Regional brain iron and gene expression provide insights into neurodegeneration in Parkinson’s disease

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The mechanisms responsible for the selective vulnerability of specific neuronal populations in Parkinson’s disease are poorly understood. Oxidative stress secondary to brain iron accumulation is one postulated mechanism. We measured iron deposition in 180 cortical regions of 96 patients with Parkinson’s disease and 35 control subjects using quantitative susceptibility mapping. We estimated the expression of 15 745 genes in the same regions using transcriptomic data from the Allen Human Brain Atlas. Using partial least squares regression, we then identified the profile of gene transcription in the healthy brain that underlies increased cortical iron in patients with Parkinson’s disease relative to controls. Applying gene ontological tools, we investigated the biological processes and cell types associated with this transcriptomic profile and identified the sets of genes with spatial expression profiles in control brains that correlated significantly with the spatial pattern of cortical iron deposition in Parkinson’s disease. Gene ontological analyses revealed that these genes were enriched for biological processes relating to heavy metal detoxification, synaptic function and nervous system development and were predominantly expressed in astrocytes and glutamatergic neurons. Furthermore, we demonstrated that the genes differentially expressed in Parkinson’s disease are associated with the pattern of cortical expression identified in this study.

Our findings provide mechanistic insights into regional selective vulnerabilities in Parkinson’s disease, particularly the processes involving iron accumulation.

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Keywords: Parkinson’s disease; iron; quantitative susceptibility mapping; genetics; transcriptomics

Abbreviation: QSM = quantitative susceptibility mapping

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Introduction

In Parkinson’s disease, bradykinesia and rigidity severity correlate approximately with the degree of nigrostriatal dopamine denervation \(^1\) and the presence of \(\alpha\)-synuclein-rich cell inclusions, \(^2\) Lewy bodies, which represent the pathological signature associated with the disease. However, \(\alpha\)-synuclein inclusions do not correlate with symptom severity \(^3\) and the pathophysiological processes that are responsible for the associated dementia seen in many elderly patients remain unknown. As such, there is an increasing need to understand why some brain regions display selective vulnerability to neurodegeneration \(^4\) and determine the mechanisms that trigger the degenerative cascade. \(^5\)

For over 25 years, it has been known that iron accumulates in the basal ganglia and substantia nigra in Parkinson’s disease \(^6-8\) with the resultant oxidative stress being of interest as an important potential driver of neurodegeneration. \(^9\)-\(^11\) Moreover, this observation has been corroborated by recent advances in the study of this disease. \(^12,13\) Iron accumulates in the brain, especially within the basal ganglia, during ageing, \(^14\) partly due to the increased permeability of the blood–brain barrier. \(^15\) High levels of iron in the tissue cause a build-up of toxic reactive oxygen species that interfere with mitochondrial function, \(^16\) damage DNA, \(^17\) catalyse dopamine oxidation reactions to produce toxic quinones \(^18\) and irreversibly modify proteins through highly reactive aldehydes. \(^19\) All these causes of cell stress ultimately lead to iron-mediated cell death. \(^20\) Intriguingly, lipofuscin formation, previously characterized in Parkinson’s disease, \(^21\) has recently been shown to be driven by the disruption of lysosomal lipid metabolism in neurons. \(^22\) This in turn leads to iron accumulation, oxidative stress and ferroptosis, \(^22\) providing a direct link between the pathological inclusions in Parkinson’s disease and excess brain iron. Excess iron also interacts directly and indirectly through free radical species with key pathological proteins associated with Parkinson’s disease by promoting the aggregation of \(\alpha\)-synuclein, \(^23\) stimulating the production of amyloid-\(\beta\) via the downregulation of furin, \(^24\) and increasing the toxicity of amyloid-\(\beta\) either directly \(^25\) or through increased tau phosphorylation. \(^26\)

Until recently, whole brain neuroimaging using conventional structural MRI has maintained only a limited role in uncovering neurodegenerative mechanisms in Parkinson’s disease as cortical atrophy is not prominent, particularly early in the disease process. \(^27\) However, quantitative susceptibility mapping (QSM) \(^28\) has emerged as a powerful new technique, which is sensitive to local sources of magnetic susceptibility and in particular to variation in brain iron content. \(^29,30\) We recently used QSM to show that brain iron is associated with disease severity in Parkinson’s disease, \(^31\) with higher levels of hippocampal iron linked to poorer cognition and higher putaminal iron linked with worse motor performance. Whether brain iron deposition is a cause of neuronal damage or a surrogate marker remains unknown, but QSM offers a new way to explore the regional effects of neurodegeneration.

We have used the spatial information about iron accumulation provided by QSM and combined it with regional gene expression profiles from the Allen Human Brain Atlas transcriptomic data \(^32\) to identify possible mechanisms for regional selective vulnerability in Parkinson’s disease. The approach of combining neuroimaging with transcriptomic data has previously been used to investigate autism, \(^33\) schizophrenia \(^34\) and Huntington’s disease. \(^35\) Here, we used partial least squares (PLS) regression to test whether cortical brain iron accumulation in Parkinson’s disease measured using QSM correlates with specific patterns of gene expression to shed light on the gene expression profiles that render specific cortical regions vulnerable to higher levels of brain iron accumulation and subsequent neurodegeneration. We also performed an enrichment analysis on the identified gene expression patterns linked to higher brain iron to determine the biological and cell processes linked with the degenerative process of aberrant iron accumulation and neurodegeneration.

Materials and methods

Participants

Ninety-six patients with Parkinson’s disease with a disease duration of <10 years [age 49–80 years, mean = 66.4, standard deviation (SD) = 7.7, 48 females] volunteered to participate from October 2017 to December 2018. The inclusion criteria were a clinical diagnosis of early to mid-stage Parkinson’s disease (Queen Square Brain Bank Criteria \(^36\) in the age range of 49–80 years. Exclusion criteria were confounding neurological or psychiatric disorders, dementia and metallic implants considered unsafe for MRI. Participants continued their usual therapy (including L-DOPA) for all assessments. No patients were taking cholinesterase inhibitors. In addition, we recruited a group of 35 age-matched controls (50–80 years, mean = 66.1, SD = 9.4, 21 females) that included some unaffected patient spouses. All participants gave written informed consent, and the study was approved by the Queen Square Research Ethics Committee. All participants underwent clinical assessments of motor function, cognition, vision, mood and sleep, as previously described. \(^31\) Motor assessments were performed using the Movement Disorder Society–Unified Parkinson’s Disease Rating Scale, \(^37\) general cognition was assessed using the Mini-Mental State Examination \(^38\) and the Montreal Cognitive Assessment \(^39\) (see Table 1 for clinical and demographic information).

MRI acquisition

MRI acquisition was as previously described. \(^31\) In brief, MRI measurements consisting of susceptibility- and T\(_1\)-weighted MRI scans were performed on a Siemens Prisma-fit 3-T MRI system using a 64-channel receive array coil (Siemens Healthcare). Susceptibility-weighted MRI signals were obtained from a 2 × 1-accelerated, 3-D flow-compensated spoiled-gradient-recalled echo sequence with flip angle, 12\(^\circ\), echo time, 18 ms; repetition time, 25 ms; receiver bandwidth, 110 Hz/pixel; matrix dimensions 204 × 224 × 160 with 1 × 1 × 1 mm voxel resolution (scan time, 5 min 41 s). T\(_1\)-weighted magnetization-prepared 3-D
rapid gradient-echo (MPRAGE) anatomical images were acquired using a 2 × 1-accelerated sequence with inversion time, 1100 ms; flip angle, 7°; first echo time, 3.34 ms; echo spacing, 7.4 ms; repetition time, 2530 ms; receiver bandwidth, 200 Hz/pixel; matrix dimensions 256 × 256 × 176 with 1 × 1 × 1 mm voxel size (scan time, 6 min 3 s).

### QSM preprocessing and spatial standardization

QSM preprocessing was as previously described. Brain masks were calculated using the BET2 algorithm in the FMRIB software library (FSL version 5.0, https://fsl.fmrib.ox.ac.uk/fsl/fslwiki; accessed Jan 2019). Background field removal was completed using Laplacian boundary value extraction and variable spherical mean-value filtering, and susceptibility maps were estimated using Multi-Scale Dipole Inversion in MATLAB. A linear model was fitted to each region of interest in all the control subjects such that:

\[ \hat{Y}_i = \alpha_i + \beta_i A \]

where \( \hat{Y}_i \) is the estimated mean susceptibility value in region \( i \), \( A \) is the subject’s age and \( \alpha_i \) and \( \beta_i \) are the fitted model parameters. The QSM susceptibility values in the controls and patients with Parkinson’s disease were then age-adjusted according to:

\[ \chi_{ik} = Y_{ij} + \beta_i(\mu_i - A_i) \]

where, for region \( i \) in subject \( j \), \( Y_{ij} \) is the unadjusted susceptibility value, \( \mu_i \) is the mean age of the control subjects, \( A_i \) is subject’s age and \( \chi_{ij} \) is the age-adjusted susceptibility value. We used the fitted parameters from the control group to age-correct the susceptibility values measured in the patients with Parkinson’s disease to avoid removing effects of disease progression with age. For the sex adjustment, as sex may affect magnetic susceptibility values differently in controls and Parkinson’s disease patients, a linear model was fitted to each region of interest for each participant group separately:

\[ \hat{Z}_{ik} = \chi_{ik} + \beta_{ik}S_k \]

where, for region \( i \) in participant group \( k \), \( \chi_{ik} \) is the estimated age-adjusted susceptibility, \( S_k \) is the sex of the subject and \( \alpha_{ik} \) and \( \beta_{ik} \) are the fitted parameters. We then adjusted for the effect of sex in each group using:

\[ \chi_{ijk} = \chi_{ik} + \beta_{ik}(\mu_k - S_k) \]

where, for region \( i \) in subject \( j \) in participant group \( k \), \( \chi_{ijk} \) is the age-adjusted susceptibility.
To generate a continuous measure of magnetic susceptibility difference in patients with Parkinson’s disease relative to the controls, Parkinson’s age- and sex-adjusted regional means were normalized to the control mean for that region by a z-score transformation using MATLAB, giving a \( 180 \times 1 \) age-adjusted QSM score vector, \( \mathbf{Y} \) (see Fig. 1 for an overview of QSM and gene expression processing). Statistically significant differences between the regions of interest were probed using \( t \)-tests and are reported at \( P < 0.05 \) and \( Q < 0.05 \).

**Estimation of regional gene expression**

We used the Allen Human Brain Atlas microarray transcriptomic data for five male and one female donors with no history of neurological or psychiatric disease (mean age 42.5 years), available from the Allen Institute for Brain Science. As data were available from six donors for the left hemisphere but only two donors for the right hemisphere, we assessed the left hemisphere samples only, as these were considered more robust. Using MATLAB, each tissue sample was assigned to one of the 180 left cortical regions of the Glasser atlas, using the Allen Human Brain Atlas MRI data for each donor, in a process that has previously been described in detail. Genes with expression levels above a background threshold of 50% were selected and gene expression data were normalized across the left cortex, as previously described. Regional expression levels for each gene were compiled to form a \( 180 \times 15,745 \) regional transcription matrix, \( \mathbf{X} \).

**Partial least squares regression**

We used PLS regression to examine the association between the healthy brain transcriptome and cortical QSM in Parkinson’s disease, as this technique is well suited to the high collinearity of gene expression data. PLS regression is a multivariate analysis technique, similar to principal component analysis, which combines dimension reduction and linear regression, producing components from \( \mathbf{X} \) (the \( 180 \times 15,475 \) predictor matrix of 180 regional mRNA measurements for 15,475 genes) that have maximum covariance with \( \mathbf{Y} \) (the \( 180 \times 1 \) regional QSM score vector). The second PLS component (PLS2) was used to weigh and rank gene predictor variables. In total, 10,000 permutations based on sphere-projection-rotations of the QSM-score cortical map were examined to test the null hypothesis that PLS2 explained no more variance in \( \mathbf{Y} \) than chance. Bootstrapping was used to estimate the variability of each gene’s positive or negative weight on PLS2 and the ratio of the weight of each gene to its bootstrapped standard error was used to rank its contribution to the PLS2. All PLS and bootstrapping analyses were conducted in MATLAB. We tested the null hypothesis of zero weight for each gene using a false discovery rate inverse quantile transformation correction to account for winner’s curse bias using R version 3.6.1. Only genes that survived this correction at \( Q < 0.05 \) were included in the enrichment analyses, and upweighted and downweighted genes were assessed separately.

**Gene ontological analysis**

We used the g:Profiler toolset, implemented in R, to perform a gene ontological (GO) enrichment analysis of the significant

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**Figure 1** An overview of the methodology used for regional QSM extraction, estimation of regional gene expression, PLS regression, gene ontological and cell type analyses. (A) Mean, signed QSM values were extracted from 180 left-cortical regions for 96 patients with Parkinson’s disease and 35 controls. (B) A QSM score, \( Y_{PD} \), was calculated for each region by a z-score transformation. (C) Allen Human Brain Atlas samples of gene expression data were mapped to the 180 left-cortical regions according to the anatomical parcellation and were used to create a matrix containing the average expression of 15,745 genes in those regions. (E) A bootstrapped PLS regression was performed using gene expression (\( \mathbf{X} \)) as the predictor variable and the QSM score (\( \mathbf{Y} \)) as the response variable. The second component of \( \mathbf{X} \) explained maximum variance in \( \mathbf{Y} \) and bootstrapped z-scores were used to rank each gene’s contribution to this component. (F) Genes that were significant at \( Q \) (corrected \( P \)) < 0.05 underwent gene ontological analyses for biological processes and expression-weighted cell-type enrichment analyses.
positively and negatively weighted genes defined by PLS2. We filtered the resulting list of GO terms by retaining only those that were significantly enriched at \( P < 0.05 \) (corrected for multiple comparisons using the gSCS algorithm) and discarded the terms associated with >2500 genes as being too general. To reduce and visualize the GO terms, we used the REViGO web page tool, which is based on semantic similarity. We performed additional GO enrichment analyses using R to mitigate against the possibility of false-positive bias for GO terms in the enrichment analyses of brain-wide transcriptomic data due to null models not accounting for gene-gene co-expression and spatial autocorrelation present within such data. Specifically, we ran GO enrichment analyses for both a random and spatial-spin permutation of our QSM score data (generated in MATLAB) and compared the associated GO terms to those arising from our main analysis.

### Cell-type analysis

As gene expression is often driven by the underlying cell-type distribution, we used expression-weighted cell-type enrichment analysis, implemented in R, to investigate whether the most strongly weighted genes were more significantly expressed in particular cell types. We took the top 20\% of the most significantly upweighted and downweighted genes identified by PLS2 and assessed their relative expression in the cell types defined in the Allen Institute for Brain Science single-cell transcription dataset against a background set of all 15745 genes included in our initial analysis. We replicated our analysis using a separate human derived dataset and performed replication analyses with the top 10, 30, 40 and 50\% of PLS2 genes to ensure that the results were not driven by the threshold selection.

### Comparison with external Parkinson’s disease post-mortem-derived gene expression data

To test whether differentially expressed genes in Parkinson’s disease and associated disease states help to explain the pattern of PLS2 expression, we matched the sets of genes identified in cortical and subcortical brain regions in Parkinson’s disease, as well as cortical regions in Parkinson’s disease dementia, and dementia with Lewy bodies, to our weighted PLS2 gene list. This included the reanalysis of one dataset using the available clinical data to differentiate between Parkinson’s disease and Parkinson’s disease dementia, with the RNA-sequencing data in each group analysed using the R package DESeq2 version 1.30.0. We used age at death and the RNA integrity number as covariates and generated lists of genes that were differentially expressed in: Parkinson’s disease relative to controls; Parkinson’s disease dementia relative to controls; and Parkinson’s disease dementia relative to Parkinson’s disease (significant at \( P < 0.05 \)). We tested whether genes in each dataset were significantly more positively or negatively weighted than was due to chance by using 10 000 random permutations of the same sample size. To maintain consistency across datasets, we only included genes with an absolute fold-change of > 1.5. The significant results are reported at Bonferroni corrected \( P < 0.0023 \) (22 comparisons across five studies).

### Data availability


### Results

#### Brain iron content is increased in Parkinson’s disease compared with control subjects

The regional cortical signed QSM scores showed increases in magnetic susceptibility in patients with Parkinson’s disease relative to controls in the frontal, posterior parietal and insular cortices, and slightly decreased susceptibility values in patients with Parkinson’s disease relative to controls in the occipital cortex (Figs 2A and 3B), indicating differences in brain iron content in these brain regions. These differences were statistically significant (\( P < 0.05 \)) in 17 Glasser regions of interest, with increased magnetic susceptibility in Parkinson’s disease in 16 regions of interest and increased magnetic susceptibility in controls compared with Parkinson’s disease in one region of interest (Fig. 2B and Table 2). The anterior agranular insular complex survived correction for multiple comparisons (FDR-adjusted \( Q < 0.05 \)). These results were qualitatively similar to the whole-brain results we obtained using absolute QSM in our previous study, where we found increased absolute QSM susceptibility values in Parkinson’s disease in the frontal and posterior parietal cortices.

#### Relating cortical brain iron in Parkinson’s disease to variation in gene expression patterns

We used PLS regression to identify the pattern of gene expression that correlated with the anatomical distribution of brain iron content, as measured using QSM. The PLS2 explained the most (20\%) variance in the QSM score (\( P < 0.01 \)) and
the PLS2 gene expression weights showed a strong positive correlation with the QSM score \( r = 0.44, P = 4.46 \times 10^{-10} \) (Fig. 3), meaning that genes that were positively weighted on PLS2 were also more highly expressed in cortical brain regions with higher magnetic susceptibilities. Similarly, genes that were negatively weighted on PLS2 showed relatively low expression in cortical brain regions with high QSM scores. The spatial profile of PLS2 weightings matched that of the QSM scores, particularly in the frontal, posterior cingulate and insular cortices. We therefore used PLS2 to rank and select significantly weighted genes, giving a set of 1622 significantly upweighted and 1068 significantly downweighted genes \( Q < 0.05 \). We assessed these sets separately (Fig. 3). The complete set of PLS2 gene weights and associated statistics are provided in Supplementary Table 1.

Using GO analyses, we found sets of biological processes associated with upweighted genes (Fig. 4A). Genes more highly expressed in regions with higher QSM values in Parkinson’s disease were enriched for GO terms relating to nervous system development, synaptic transmission and signalling and the detoxification of and stress response to metal ions (Table 3). We also found these genes to be enriched for the REACTOME pathways ‘Metallothioneins bind metals’ and ‘Response to metal ions’. In contrast, we did not find any GO biological processes to be enriched in the downweighted gene set. Full tables of the GO terms associated with up- and downweighted genes are provided in Supplementary Tables 2 and 3.

Our additional control analyses confirmed that these findings were specific to the QSM data, rather than spuriously arising from gene-gene co-expression. Specifically, the control analysis using a spatial spin of the cortical data revealed upweighted genes that were not enriched for any GO biological processes, and downweighted genes enriched for biological processes relating to immune cell proliferation and viral transcription. See Supplementary Tables 4 and 5 for the full list of the GO terms associated with the up- and downweighted genes from the control spatial-spin null analyses. The second control analysis was performed using a random-null model and generated an upweighted gene list that was not significantly enriched for any GO biological processes and a downweighted list that was enriched only for the term ‘response to lipopolysaccharide’. These control analyses provide additional support that the gene lists arising from our experimental analysis relating spatial variation in QSM with gene expression are specific to this relationship and have not arisen spuriously due to gene-gene co-expression or spatial autocorrelation.

**Cell types linked with variation in brain cortical iron content**

We assessed whether the most significantly upweighted and downweighted genes associated with brain iron, as measured using QSM, were more strongly expressed in specific cell types. Cell-type expression was defined using the human-derived single-nucleus data from the Allen Institute for Brain Science and increased expression assessed using expression-weighted cell-type enrichment analysis. We found that the top 20% of upweighted genes (324 genes) were significantly enriched in astrocytes, glutamatergic and GABAergic neurons and oligodendrocyte precursor cells, implying a relatively higher proportion of these cell types in healthy brains in regions with a higher iron contents in Parkinson’s disease (Fig. 4B). We found that the top 20% of downweighted genes (214 genes) showed significantly greater expression in GABAergic and glutamatergic neurons, implying a higher proportion of these cells in healthy brains in regions with lower iron contents in Parkinson’s disease. Adjusting the expression-weighted cell-type enrichment analysis threshold to include the top 10, 30, 40 or 50% of upweighted or downweighted genes did not alter the cell transcription profile seen in either case, nor did defining cell-type expression using a different human-derived single-cell nucleus dataset (Supplementary Fig. 1).

**Association with Parkinson’s disease post-mortem-derived gene expression data**

We found that genes upregulated in Parkinson’s disease cortex relative to controls, as well as genes upregulated in
cortex in Parkinson’s disease dementia relative to control subjects,\textsuperscript{60} were more negatively weighted than expected by chance in our analysis ($P = 0.0002$, $P < 0.0001$, respectively). Similarly, genes identified as being upregulated in cortex amongst individuals with dementia with Lewy bodies relative to controls\textsuperscript{61} were significantly more downweighted than expected by chance in our analysis ($P < 0.0001$), whereas genes found to be downregulated were significantly more upweighted ($P < 0.0001$). Additionally, we found that downregulated genes identified in Parkinson’s disease substantia nigra\textsuperscript{61} were more positively weighted than was due to chance in our analysis ($P < 0.0001$). Full results from the above weighting analysis can be found in Supplementary Table 8. Together, these results suggest that the gene expression profiles we identified based on susceptibility to cortical iron deposition and using baseline gene expression (as determined by the Allen Atlas dataset) are more perturbed in Parkinson’s disease than expected by chance, and this finding is replicable across independent sample cohorts and experimental approaches. Furthermore, these findings hold at later disease stages linked with Parkinson’s disease dementia and dementia with Lewy bodies.

**Discussion**

We demonstrated that the regional increases in magnetic susceptibility that are most likely due to brain iron accumulation are associated with the expression of genes involved in metal detoxification and synaptic function. Regional...
differences in the probable proportion of astrocytes, glutamatergic and GABAergic neurons and oligodendrocyte precursor cells were also present. Increased QSM susceptibility values were found in prefrontal and posterior parietal cortices, consistent with our previous findings, and were also present in insular and cingulate cortices, consistent with our previous

Our finding of gene expression related to the detoxification of other heavy metals is intriguing. This is likely to be explained by the fact that iron-copper homeostasis is controlled by proteins involved in both the metabolism of iron and copper. As well as iron, altered levels of copper occur in Parkinson’s disease. Reduced copper is seen alongside increased iron in the substantia nigra in Parkinson’s disease, whereas increased copper has been reported in the CSF. Free copper can catalyse reactions generating reactive oxygen species in a manner similar to iron and has similar potential to cause cell damage. However, some copper binding and transport proteins, notably metallothioneins, may have a neuroprotective role. The altered binding of copper by cuproproteins in Parkinson’s disease could contribute to the inability of cells to deal with an increased oxidative load, which is in turn aggravated by a large pool of labile iron. Copper has a high binding affinity with α-synuclein and is more potent than iron in aggregating α-synuclein, as well as being implicated in the formation of amyloid-β plaques and neurofibrillary tangles. When bound to α-synuclein it also has the capacity to reduce ferric to ferrous iron, facilitating further reactive oxygen species.

Disturbances in synaptic function are known to play a role in vulnerability and progression in Parkinson’s disease. A number genes mutated in familial forms of Parkinson’s disease are involved in synaptic function and autophagy, and genes implicated in sporadic Parkinson’s disease also relate to mechanisms of synaptic homeostasis. Aberrant synaptic autophagy and excess synaptic pruning may lead to downstream synaptic loss, which is expected to precede neurodegeneration.

The distribution of gene expression could be explained partly by variation in the distribution of particular cell types,
leading us to perform an expression-weighted cell-type enrichment analysis. We found that upweighted genes showed significantly greater expression in astrocytes and glutamatergic and GABAergic neurons and oligodendrocyte precursor cells. The finding of relative enrichment of astrocytes in regions with high levels of brain iron is intriguing, as astrocytes play an important role in brain iron uptake and metabolism, and in distributing iron to other neuronal cells. Enrichment in glutamatergic neurons is also of note. Higher levels of brain iron have been reported to correlate with higher N-methyl-D-aspartate receptor overactivity, and the increased activity of these receptors may also stimulate the release of iron from the lysosome. Glutamate-mediated excitotoxicity has been linked with neurodegeneration in Parkinson’s disease. Glutamate induces Parkin accumulation in mitochondria in a calcium- and N-methyl-D-aspartate-receptor dependent manner, and α-synuclein may perturb intracellular calcium levels via α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors.

We compared the pattern of cortical gene expression from our analyses against differentially expressed gene lists identified in a number of external post-mortem datasets in Parkinson’s disease, Parkinson’s disease dementia and dementia with Lewy bodies. We found that genes altered in these conditions were linked with the profile of healthy brain gene expression underlying higher magnetic susceptibility in Parkinson’s disease. This provides further evidence for processes involving brain iron in the selective vulnerabilities driving degeneration in Parkinson’s disease and Lewy body dementia.

Our findings demonstrate that regions showing increased iron deposition in Parkinson’s disease have a functional abundance of cells and proteins involved in the homeostasis and detoxification of metals in the healthy brain as well as in synaptic function. Any disturbance of such pathways, as has been reported in Parkinson’s disease and models of Parkinson’s disease, could potentially render these regions vulnerable to excess iron accumulation, aberrant iron processing and subsequent downstream oxidative stress, proteinopathy and cell death.

**Limitations**

Our main analysis used gene expression data derived from donors unaffected by neurological or psychiatric disease. Spatial expression profiles are therefore based on unaffected individuals and conclusions only apply to this intrinsic variability, not to changes in gene expression that occur in Parkinson’s disease.

Because of the availability of data in the Allen Human Brain Atlas, we used only left hemisphere data, as data from the right hemisphere was only available from two brains. There is a laterization effect for the distribution of QSM susceptibility, with higher magnetic susceptibilities in the right compared with the left hemisphere. Differences between hemispheres could be examined in future work when gene expression data are available for both hemispheres from a larger number of donors. When Parkinson’s disease gene expression data with widespread cortical coverage become available, it will be of interest to relate gene expression in Parkinson’s disease directly to brain iron levels. We were unable to validate our PLS analysis with any additional healthy gene expression databases as those currently available are derived from only a small number of cortical regions.

We used QSM to measure increased iron deposition in Parkinson’s disease and investigated the genetic makeup of implicated regions, as there is strong evidence implicating neurodegeneration as a process downstream of iron-induced oxidative stress. However, QSM is unable to capture all drivers of neurodegeneration, especially those that occur...
Iron deposits have been shown to co-localize with amyloidoid and tau pathology. How these contribute to one another and lead to neurodegeneration is not known. Future studies of this kind could incorporate contemporaneous data on brain amyloid and tau, e.g. using amyloid or tau PET, with QSM and transcriptomic data to shed light on the biological processes which may contribute to this.

Our cell-type analysis was performed using the Allen Institute for Brain Science single nucleus RNA-sequencing dataset, which is based on the profiling of cortical brain samples. Future analyses could specifically examine the role of cells in other brain regions, for example midbrain dopaminergic neurons.

Participants in our study were scanned whilst taking their usual dopaminergic medications. There is no evidence that L-DOPA causally affects brain tissue iron content or magnetic susceptibility, and L-DOPA does not alter cortical iron levels in a mouse model of Parkinson’s disease. In humans, magnetic susceptibility in subcortical regions of interest analyses do correlate with L-DOPA dose, but whether this relates specifically to the effects of L-DOPA or is a function of disease severity is not yet known. The effects of being ON versus OFF L-DOPA on cortical QSM magnetic susceptibility could specifically be examined in future work. Finally, our study was cross sectional. Longitudinal studies of brain iron accumulation in patients with Parkinson’s disease progressing to dementia will provide further insights into the temporal order of brain iron accumulation in specific brain regions and the associated biological processes.

### Conclusions

Regional increases in magnetic susceptibility in Parkinson’s disease, most likely due to brain iron accumulation in frontal, cingulate and insular cortices, are associated with a distinct gene expression profile. In these regions, we found higher intrinsic levels of gene expression relating to heavy metal detoxification and synaptic function as well as a probable relative abundance of astrocytes and glutamatergic neurons. These findings shed light on the processes driving neurodegeneration in Parkinson’s disease and the selective vulnerabilities of brain regions that are most affected, providing potential insights into future therapeutic targets to slow the progression of neurodegeneration in Parkinson’s disease.

### Funding

G.E.C.T. is supported by a PhD studentship from the Medical Research Council (MR/N013867/1). A.Z. is
supported by an Alzheimer’s Research UK Clinical Research Fellowship (2018B-001). M.R. is supported by a UK Medical Research Council Tenure-track Clinician Scientist Fellowship (MR/N008324/1). K.S. is supported by ERC Consolidator Grant DiScO MRI SFN 770939. A.L.G.-M. is supported by Fundación Séneca (21230/PD/19). P.M. is supported by the National Institute for Health Research. R.S.W. is supported by a Wellcome Clinical Research Career Development Fellowship (205167/Z/16/Z). Recruitment to the study was also supported by Parkinson’s UK and the Cure Parkinson’s Trust. The study was further supported by UCLH Biomedical Research Centre Grant (BRC302/NS/RW/101410) and by grants from the National Institute for Health Research.

Competing interests

R.S.W. has received personal fees from GE healthcare. J.A.-C. has equity and a full-time appointment at Tenoke Limited.

Supplementary material

Supplementary material is available at Brain online.

References


