Extracranial arterial wall volume is increased and shows relationships with vascular MRI measures in idiopathic Parkinson’s disease

Stephen Ball\textsuperscript{a}
Sarah Al-Bachari\textsuperscript{b,d,e}
Laura M. Parkes\textsuperscript{c,e}
Hedley C. A. Emsley\textsuperscript{d,f,g}
Charles N. McCollum\textsuperscript{a}

\textsuperscript{a}Academic Surgery Unit, Institute of Cardiovascular Sciences, University of Manchester
\textsuperscript{b}Department of Neurology, Salford Royal NHS Foundation Trust, Salford, UK
\textsuperscript{c}Division of Informatics, Imaging and Data Sciences, School of Health Sciences, Faculty of Biology, Medicine and Health, University of Manchester, UK
\textsuperscript{d}Faculty of Health and Medicine, Lancaster University, UK
\textsuperscript{e}Division of Neuroscience and Experimental Psychology, School of Biological Sciences, Faculty of Biology, Medicine and Health, University of Manchester, UK
\textsuperscript{f}Department of Neurology, Royal Preston Hospital, Preston, UK
\textsuperscript{g}Faculty of Biology, Medicine and Health, University of Manchester, UK

Correspondence to:
Mr Stephen Ball
University Hospital of South Manchester,
Academic Surgery Unit, Manchester, UK.
Email: stephen.ball@manchester.ac.uk
Tel: +44 (0)161 291 5853
Fax: +44 (0)161 291 5854

Sources of Funding: The sources of funding were the Manchester Surgical Research Trust (Registered Charity No. 702313), the Sydney Driscoll Neuroscience Foundation (Registered Charity No. 1129387), and a local stroke neurology research fund. The funding sources had no involvement in the design, collection or interpretation of the data.

Keywords:
Idiopathic Parkinson’s disease; wall volume; cerebral emboli; white matter lesion volume; arterial arrival time.
Abstract

**Objective**

Idiopathic Parkinson’s disease (IPD) is the second most common neurodegenerative disorder, often complicated by dementia. Cardiovascular risk factors and spontaneous cerebral emboli (SCE) are strongly associated with Alzheimer’s (AD) and vascular dementia (VaD). We measured SCE in the middle cerebral artery and arterial wall volume in the extracranial arteries in patients with IPD and controls, and explored the relationships with structural and physiological MRI brain neurovascular measures.

**Patients and Methods**

Arterial wall volume over 2cm of the axillary and internal carotid arteries (ICA) bilaterally was measured by 3-D tomographic ultrasound in 15 IPD patients and 16 age/gender matched controls. SCE were counted by Transcranial Doppler (TCD) using international consensus criteria. Venous to arterial circulation shunting (v-aCS), usually through a patent foramen ovale (PFO), was measured using a TCD technique with intravenous microbubble contrast. Structural and physiological MRI brain neurovascular measures, acquired separately, comprised white matter lesion volume (WMLV), cerebral blood flow (CBF) and arterial arrival time (AAT).

**Results**

Mean (95% CI) axillary and ICA wall volume was higher in IPD patients at 523mm$^3$ (446, 600) and 455mm$^3$ (374, 536) respectively compared with 412mm$^3$ (342, 483) and 408 mm$^3$ (362, 454) in controls being significant for the axillary artery (p=0.04).

Cerebral WMLV was related to mean arterial wall volume for both axillary ($r=0.555$, $p=0.009$) and ICA ($r=0.559$, $p=0.026$) in all participants.

SCE were detected in four IPD patients and three controls ($p=1.00$). Two IPD patients and three controls were positive for a v-aCS equivalent to PFO ($p=0.477$).

**Conclusion**

Although frequent in AD and VaD, neither SCE nor v-aCS were associated with IPD. This is the first study to demonstrate arterial wall volume is increased in IPD and relates to WMLV.
Introduction

Idiopathic Parkinson’s disease (IPD) is the second most common neurodegenerative disorder worldwide after Alzheimer’s disease (AD). In the UK, IPD has an overall prevalence of 1 in 500, but there is a striking increase with age. Pathological hallmarks include progressive degeneration of dopaminergic (DA) neurones projecting from the substantia nigra (SN) to the striatum and intracytoplasmic inclusions called Lewy bodies comprising aggregated forms of the protein α-synuclein within surviving SN DA neurones. (1) Clinical diagnosis rests on the cardinal motor features of bradykinesia, rest tremor, rigidity, and postural instability, although IPD causes a number of non-motor features including disturbances of sleep, mood and cognition. Neurodegenerative disorders are increasingly believed to be multifactorial with many factors leading to neuronal death. (2) Accumulating evidence suggests that the neurodegenerative process is also influenced by neurovascular changes which constitute more than simply the presence of comorbid cerebrovascular disease (CVD) as a consequence of ageing. Blood-brain barrier (BBB) damage is implicated in the pathogenesis of IPD, with normal BBB structure and function being dependent on the integrity of the neurovascular unit (NVU), which comprises pericytes, glial cells, neurones and basal lamina, and tightly regulates the blood-brain interface. For example, string vessels comprising collapsed basement membrane lacking endothelium, with no function in circulation, are increased in IPD, suggesting a possible role for cerebral hypoperfusion in the neuronal degeneration of IPD, which needs further investigation. (3)

Studies on the prevalence of CVD in IPD have generated conflicting results with reports of increased, (4-8) decreased (9-11) or unchanged prevalence (12-16), by comparison with controls. Recently, MRI measures of neurovascular status have found prolonged arterial arrival time, thought to be related to age driven structural cerebrovascular changes, in patients with IPD compared to age and cardiovascular risk matched controls. (17)

We have previously demonstrated that cerebral emboli have a role in the pathophysiology of AD and vascular dementia (VaD), two other common neurodegenerative disorders. (18-23) Cerebral emboli showed similar frequency, and similar association with cognitive decline, in both AD and VaD, suggesting some commonalities in pathophysiology. This could be mediated via disturbance of the cerebral microcirculation as a result of cerebral microemboli, potentially associated with microglial activation and neuroinflammation. In this study we have investigated cerebral emboli and extracranial arterial wall volume in patients with IPD,
and explored the relationships with structural and physiological MRI brain neurovascular measures.

**Patients and Methods**

Participants enrolled in an MRI study investigating structural and physiological neurovascular measures were invited to participate. These included both patients with IPD and age and gender matched healthy controls. Eligibility criteria for IPD participants were a clinical diagnosis of IPD fulfilling UK Parkinson’s disease society brain bank criteria.

Exclusion criteria included: Clinical or radiological features suggesting secondary or atypical parkinsonism; Features suggesting vascular parkinsonism; History of TIA or stroke; Other Focal neurological signs; Cognitive dysfunction; Evidence of infection within the previous 6 weeks or the presence of a concomitant inflammatory condition. Local ethical approval was granted and patients underwent written informed consent.

3D tomographic ultrasound (3D t-US) was used to measure arterial wall volume, as an accurate measure for intimal thickness, bilaterally over 2cms of the axillary and internal carotid arteries (ICA). A magnetically tracked freehand t-us system (Curefab GmbH, Munich, Germany) was attached to a Philips iu22 duplex ultrasound (Philips, Bothwell, USA) to track the transducer orientation and position in time and space. Multi-planar reconstructions (MPR) were computed with an ultrasound volume from the 2D ultrasound frames almost instantly.

Firstly, a plain 2D scan was performed in B-Mode to gain a signal and grade stenosis using a combination of grey-scale and velocity. A 2cm segment of each artery was then scanned in the transverse plane and the images captured on the 3D software. The proximal 2cms of the ICA and 2cms of the axillary artery which produced the clearest images were chosen for analysis. Volume was calculated by two independent observers using manual planimetry. The outer and inner vessel walls were identified and circumscribed for each transverse slice with an inter-slice distance of 1mm. The volume was then calculated automatically by summing the areas in each slice and multiplying it by the inter-slice distance.

By insonating the middle cerebral artery (MCA) through the transtemporal window, cerebral emboli can be detected and their frequency counted. All patients underwent one hour of continuous transcranial Doppler (TCD) insonation of the middle cerebral artery using a 2-MHz pulsed-wave Doppler probe. Patients were observed for any movement or potential artefacts and the data analysed by two observers, blinded to each other’s results. The
international consensus criteria (24) for emboli detection were used, which specify that embolic signals should be transient (lasting <300 millisec), at least 3 dB higher than the background blood flow, unidirectional, within the Doppler spectrum, and accompanied by an audible ‘snap’, ‘chirp’, or ‘moan’.

The presence of a v-aCS was investigated using an emulsion of air microbubbles in saline as an ultrasound contrast medium following the completion of one hour TCD. The bubble suspension was rapidly injected intravenously under three conditions, each separated by one minute: (i) resting quietly; (ii) coughing repeatedly during injection and for a further 10 sec; (iii) performing a standardised Valsalva manoeuvre with 5 sec release after injection. The presence of a v-aCS equivalent to a Patent Foramen Ovale (PFO) was defined as 15 or more embolic signals with the first within 12 cardiac cycles of contrast administration.

Magnetic resonance imaging was undertaken on a 3T Philips Achieva MRI Scanner. The full MRI methodology has been reported previously.(17) In brief, in this study we used images from: 1) T2-weighted FLAIR (repetition time TR 11s, inversion time TI 2.8s, echo time TE 120 ms) from which an estimation of white matter lesion volume (WMLV) (a widely used marker of cerebral small vessel disease) was determined using the lesion segmentation toolbox in SPM8. 2) Arterial spin labelling (ASL), using pulsed labelling and multiphase readout at 4 readout times of 800, 1400, 2000, 2600 ms, TR: 3500 ms; TE 22ms; flip angle 40 degrees with 60 control and label pairs. Voxel-wise fitting to a single blood compartment model enabled quantification of cerebral blood flow (CBF) and arterial arrival time (AAT), the time taken for blood to travel from the labelling slab to the tissue of interest. Whole brain values for CBF and AAT were calculated using a simple threshold mask based on the ASL control images on an individual basis.

Analyses were performed using SPSS version 22. A Generalised Estimating Equation longitudinal regression analysis was used to compare carotid, axillary and temporal artery values between IPD patients and controls. Sets of readings for both the left and right side were obtained from each of 2 observers.

A two-sided 5% significance level was used throughout the analysis. To assess the relationship between total wall volume and the main outcome variables, namely WMLV, AAT and CBF, Pearson correlation, using Log transformed variables, was used and represented graphically. Linear regression was used when adjusting for variables. Smoking was used as a proxy for gender as no females smoked. Comparisons of the outcome variables
between patients with IPD and controls were made using independent t-tests and Mann-
Whitney U tests.

Reliability of the two observer values was assessed using a combination of correlational and
bland-Altman analysis.

**Results**

Fifteen patients with IPD (mean (sd) age 67 (8)) and 16 age and gender matched healthy
controls (mean (sd) age 66 (7)) were enrolled. Cardiovascular risk factors did not differ
significantly between the two groups (Table 1). Table 2 summarises the outcome variables in
the two groups.

**Arterial Wall Volume**

As there was good agreement between the two observers, demonstrated by a correlation of
>0.8 for all but the left ICA (0.74), a mean of the two values was taken for each artery and a
mean of left and right for each artery.

Mean (95% CI) wall volume of the axillary artery was significantly higher in IPD patients at
522.8mm$^3$ (446.0, 599.7) compared to controls at 412.5mm$^3$ (342.2, 482.7) with a difference
of 110.4mm$^3$ (6.3, 214.5) ($p = 0.04$). Mean wall volume of the ICA was also higher in IPD
patients at 455.0mm$^3$ (373.9, 536.2) compared to controls at 407.7mm$^3$ (361.58, 453.8) but
this difference of 47.4mm$^3$ (-46.0, 140.8), did not reach statistical significance ($p = 0.32$).

**White Matter Lesion Volume**

Combined analysis of all 31 participants revealed a significant moderate positive correlation
between WMLV and mean arterial wall volume for both the ICA ($r=0.436$, $p = 0.026$) and
axillary arteries ($r=0.504$, $p=0.009$). (Figure 1) After adjustment for age and smoking the
positive correlation remained significant for the axillary artery ($p=0.009$) but not for the ICA
($p=0.064$).

When analysing the subgroups, the correlation between WMLV and mean arterial wall
volumes in the IPD group remained positive but weak at 0.142 for the axilla and 0.292 for the
ICA. (Figure 2) Neither reached statistical significance before and after adjustment for age
and smoking. In the control group, significant positive correlations were reflected for both the
ICA ($r=0.651$, $p=0.012$) and axillary artery ($r=0.722$, $p=0.004$). This significance remained
for the ICA group (p=0.015) but was lost for the axillary artery (p=0.094) when adjusting for age and smoking.

**Whole brain Arterial Arrival Time and Cerebral Blood Flow**

Combined analysis of all 31 participants revealed a significant moderate positive correlation between whole brain AAT and mean wall volume of the axillary artery (r=0.403, p=0.041) and a weak non-significant correlation with the ICA (r=0.236, p=0.245). (Figure 3) Following adjustment for age and smoking, the significance with the axillary artery wall volume was lost (p=0.085).

Subgroup analysis revealed weak and moderate correlations between whole brain AAT and mean arterial wall volume of both the ICA (r=0.238, p=0.457) and axillary artery (r=0.517, p=0.085) in the IPD group. (Figure 4) When adjusted for age and smoking the relationship between whole brain AAT and axillary artery wall volume became significant (p=0.022). In the control group the correlation with whole brain AAT was weak and non-significant for both the ICA (r=0.173, p=0.555) and axillary artery (r=0.234, p=0.420).

No significant association was observed between mean axillary or ICA wall volume and CBF.

**Cerebral Emboli**

TCD monitoring for SCE in the MCA was performed in 12 IPD patients and 12 controls with SCE detected in only 4 (33%) IPD patients and 3 (25%) healthy controls (p=1.00). Two (17%) IPD patients and three (25%) controls were positive for a v-aCS equivalent to PFO (p=0.477); similar to the prevalence of PFO in the adult population.

**Discussion**

To our knowledge, this is the first study to investigate extracranial arterial wall volume in idiopathic Parkinson’s disease and any potential association with structural and physiological MRI brain neurovascular measures. In addition, we investigated the presence of cerebral emboli and evidence of v-aCS in IPD, by comparison with controls.

We have found significantly higher axillary artery wall volume in patients with IPD than in healthy controls. Previously, Rektor et al. reported significantly increased intima-media thickness (IMT) in the common carotid artery (CCA) of patients with IPD compared to controls, concluding that this might suggest a link between generalised atherosclerotic
disease, of which increased CCA IMT is a known marker, and IPD. (25) A positive
correlation was observed between axillary artery wall volume and WMLV in the combined
analysis of all participants, and in the analysis of the control group but not in the IPD group,
suggesting that there may be differences in underlying pathophysiological factors in healthy
ageing and in IPD. Following adjustment for age and smoking, a significant positive
correlation was observed between AAT and axillary artery wall volume in the IPD group.
This relationship was not seen in the control group. Again, this seems to imply a divergence
in underlying pathophysiology.

The question of whether there is any causal relationship between atherosclerotic disease, or
between conventional cerebrovascular disease (CVD), and IPD, is unclear. It remains
possible that CVD represents a co-morbidity alongside a predominantly age-related
neurodegenerative disorder. However, the present observations of increased axillary artery
wall volume, and the suggestion of a relationship between axillary artery wall volume and
AAT does merit further examination.

Recent findings of alterations in physiological neurovascular measures in IPD – including our
own observations of prolonged cerebral AAT, are currently attributed to structural vascular
changes such as increased vessel tortuosity, increased rarefaction and arteriolar wall damage,
so any potential new insights into novel markers of extracranial arterial changes in IPD, as in
the present study, are valuable. (17) Previously we have reported a higher prevalence of
radiological and clinical cerebrovascular disease in patients with IPD compared to
controls. (26) Emerging data indicate that over 60% of patients with recent onset IPD have
high or medium vascular risk, which is associated with a worse motor and cognitive
phenotype. Statins are underused, representing a missed opportunity for vascular secondary
prevention. (27) Improved recognition of the burden of systemic vascular changes in IPD,
how this may interact with the neurodegenerative process and treatment response, as well as
provide opportunities for treatment, is needed.

There is an abundance of literature concerning the aetiology of white matter lesions and their
involvement in ageing and in neurological disease, especially dementia. (28-35) It has to be
remembered that the pathophysiology of WMLs is unclear and could be multifactorial. The
present study found a significant correlation between axillary artery wall volume and WMLV
which persisted following adjustment for age and smoking. Our own recent data suggest
differences in WML burden between IPD motor phenotypes, with an apparently greater
burden in patients with the more severe postural instability and gait disorder pattern, and with
greater cognitive impairment.\(^{36}\) The lack of association of ICA wall volume and WMLs in
the current study is perhaps surprising given the axillary artery findings, but nonetheless, a
lack of association has been reported before.\(^{37}\) This is likely consistent with the fact that
most WMLs are probably not due to atherothromboemboli, and the current finding of axillary
artery changes is more indicative of more widespread changes in the vasculature with ageing,
including in IPD.

With respect to the TCD findings in the present study, the results do contrast with our
findings in AD and VaD, where an association was observed with spontaneous and
paradoxical cerebral emboli. Again, this would argue against a significant role for cerebral
emboli as part of the neurovascular process in IPD, although it is acknowledged that our
negative finding may be influenced by our small sample size.

This present study was relatively small, which did not afford an opportunity to explore the
relationships with different IPD phenotypes, including motor phenotypes and non-motor
symptoms such as cognitive impairment and dementia. Large, longitudinal studies of clinical
and imaging measures of the extracranial and intracranial vasculature, including the
microvasculature, will be required in order to advance our knowledge in this area. However,
our results provide some indication that extracranial arterial wall volume may correlate with
intracranial small vessel disease, including in patients with IPD.

**Conflicts of interest:** None

**Acknowledgements:** None
References:


Table 1: Cardiovascular risk factors between the two groups

<table>
<thead>
<tr>
<th>CV Risk Factor</th>
<th>IPD</th>
<th>Control</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hypertension</td>
<td>6</td>
<td>6</td>
<td>0.462</td>
</tr>
<tr>
<td>Diabetes</td>
<td>1</td>
<td>2</td>
<td>0.619</td>
</tr>
<tr>
<td>Hypercholesterolaemia</td>
<td>3</td>
<td>9</td>
<td>0.098</td>
</tr>
<tr>
<td>Ischaemic Heart Disease</td>
<td>2</td>
<td>2</td>
<td>0.660</td>
</tr>
</tbody>
</table>

Table demonstrating no difference between the two groups with regards cardiovascular risk factors.
Table 2: Comparison of outcome variables between the two groups

<table>
<thead>
<tr>
<th></th>
<th>IPD</th>
<th>Control</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean ICA (sd) Wall</td>
<td>455 (97.1)</td>
<td>407.4 (166)</td>
<td>0.32</td>
</tr>
<tr>
<td>Volume</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (sd) Axillary Wall</td>
<td>522.8 (157.1)</td>
<td>412.5 (148)</td>
<td>0.04</td>
</tr>
<tr>
<td>Volume</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mean (sd) WMLV</td>
<td>5.29 (6.2)</td>
<td>6.49 (11.3)</td>
<td>0.410</td>
</tr>
<tr>
<td>Mean (sd) AAT</td>
<td>1491 (137.9)</td>
<td>1421.6 (152.6)</td>
<td>0.227</td>
</tr>
<tr>
<td>Mean (sd) CBF</td>
<td>45.8 (8.5)</td>
<td>46.2 (5.3)</td>
<td>0.797</td>
</tr>
<tr>
<td>+ve Emboli (n=12)</td>
<td>4</td>
<td>3</td>
<td>//</td>
</tr>
<tr>
<td>+ve V-acs (n=12)</td>
<td>2</td>
<td>3</td>
<td>//</td>
</tr>
</tbody>
</table>

Table demonstrating that the mean wall volume of both the internal carotid artery (ICA) and axillary artery is increased in patients with idiopathic parkinsons disease (IPD) and significantly so for axillary wall volume.
Figure 1: Scatter plots demonstrating the relationship between mean arterial wall volume (mm$^3$) and white matter lesion volume (WMLV) and arterial arrival time (AAT). There is a significant moderate correlation between WMLV and both mean internal carotid artery (ICA) wall volume (A) and mean axillary wall volume (B). There was no association between AAT and ICA mean wall volume (C) but a moderate significant association between with mean axillary wall volume (D).
Figure 2: Scattergrams demonstrating the relationship between mean artery wall volume and LogWMLV and LogAAT in IPD patients

Figure 2: Scatter plots demonstrating a lack of relationship between mean arterial wall volumes of both the internal carotid artery (ICA) and axillary artery with white matter lesion volume (WMLV) (A+B) and arterial arrival time (AAT) (C+D) in patients with idiopathic parkinsons disease (IPD).