ions will be responsible for the reduced contractility of the muscle cell itself. The COX-2 resulting in an increase of prostaglandins in the extracellular space produces vasodilation. The increase of ATP determines increase in the concentration of adenosine that in turn activates the receptors A2b placed both on the membrane astrocytic that the membrane of the smooth muscle cell. The activation of these receptors leads to the increase of the concentration of cAMP with consequent vasodilatation. The CO2 is believed to work with PH-mediated mechanism. The adjustment uses a neurogenic component extrinsic and intrinsic.

Extrinsic component: Several studies have demonstrated the innervation of vessels extraparenchymal located at the base and the surface of the brain, the ganglia of origin of these endings perivascular and vasoactive substances released from the jar. It is believed that the extrinsic innervation finds its origin in the superior cervical
ganglion of the sympathetic fibers which originate containing norepinephrine and neuropeptide Y from the sphenopalatine and otic ganglion from which they came parasympathetic fibers containing vasoactive intestinal peptide (VIP) and nitric oxide synthase (NOS) and the trigeminal ganglion, from which originate sensory fibers containing calcitonin gene-related peptide (CGRP), substance P, the neurokinin A and pituitary adenylate cyclase-activating polypeptide. Once inside these vessels into the brain parenchyma lose the autonomic component and finished the spaces of Virchow-Robin vessels respond to a central neuronal control (intrinsic innervation). The role of the sympathetic system is mainly linked to the ability to move the upper limit of the curve of self-regulation in response to increases in systemic arterial pressure in the protection of brain tissue. The parasympathetic system, which mediates vasodilator responses, does not appear to play a significant role on the physiological mechanism of autoregulation of cerebral blood flow while it seems implicated in pathological conditions such as ischemic stroke and migraine. The trigeminal-vascular system, whose fibers are the only sensory innervation of brain vessels, appears to mediate CGRP-mediated vasodilator responses to vasoconstrictor stimuli. The trigeminal vascular afferents play an important role in the pathogenesis of migraine.

Intrinsic component: the intrinsic regulation of small intraparenchymal cerebral vessels is mediated by the neurovascular unit, which receives nerve fibers from basic subcortical systems and local cortical interneurons, although it should be noted that it is not yet well-defined neuronal network that acts functionally complete in synergy with the neurovascular unit. Neural networks, to date the best studied, which project to the small cerebral vessels and also to the surrounding astrocytes are cholinergic pathways from the basal nucleus of Meynert, noradrenergic pathways from the locus coeruleus and serotonergic pathways from the raphe nucleus. Electrical or chemical stimulation of these nuclei elicits profound
changes in cortical cerebral blood flow. M5 muscarinic receptors present on the vascular wall (endothelium and smooth muscle cells) that astrocytes mediate vasodilator responses.

Serotonergic 5-HT 1b median vasoconstrictive responses. It been shown that stimulation of noradrenergic neurons of the locus coerulus causes reduction of cortical CBF and that the target of perivascular cortical noradrenergic fibers is represented mainly by astrocytes. It is also shown that by stimulating specific subsets of cortical GABAergic interneurons are generated vasoconstrictive or vasodilatory responses of local microvessels; in particular interneurons producing VIP or NOS causing vasodilation, while those producing somatostatin caused vasoconstriction.

There are several conditions associated with damage to cerebral autoregulation (CA) and neurovascular coupling (NVC): vascular risk factors such as hypertension,
diabetes mellitus, smoking, obesity, stroke and degenerative diseases such as the Alzheimer’s disease.

**3. Cerebral Vasoreactivity: methods of evaluation**

Cerebral Vasoreactivity (CVR) reflects the ability of small cerebral vessels to adapt to changes in the microenvironment, to metabolic demand and to pressure changes, through the regulation of local blood flow. In neuroscience, there are several non-invasive techniques and vasomotor stimuli for the estimation of CVR; between the techniques include the TCD (transcranial Doppler), PET (positron emission tomography), SPECT (Positron Emission single-photon), the NIRS (infrared spectroscopy), the CASL MRI 3D (3D-continuous arterial spin labeling), the MRI BOLD (blood oxygenation level dependent) and Angio-RM 2D time of flight.
These techniques, however, are based on different parameters (blood flow or velocity of flow at the level of the great vessels, volume of flow, tissue perfusion), use different variables and have a different temporal and spatial resolution. Consequently, the estimation of CVR may differ widely between the different approaches. The main vasoactive stimuli are represented by chemical stimuli (CO2, O2), pharmacological (acetazolamide) or causing a transient reduction in blood pressure. It is believed that CO2 represents the stimulus more appropriate in terms of ease of use and in terms of standardization.

3.1 Vasoreactivity to CO2
Carbon dioxide (CO2) is a potent vasodilatory stimulus that modulates vascular tone during normal breathing during sleep and during hyperventilation. The CO2-VCR is measured in terms of changes in cerebral blood flow (CBF) or flow velocity (BFV) in response to vasoconstrictive stimuli (hypocapnia and hyperventilation) or vasodilators (hypercapnia, acetazolamide). The VCR is calculated as the percentage change in flow due to changes in CO2. Blood flow increases linearly over the physiological range of CO2 between 25-50 mmHg.
The method most commonly used in the clinic to measure the CO2-VCR is transcranial Doppler for the ease of execution. The advanced imaging techniques sovramenzionate than the TCD used to assess vasoreactivity in specific brain areas and vascular but lack in temporal resolution. The tests used for the assessment of CO2-VCR using transcranial doppler are represented by the Breath-holding, the test with a mixture of CO2, by re-breathing and test with acetazolamide.

Breath-holding test: this test for the measurement of CVR was demonstrated for the first time since Ratnatunga and Adiseshiah in 1990 and subsequently used clinically to assess the risk of stroke in pz suffering from carotid artery stenosis.\(^{15,16}\)

Methodology: after a period of normoventilation, asks the patient to hold their breath for at least 30 seconds. It is important that the apnea beginning with a normal breath to avoid the phenomenon of Valsalva and therefore the increase of the intracranial pressure. Through a bilateral monitoring by TCD are measured values of mean flow velocity (VMF) at the level of the stretch M1 middle cerebral artery bilaterally in rest conditions and after 4 seconds of breath at the end apnea. During the experiment are monitored by a capnometer the pCO2, the blood pressure (BP) and heart rate (HR). The Breath Holding index (BHI) is obtained by dividing the percentage increase of the speed of flow that occurs during the BH for the period of time (seconds) in which the subject is able to hold his breath after a normal inhalation according to the formula:

\[
\text{BHI: } \left( \frac{\text{MFV}_{\text{BH}} - \text{MFV}_{\text{base}}}{\text{MFV}_{\text{base}}} \right) \times 100 \text{ / sec of BH}. 
\]

This method has been validated in numerous studies, it is also a non-invasive, well tolerated and easy to perform. The main limitations of the method are represented by:

. need for adequate cooperation of the subject.
. insufficient change in the pCO2 (prolonged apnea causes in fact an increase of only 3-4 mmHg pCO2)
. not perfect reproducibility: interindividual variability with which it establishes hypercapnia and apnea post does not make clear correlation between blood levels of PCO2 and time of apnea; apnea is not the only factor that influences the PCO2 "current" are called upon other factors such as the metabolism of the subject, the size of the lungs (influenced by age, sex, weight) and the presence of bronchopulmonary disease. However, the use of capnometer can sensitize the method.

Testing with inhalation of CO2: in the reactivity test is used to CO2 inhalation of mixtures of CO2 at a concentration that varies from 2, 5 or 7%.

By registering with bilateral TCD, are measured values of average velocity in the middle cerebral arteries of the two sides at rest and at the end of the test. During the experiment is also measured by a capnometer pCO2, and monitored the PA and FC. At each steady state value of MFV is compared with the value of the end-tidal CO2. The cerebral vasomotor response (VMR) to CO2 is calculated as the percentage increase of MFV measured after 90 seconds of inhalation of the mixture of CO2 compared to the rest condition in accordance with the formula:

\[ \frac{(MFV_{MFVCO2} - MFV_{base})}{MFV_{CO2}} \times 100. \]

The test also provides a simple method to CO2, non-invasive and validated.\(^{(17)}\)

The re-breathing consists in allowing the patient to breathe the same air that is exhaled within a closed system. With this method you do not have reached a steady state of blood concentration of CO2 and it is difficult to correlate changes in flow with changes in PaCO2.

The acetazolamide test consists of a test drug that involves the intravenous infusion of 500 mg or 2 g of acetazolamide, a carbonic anhydrase inhibitor, the enzyme that catalyzes both dissociation and the formation of carbonic acid (H2CO3); the inhibition of this enzyme is responsible for a marked cerebral acidosis. The reduction of brain pH thus obtained causes vasodilation. The effects are expressed within the
first minute of infusion and end after 30 minutes, reaching a plateau at 15 minutes. The drug in its intravenous formulation is not yet available in Italy. Compared to the previous two techniques, it is an invasive method, a longer duration and with important side effects.\(^{(18)}\)

In addition, another limitation inherent in the TCD is to give an overall measure of cerebrovascular reactivity going to sample the proximal vessels; methods of functional magnetic resonance imaging (fMRI) are able to analyze functional changes in regional cerebral perfusion.

4. Cerebral hemodynamics and cognitive functions

In recent years the relationship between cerebral hemodynamics and cognitive performance has become a major challenge in clinical practice in relation to the worldwide growth of the elderly population. The presence of cardiovascular risk factors in middle-aged individuals is considered to be a potential marker for the early identification of individuals at risk of dementia. Hypertension and diabetes mellitus are risk factors for vascular prevalent; hypertension affects more than one-third of the world population and 366 million people are suffering from diabetes. Among the subjects that have more than 65 years of age approximately 70% are suffering from hypertension and 26% diabetes. Although the pathophysiological link between cerebral hemodynamics, reduced cerebrovascular reserve and dementia is complex, several epidemiological studies, neurophysiological and clinical-instrumental showed that microvascular damage associated with hypertension, diabetes, and other vascular risk factors, advanced age is responsible alteration of the neurovascular coupling resulting in regional hypoperfusion and neurodegeneration. Genetic factors, inflammation, autonomic disorders, neurodegeneration (m. Alzheimer's, Parkinson-dementia, dementia with Lewy
bodies), ischemic and hemorrhagic stroke are other overriding factors in the
development of dementia.

Studies measuring CBF in response to hypercapnia with transcranial doppler\textsuperscript{(19)} and method of inhalation of 133 Xe\textsuperscript{(20)} showed reduced vasoreactivity in subjects with type 2 diabetes compared with controls healthy. Other studies have shown an increased pulsatility index in diabetic subjects highlighted an increase in vascular resistance and microvascular damage. Studies carried out with high-resolution MRI (8 TESLA) showed the damage of the small vessels, microinfarcts, microhemorrhages, iron deposits and diffuse leukoaraiosis (white matter hyperintensities WMHS) in subjects with hypertension.

From a histopathological point of view of the deep periventricular white matter lesions showed partial demyelination, astrogliosis, dilatation of the perivascular spaces, activated macrophages and fibrohyalinosis\textsuperscript{(21)}. Isolated lesions and spread of the deep white matter were associated with heart attacks arteriolar sclerosis incomplete and complete\textsuperscript{(22)}. The small vessel disease brain and the WMHS were associated with cognitive impairment in several studies\textsuperscript{(23,24)}; it is also considered that the same factors affecting the white matter may cause hypoperfusion at the level of the gray matter and functional disconnection cortico-subcortical and that this process may precede cognitive decline even many years. More studies have shown the strong association between WMHS with diabetes and hypertension. Although vascular inflammation related to glucose toxicity, lipotoxicity and neurotoxicity plays an important role in damaging the neurovascular unit and the blood-brain barrier by promoting neuronal loss. The combined action of vascular risk factors cause oxidative stress in the mitochondria and endoplasmic reticulum causing the release of pro-inflammatory cytokines and activation of pro-apoptotic pathways; these events cause endothelial dysfunction with decrease of NO synthesis and impaired cerebrovascular reactivity. Recently it has been shown that the
presence of intercellular adhesion molecule (ICAM) and vascular cell adhesion molecule (VCAM) were associated with impaired cerebral vasoreactivity and atrophy of the gray matter in several regions of the brain in type 2 diabetic standing and in hypertensive subjects \(^{(25)}\) and that adhesion molecules were also associated with worsening of cognitive sphere.

In recent years, much emphasis is given to the relationship between cerebrovascular damage and neurodegeneration, in particular on the role (causal summation or synergistic) of vascular risk factors and the neurovascular unit damage on neuronal loss.

There is extensive evidence epidemiological, clinical, neuropathological and pharmacological evidence that an "overlap" between vascular dementia and Alzheimer's disease (AD).

Vascular risk factors such as hypertension, diabetes, hypercholesterolemia, and obesity are associated with an increased risk of AD and accelerate the clinical expression. White matter lesions (white matter lesions) and lacunar infarcts are associated with amyloid plaques and neurofibrillary tangles in the more than 40% of elderly patients with dementia. The control of vascular risk factors in patients with AD slows the progression of WMLs \(^{(26)}\). The use of drugs angiotensin receptor blockers reduce the risk of AD \(^{(27)}\). Several studies have shown alterations of cerebral vasoreactivity in Alzheimer's dementia \(^{(28,29,30,31)}\); AD patients with impaired cerebrovascular reactivity to CO2 undergo a more rapid cognitive decline.

Experimental studies have shown that β-amyloid (Abeta) has important effects on the vessels and that hypoxia-ischemia is a potent modulator dell'amiloidogenesi brain \(^{(32)}\). The blood-brain barrier plays a critical role in the clearance of β-amyloid \(^{(33,34)}\). Genetic factors (such as the ApoE4 genotype) can alter the regulation of cerebral blood flow, contributing to white matter lesions (WMLs), and deposition of amyloid β-. Ischemia can promote the accumulation of β-amyloid through reduced
vascular clearance of this peptide; also ischemia-hypoxia promotes the cleavage of Aβ from APP precursor (Amyloid Precursor Protein) through the up-regulation of β-secretase and egressione; the increased production and decreased clearance of Aβ can increase the deposition in the brain and encourage the formation of amyloid plaques and amyloid angiopathy.

In turn, elevated brain levels of Aβ and amyloid plaques are found in patients with cerebrovascular insufficiency and cognitive impairment (35) and it is assumed that the damage of the vessel wall may damage the Aβ-induced vasomotor regulation and cerebrovascular reserve; in favor of this hypothesis, some studies have shown that focal ischemia produced large infarcts in mice over-expressing APP (36,37) in relation to Aβ-mediated vascular dysregulation. Other pathophysiological hypothesis argue that the deficit of the cholinergic neurons of the basal nucleus of Meynert produces a deficit of intrinsic neurogenic control of CBF.

From the most recent acquisitions can therefore be assumed that there is a synergistic pathogenetic or at least a synergistic effect between vascular damage and neurodegenerative disease.
Conceptual model of pathogenesis: the vascular risk factors through processes of oxidative stress and inflammation cause alteration neurovascular unit (resulting in hypoperfusion) and neurotoxicity with evolution in atrophy and functional decline (Novak V.2012).

Conceptual model of synergism between vascular dysfunction and neurodegeneration in AD. (38)
Limited data in the literature between cerebrovascular reactivity (VCR) and Parkinson's disease. The studies were structured framing the deficit cerebral autoregulation as a dysfunction of the autonomic system, with results only partially concordant \(^{[39,40]}\). A recent study that investigated the VCR using functional magnetic resonance imaging (3T blood oxygenation level-dependent-BOLD) and hypercapnic stimulus showed no significant differences before and after loading of levodopa, among patients with Parkinson's disease and healthy controls \(^{[41]}\); However, the study population excluded subjects with cognitive impairment.
5. Objectives of the study

Given the recent findings that identify a possible association between alterations in cerebral hemodynamics and cognitive performance decay of both primary and secondary origin, the objective of this study was to evaluate the possible relationship between cognitive impairment and alterations in cerebrovascular reactivity in patients suffering from Parkinson's disease.

6. Materials and Methods

It was assessed a population of 34 patients, 22 male and 12 female subjects suffering from Parkinson's disease according to UK Brain Bank Criteria pertaining to the Parkinson's Disease Center of the Department of Neurology and Psychiatry of the' University of Rome "Sapienza ". Of each patient were recorded medical history-personal data such as gender, age, age of onset of motor symptoms and dosage of dopaminergic therapy. Were not considered were to be enrolled patients with signs and symptoms of atypical parkinsonism, comorbidity, neurological pathologies can change the cerebrovascular reactivity as previous cerebrovascular events, changes in carotid atheromatous with significant hemodynamic intracranial stenosis, moderate to severe hypertension, diabetes mellitus; were also excluded patients with no or questionable response to dopamine replacement therapy, cognitive impairment (MMSE <24) occurred within one year of onset of motor symptoms, in patients treated with anticholinergic drugs, major depression. The study population was classified into two groups according to the presence of mild cognitive impairment (MCI) according to Diagnostic Criteria for PD-MC (42). All pieces have given their informed consent. Each patient was subjected to:

1. clinical, neuropsychological and motor assessment:

All patients were assessed 60-90 minutes after the administration of L-Dopa and subjected to clinical assessment of motor symptoms by UPDRS part III, H & Y.
Cognitive assessment:
- Mini Mental State Examination (Folstein, 1975) for global cognitive function.
- Ten point clock test.
- Test for the assessment of memory functions: immediate and delayed memory for prose.
- Test for the assessment of executive / attentional: TMT-A and TMT-B, Digit Span
- Assessment of phonological verbal fluency.
- Test for the assessment of visuo-spatial functions: Copy Immediate and Deferred Figure of Rey, test figures tangled.
Praxic-test functions.
In addition, the test was administered structured SCOPA-AUT for evaluation of autonomic functions.

**6.1 MCI in Parkinson's disease**

A diagnosis of MCI is distinct from that of dementia. A person with a diagnosis of MCI has altered cognitive function with objective and subjective indicators of deterioration, without there being necessarily an impact on the complex functions of everyday life.

The existence of a continuum that goes from normal cognitive aging to frank dementia (Alzheimer's disease) and passing through a phase of failure has been postulated in 1995 by Petersen. With a priori hypothesis and applying the same methodological criteria used for Alzheimer's disease, Caviness (43) has incorporated the concept of MCI in Parkinson's disease. A review of 'critique' of this concept was published in 2011 by the Movement Disorders Society (MDS). The condition, which is considered a risk factor for the development of dementia, is found to be rather frequent (27% prevalence) and heterogeneous in terms of number and cognitive domains involved.
However, we must stress that major factors contributing to the overestimation of cognitive impairment in Parkinson's disease: depressive symptoms, pre-morbid personality, motor performance, social isolation, side effects attributable to treatment, and the presence of methodological biases, both within and between groups psychometric analyzes, can limit the validity of the construct PD-MCI \(^{44}\), in particular a pathology where it would seem evident that the cognitive deterioration cannot recognize a single disease process. Therefore, we recognize that the term MCI in PD (PD-MCI) should not be accepted without these appropriate conditions\(^{45}\).

Criteria for the Diagnosis of PD-MCI, from Litvan I (Mov Dis, 2012)

I. **Inclusion criteria**
   - Diagnosis of Parkinson’s disease as based on the UK PD Brain Bank Criteria
   - Gradual decline, in the context of established PD, in cognitive ability reported by either the patient or informant, or observed by the clinician
   - Cognitive deficits on either formal neuropsychological testing or a scale of global cognitive abilities (detailed in section III)
   - Cognitive deficits are not sufficient to interfere significantly with functional independence, although subtle difficulties on complex functional tasks may be present

II. **Exclusion criteria**
   - Diagnosis of PD dementia based on MDS Task Force proposed criteria
   - Other primary explanations for cognitive impairment (e.g., delirium, stroke, major depression, metabolic abnormalities, adverse effects of medication, or head trauma)
• Other PD-associated comorbid conditions (e.g., motor impairment or severe anxiety, depression, excessive daytime sleepiness, or psychosis) that, in the opinion of the clinician, significantly influence cognitive testing

III. Specific guidelines for PD-MCI level I and level II categories

A. Level I (abbreviated assessment)

• Impairment on a scale of global cognitive abilities validated for use in PD or

• Impairment on at least two tests, when a limited battery of neuropsychological tests is performed (i.e., the battery includes less than two tests within each of the five cognitive domains, or less than five cognitive domains are assessed)

B. Level II (comprehensive assessment)

• Neuropsychological testing that includes two tests within each of the five cognitive domains (i.e., attention and working memory, executive, language, memory, and visuo-spatial)

• Impairment on at least two neuropsychological tests, represented by either two impaired tests in one cognitive domain or one impaired test in two different cognitive domains

• Impairment on neuropsychological tests may be demonstrated by:
  • Performance approximately 1 to 2 SDs below appropriate norms or
  • Significant decline demonstrated on serial cognitive testing or
  • Significant decline from estimated premorbid levels
Subtype classification for PD-MCI (optional, requires two tests for each of the five cognitive domains assessed and is strongly suggested for research purposes)

- PD-MCI single-domain—abnormalities on two tests within a single cognitive domain (specify the domain), with other domains unimpaired or
- PD-MCI multiple-domain—abnormalities on at least one test in two or more cognitive domains (specify the domains)

6.2 Ecocolodoppler of the neck vessels, transcranial Doppler ultrasound at baseline and Breath-Holding unit test using a GE LOGIQ P5 pro.

TCD AND WITH CEREBRAL Vasoreactivity BREATH- HOLDING TEST

For the assessment of cerebral vasoreactivity was used a probe of 2 MHz with recording of the flow through the acoustic window of time, the proximal segment (M1) of the middle cerebral artery and the proximal segment (P1) posterior cerebral artery by a experienced operator and not informed about the clinical characteristics of the subject to be examined. The measurements were made in a quiet room, with the subject lying in the supine position, in the absence of visual stimulation or sound. Blood pressure and heart rate were measured at the beginning and at the end of each experiment. The values of mean flow velocity (VMF) at the M1 portion of middle cerebral artery and posterior cerebral artery P1 segment were recorded at baseline and after 4 seconds of breath at the end of an apnea lasting 30 seconds (stimulus to get the hypercapnia). A capnometer (Capnodig; Drager, Lubeck, Germany) recorded the pCO2 during the entire duration of the maneuver to confirm the increase of the gas in the blood.

Breath Holding Index was calculated according to the formula:

\[
\text{BHI} = \left(\frac{\text{MFVBH} - \text{MFVbase}}{\text{MFVbase}}\right) \times 100 \text{ / sec of BH}
\]
Each subject was submitted to the test 3 times at 5 minute intervals to allow the VMF to return to baseline; for each individual value of BHI obtained the first evidence is consistent with that of the other two tests. The BHI used in the analysis is the average of the three tests. With the Doppler ultrasound examination of the neck vessels and transcranial Doppler ultrasound were excluded significant stenosis and has been calculated pulsatility index of Gosling (PI) to assess indirectly the resistances in the microcirculation.

**Magnetic Resonance Imaging of the brain**

The neuroimaging protocol included standard sequences (FLAIR, DP-T2-T1-weighted), using 1.5 T MRI unit (Siemens, Erlangen, Germany). The lesion load was assessed using T2 and Flair and applying the semiquantitative visual scale CHS \(^{46}\).

### 7. Statistical Analysis

For the analysis of the data was the software SPSS 15. It was used the Student t for continuous variables and the chi-square test for non-continuous variables.

### 8. Results

The groups concerned could be homogeneous for age and years of illness; there were no statistically significant differences in the degree of impaired autonomic and motor disabilities, although patients with PD-MCI present average values greater than SCOPA-AUT and UPDRS Part III; according to matching procedures the MMSE was higher in subjects with cognitive impairment. The neuro-cognitive analysis of domains examined (conceptualization, executive functions, memory, attention, visual-spatial functions) showed a prevalent alteration of attention and visual-spatial functions in the population studied. The analysis of neuroimaging has documented a comparison of the degree of suffering microvascular assessed by semiquantitative
visual scale CHS [Table 1] and indirectly with the pulsatility index between the two groups of patients.

The hemodynamic parameters measured by TCD showed no significant differences at baseline of VMF MCA and PCA. After hypercapnic stimulus there was a statistically significant difference in cerebrovascular reactivity, resulting in reduced PD-MCI group (BHI- MCA PD vs PD-MCI: 1.6±0.4 vs 1.1 ±0.4  p <0.01; BHI-PCA PD vs PD-MCI: 1.0 ± 0.5 vs 0.6 ± 0.3 p <0.05) . [Table 2]

<table>
<thead>
<tr>
<th></th>
<th>PD n°=18</th>
<th>PD-MCI n°=16</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Genere (M/F)</td>
<td>(11/7)</td>
<td>(11/5)</td>
<td></td>
</tr>
<tr>
<td>Età media</td>
<td>70.1±6.8</td>
<td>70.5±6.9</td>
<td>n.s.*</td>
</tr>
<tr>
<td>Durata di malattia</td>
<td>7.3±3.6</td>
<td>8.8±6.5</td>
<td>n.s.*</td>
</tr>
<tr>
<td>UPDRS parte III</td>
<td>17.3±5.7</td>
<td>20.8±5.1</td>
<td>n.s.*</td>
</tr>
<tr>
<td>LED (levodopa equivalent dose)</td>
<td>673.8 ±296.1</td>
<td>856.6±617.2</td>
<td>n.s.*</td>
</tr>
<tr>
<td>SCOPA-AUT</td>
<td>13,5±6.9</td>
<td>17,6±7.0</td>
<td>n.s.*</td>
</tr>
<tr>
<td>SCOPA-AUT Item 14-16</td>
<td>1,4±1,5</td>
<td>1,9±1,6</td>
<td>n.s.*</td>
</tr>
<tr>
<td>Carico lesionale CHS</td>
<td>2,3±0,9</td>
<td>2,4±0,9</td>
<td>n.s.*</td>
</tr>
<tr>
<td>Ipertensione arteriosa (SI/NO)</td>
<td>(9/9)</td>
<td>(6/10)</td>
<td>n.s.*</td>
</tr>
<tr>
<td>MMSE</td>
<td>28,1±1,2</td>
<td>23,9±2,5</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

**TABLE 1 Demographic data and clinical**

*n.s.= not significant*
<table>
<thead>
<tr>
<th></th>
<th>PD n°= 18</th>
<th>PD-MCI n°= 16</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>BHI - MCA</td>
<td>1.6±0.4</td>
<td>1.1±0.4</td>
<td>&lt; .01</td>
</tr>
<tr>
<td>BHI - PCA</td>
<td>1.0±0.5</td>
<td>0.6±0.3</td>
<td>&lt;.05</td>
</tr>
<tr>
<td>PI - MCA</td>
<td>1.0±0.1</td>
<td>1.0±0.2</td>
<td>n.s.*</td>
</tr>
<tr>
<td>PI - PCA</td>
<td>0.9±0.1</td>
<td>0.9±0.2</td>
<td>n.s.*</td>
</tr>
<tr>
<td>VMF - MCA</td>
<td>34.0±10.5</td>
<td>33.3±7.8</td>
<td>n.s.*</td>
</tr>
<tr>
<td>VMF - PCA</td>
<td>13.5±6.9</td>
<td>17.6±7.0</td>
<td>n.s.*</td>
</tr>
</tbody>
</table>

**TABLE 2. Rating transcranial**

*n.s.= not significant*

<table>
<thead>
<tr>
<th>t-student; p &lt; 0.01</th>
<th>PD – Ctrl</th>
<th>PD - MCI</th>
</tr>
</thead>
<tbody>
<tr>
<td>BHI - MCA</td>
<td>1.6±0.4</td>
<td>1.1±0.4</td>
</tr>
</tbody>
</table>
t-student; p < 0.05

<table>
<thead>
<tr>
<th></th>
<th>PD - Ctrl</th>
<th>PD – MCI</th>
</tr>
</thead>
<tbody>
<tr>
<td>BHI - PCA</td>
<td>1.0±0.5</td>
<td>0.6±0.3</td>
</tr>
</tbody>
</table>
9. Discussion

The proposed study represents the first evaluation of cerebrovascular reactivity in patients with no-Alzheimer degenerative cognitive impairment; the results are consistent with a reduction of vasomotor reserve, identified by BHI, in patients with Parkinson's disease (PD) and cognitive impairment, when compared to a control group of patients with PD.

Recent acquisitions, with appropriate methodological approaches and samples, suggest that some parameters of cerebral vasoreactivity, after hypercapnic stimulus, is compromised in patients with primary dementia and, possibly, even in patients with MCI.

Therefore, the data for which the vascular dysregulation is a feature frequently observed in neurodegenerative diseases such as Alzheimer's dementia, although difficult to define, emphasizes the potential role of involution brain at the base of the reduced reactivity cerebrovascular.

In fact, the structural and functional integrity of brain tissue depends on the delicate balance between neuronal energy demands tax the activity and substrate availability through the bloodstream, indicating how the neurovascular coupling is a crucial mechanism for maintaining the functional integrity, mutually depends on the elements that constitute it.
Therefore we cannot exclude that the involution of the structures primarily involved in the disease process, in particular by extending the cortex, could lead to inadequate modulation of cerebral vasoreactivity, probably resulting in an impairment of the physiological signaling between the elements of the neurovascular unit.

In line with this hypothesis, some authors have found, through the use of Voxel-Based Morphometry (RM automated method), circumscribed areas of cortical atrophy already evident in patients with MCI and PD (47).

Moreover, as the real impact of Alzheimer-like neuropathological alterations in dementia of the Parkinson's disease is yet to be determined, it could be argued that the data obtained can be attributed to pathophysiological mechanisms similar to those proposed in Alzheimer's disease, recognizing a loss of Aβ-induced vascular wall. However, studies carried out by means of PET-imaging showed a high degree of β-amyloid load in only a small percentage of patients with PDD (48); Therefore, further evidence is needed to establish the real contribution of the pressure taupatic cone on neurovascular dysregulation in patients with PD.
Another pathophysiologic mechanism potentially implicated in the reduction of cerebrovascular reactivity is represented by a deficiency of the cholinergic system; in fact it would seem the neurotransmitter implicated in the neurogenic control of cerebral blood flow. Several lines of evidence using PET studies emphasize the impairment of the cholinergic system in patients with Parkinson's disease and cognitive impairment (49,50) and as such degeneration appears spatially consistent with a reduction in metabolic activity (51).

The impact of vascular disease on cognitive dysfunction in Parkinson's disease has not been widely accepted; the possible contribution of vascular degeneration has been suggested in some studies of ultrastructural evaluation of the cerebral microcirculation in patients with PD, assuming alterations similar to those observed in Alzheimer's disease (52,53). Although the selection of the sample in the present study has been carried out in order to minimize the possible confounding role of cardiovascular risk factors and the groups in question were found to be homogeneous with respect to the parameters established for estimating the
vascular damage (white matter lesion load, index pulsatility), we cannot exclude that the observed data can recognize the contribution of ultrastructural alterations in the microcirculation.

The pathophysiological mechanisms that modulate the relationship between cerebral hemodynamics, cerebrovascular reserve and cognitive deficits remain poorly characterized. However, the emerging model of an altered neurovascular coupling resulting in dysfunction of the microcirculation, regional hypoperfusion and neurodegeneration could integrate the relationship between vascular changes and cognitive decline\(^{(54)}\).

The previously conducted studies on cerebrovascular reactivity in patients with Parkinson's disease have interpreted this aspect as a potential expression of autonomic dysfunction, but does not establish a specific profile or hemodynamic parameter.

In our study, the autonomic aspects were evaluated by SCOPA-Aut, observing, although not reaching statistical significance, mean scores higher in the group of patients with cognitive impairment. In line with other authors, one could hypothesize that alterations in cerebrovascular reactivity may be secondary to a dysregulation in neurogenic vascular control in the context of a systemic autonomic dysfunction. However, it is possible that the relationship between autonomic disorders and cerebrovascular reactivity may represent the expression of a more obvious spread of the disease process, rather than an interdependent phenomenon.

Furthermore, previous studies carried out with different methods, have found no change in the cerebral reactivity in relation to the administration of L-Dopa; however in our population MCI-PD patients tend to assume major doses of LED and, considering the tendency to have a higher value all'UPDRS although did not reach a statistical significance, we cannot exclude that the severity of the disease per se has contributed to the results observed.
Some limitations of the study require cautious interpretation in the generalization of the observed data, in particular:
1) the small sample size
2) as expressed previously, the BHI for the evaluation of cerebrovascular reactivity, while being generally well tolerated and widely validated, presents inherent limitations to the method
3) The integration with more sophisticated neuroimaging methods could provide more sensitive
4) for the definition of MCI in Parkinson's disease is a construct limited.

10. Conclusions
The neurovascular coupling is a module whose function is required for the integrity of the structures that constitute it. The evidence of abnormal cerebrovascular reactivity in patients with Parkinson's disease and cognitive impairment, compared to the control group of PD patients, may indicate how the involution of the primitive neural systems may lead to a negative modulatory effect neurovascular homeostasis. Further evidence is needed to adequately characterize the role of hemodynamic factors in neurodegenerative processes.
REFERENCES


