Nigella Sativa Prevented Parkinson's-Like Motor Functions Impairment, Dopamine Depletion and Neuronal Degeneration in the Striatum of Mptp-Induced Balb/C Mice

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Research Article

Keywords: Nigella sativa, MPTP, BALB/c, Parkinsonism, Selectivity, Striatum, Frontal cortex

DOI: https://doi.org/10.21203/rs.3.rs-522823/v1

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Abstract

Background

Parkinsonism is a neurological disease characterised by dopaminergic neuron degeneration in the substantial nigra and dopamine deficiency in the brain, with motor and psycho-cognitive implications, while limitations masked the efficacy of the available drugs, thus the need to find alternatives with less side effects are essential. *Nigella sativa* is a multi-potent plant with therapeutic potentials in the brain and other body organs. This study investigated the effects of *Nigella sativa* oil (NSO) on the cognitive and other Parkinsonism endophenotypes elicited by MPTP in the BALB/c strain mice.

Materials and Methods

Body weights, brain-body ratios, recognition memory (through novel object recognition test), as well as fronto-cortical, striatal and cerebellar dopamine and neuronal density were assayed in thirty-two (32) male BALB/c mice (18 g – 25 g). They were randomized into four groups exposed to; normal feed, 18 mg/kg MPTP i.p, 1 ml/kgbw NSO p.o., and NSO + MPTP respectively, for 5 consecutive days. Behaviours were analysed 24 hours after the last exposure, subsequently euthanized, the brains removed and processed for biomarkers analysis and histochemistry.

Results

Parkinsonism-like traits such as mild tremor, down-regulation of striatal and fronto-cortical dopamine and neurons were recorded in the BALB/c mice administered with MPTP only. However, significant increase (p < 0.05) in appetite, body weight, brain-body weight ratio, and recognition memory was also recorded in the MPTP-administered mice, though *Nigella sativa* was significantly prophylactic against the negative Parkinsonic features, and ‘moderative’ of the up-regulations induced by MPTP.

Conclusion

While this suggests selective MPTP sensitivity and resistance in BALB/c strains, this study recommends the investigation of possible (though ironic) beneficial potentials of MPTP.

1. Introduction

Parkinson's disease (PD) is a degenerative disorder of the central nervous system that mainly affects the motor system (NINDS, 2019). It is characterised by progressive neuronal degeneration which predominantly affects the dopaminergic neurons in the nigrostriatal system and several other regions of the brain (Roberts et al., 1994). The exact cause of Parkinson's disease (PD) is unknown, but epidemiological studies suggest an association with environmental toxins. Early in the disease, the most
obvious symptoms are rigidity, difficulty with walking, affected thinking and other behavioural problems (NINDS, 2019). Other symptoms include sensory, sleep, and emotional problems. The main motor symptoms are collectively called "Parkinsonism", or a "Parkinsonian syndrome" (Williams & Litvan, 2013).

The striatum is a principal component of the basal ganglia, mostly studied for its roles in facilitating voluntary movement, rewards and motivations, but with now recognised influences in cognition and other behaviours (Dingman, 2015). Furthermore, the neuronal degeneration and behavioural deteriorations in Parkinson's disease is evidently shown to be unlimited to the basal ganglia due to the observed non-motor symptoms, but to other high function areas of the brains, like the frontal cortices and cerebellum with established structural and functional communications with the basal ganglia (Drag et al., 2009) (Ichinohe et al., 2000).

The cerebellum has also been reported with pathological changes in Parkinson's disease, even though its major roles are thought to include pathological and compensatory effects. This is as morphological and functional or modulations were detected in the cerebellum in relation to tremor, akinesia/rigidity, gait disturbance, dyskinesia and some non-motor symptoms(Wu & Hallett, 2013).

Dopamine supplementation remain the most efficient therapy in management of, and/or slowing the progressions of Parkinson's diseases, with extensive undesired outcomes. Thus, the search for alternative or supplementary regimen, to further slow the disease progression, prevent, rescue or salvage the degenerating dopaminergic systems. *Nigella sativa* is one of the most widely used medicinal plants across the world (Ahmad et al., 2013), with a long history of use across ancient Egyptian, Indian (Unani), Greek (Ayurvedic), Roman and Islamic cultures (Al-Naqeep et al., 2011). With therapeutic effects on several body systems in humans and animals (Assi et al., 2016), *Nigella sativa* has also proven useful in the treatment of psychiatric disorders (Randhawa & Alenazi, 2016). However, due to the paucity of information on its impacts on MPTP-induced Parkinson-like symptoms, particularly on striatal, fronto-cortical and cerebellar parameters, this research aimed at filling this knowledge gap.

This study thus investigates the possible neuro-therapeutic roles of *Nigella sativa* oil in Parkinson's-like endophenotypes, through the analyses of dopamine levels, neuronal density and histopathology of the striatum, frontal cortex and cerebellum and associated functions in MPTP exposed BALB/c mice.

## 2. Materials And Methods

### 2.1. Acquisition of research materials

Sixty (60) mls of purified *Nigella sativa* oil was procured from Hemani® International, Pakistan. Similarly, 1-methyl-4-phenyl-1-2-3-6-tetrahydropyridine (MPTP) was procured from Medchem Express, New Jersey, USA.

The forty-eight (48) adult male BALB/c mice used for the study (with weight ranging from 18 g to 25 g) were in-bred at the Neurophytotherapy research lab (NPTRL) in the faculty of Basic Medical Sciences,
where they were housed in well ventilated cages improvised at the lab and under controlled temperature and humidity.

2.2. Experimental Design and Dosing

The mice were allowed to acclimatise for one week, randomly divided into four (4) groups, containing twelve mice each, and exposed to; normal feed (Control group), 18 mg/kg MPTP intraperitoneal (MPTP model group), 1 ml/kgbw NSO orally (NSO group), and NSO + MPTP (MPTP model and NSO intervention group) respectively, for 5 consecutive days.

Following respective administrations, the mice were placed on video surveillance for one hour each, to observe their behavioural responses to the respective regimens, especially the MPTP.

2.3. Measurement of Body Weight

The body weights of the animals were measured at the commencement and end of the study using a digital weighing balance (KERRO®, 2016; 0.1g accuracy) in order to check for weight gain and/or loss across the groups.

2.4. Novel Object Recognition (NOR) Test

The Novel Object Recognition (NOR) test was performed for each group of mice, 24 hours after the final administration. The test involved the habituation, training and testing phases as adapted from the (SBFNL(b), 2019) protocol. During the habituation phase, each mouse was introduced into an open field maze (38 cm x 38 cm x 38 cm) to get habituated for 5 mins. During the training phase, each mouse was placed into the open field maze for another 5 mins, this time containing objects A and B, both of which are similar in shape, colour, size and height. They were placed diagonally within the maze and equidistant from each other. During the testing phase however, the mice were again introduced into the open field maze, but this time, with object B already replaced with a novel object C at the same location. This phase also lasted for 5 minutes. The test was tracked with a video camera, and the time spent by each animal in exploring the novel object C was recorded against the time spent exploring the familiar object A and the discrimination index was determined as a percentage of the novel object exploration time over the total time spent exploring both objects.

2.5. Open Field Test

The open field test was performed as adapted from the (SBFNL(c), 2020) protocol. Mice were picked at random from the group concerned and placed at the centre of the maze to commence the test. Movement of the mice over the maze grids was surveilled and the number of lines crossed, grooming postures, walling postures and hinding (rearing) postures were recorded. Each mouse spent a total of 5 minutes in the maze.

2.6. Balance beam walk test
The balance beam walk test was performed in line with earlier defined protocols (Luong et al., 2011). The mice were picked randomly and placed on the end of the rods facing away from the balance beam. The time it took the animal to turn $180^\circ$ and the time taken for the animal to reach the safety zone at the attachment point of the rod were both recorded. Each mouse was also given 5 minutes to undergo the test.

**2.7. Sacrifice**

Following administration and neurobehavioral assays, the animals were euthanized. Each group of mice was randomly divided into two categories. The first were sacrificed by prompt decapitation, for immediate excision and homogenization of the striatum, frontal Cortex and cerebellum in 0.1 M phosphate buffered saline (PBS) for further respective neurotransmitter assay. The others were euthanized with 0.1 ml ketamine hydrochloride (i.p.), followed by transcardiac perfusion with 0.9 % normal saline for two minutes, and then 0.1 M phosphate buffered saline (PBS). Perfused brain tissues from the latter were subsequently fixed in formal calcium solution for twenty-four hours and taken through the routine H&E processing.

**2.8. Brain Weights**

The respective weights of the excised brains were measured, and the relative brain weights (RBW) of the animals across the groups was calculated using the formula:

$$RBW = \frac{\text{Brain Weight}}{\text{Body Weight at sacrifice}} \times 100$$

**2.9. Dopamine assay**

For the determination of Dopamine, the tissue homogenate was mixed directly with 6-Aminoquinoly-N-hydroxy-succinimidy carbonate (AQC) at room temperature (Bergh et al., 2016; Gottås et al., 2015). The resultant samples were reacted with ninhydrin reagent to form a colour reaction. A 100 ml of ninhydrin reagent was prepared by mixing 16 ml of 0.6 M with 58.8 g/Litre Phosphoric acid, 64 ml of glacial acetic and 1 g ninhydrin. Thus, 550 µl of ninhydrin reagent was added to every 50µl of the sample (mixture containing AQC) in 5 ml screw capped pyrex tubes. Known concentrations of Standard solution of the neurotransmitter (Dopamine) were prepared as the sample. The tubes were heated for one hour in 100°C water bath. Afterwards, the tubes were cooled at room temperature and 160 ml of formic acid was added in these tubes. Optical densities were thereafter read at 310 nm wave length concentrations of the tested sample. Dopamine concentration was then calculated as follows:

$$\text{Dopamine (µgg}^{-1}) = \frac{\text{Absence Reading} \times \text{slope (Standard)}}{\text{Dilution factor} \times 10,000}$$

**2.10. Statistical analysis**

Data acquired were expressed as mean ± standard error of mean, and were analyzed with one-way analysis of variance (ANOVA) using the GraphPad Prism (version 7.0) software. Statistical significance was set at $p < 0.05$. 

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3. Results

The following are the recorded results at the end of the research.

3.1. Physical observations

Following the first administration session, the MPTP mice expressed mild tremor, restlessness and reduced locomotion across distances within the cage, while their food and water consumption rates increased appreciably. The feed consumption rate for the NS group was however lower than the control, even though, all other groups apart from the MPTP maintained their normal levels of activity after their respective administrations throughout the study. The activity levels of the NS + MPTP mice however increased slightly upon commencement of MPTP administration, even though this did not reach the level of the MPTP mice. Furthermore, an idiopathic scoliosis (IS) developed in one of the MPTP mice on the fourth day of administration, as illustrated in appendix I. The animal was thereafter isolated for observation and possible recuperation. However, the animal never recovered until it died three days afterwards.

3.2. Body Weight Changes

We describe the mean difference in body weights (final weight – initial weight) of the animals (Fig. 1), where the MPTP mice had the highest and thus significantly higher mean body weight gain than mice of all other groups. Also, NS group had the least gain in weight when compared with the Control group while NS + MPTP group had a significantly lesser body weight gain than the MPTP group even though higher than the NS group.

3.3. Relative Brain Weight

The following are percentages of brain-to-body weight ratios obtained from the present study (Figure 2). While the administration of MPTP led to a significantly higher (and in fact highest) relative brain weights as seen in MPTP group, pre-administration of *Nigella sativa* rather prevented and counter-acted this hike, resulting in an insignificantly lower RBW in the NS+MPTP group than the control and MPTP groups. The NS group however maintained a slightly higher RBW than the Control and NS+MPTP groups.

3.4. Novel Object Recognition Test

A significantly higher novel exploration time was observed for the induced Parkinson's-like group (MPTP) than all other groups (Fig. 3). Pre-treatment of such Parkinson's-like group with *Nigella sativa* as seen in ‘NS + MPTP’ yielded a novel exploration time similar mice administered the treatment alone (NS) without the induction of Parkinson's. NS + MPTP and MPTP however had higher exploration time than the control mice.

3.5. Open Field Maze Test
Of the open field parameters assessed (Fig. 4), there was a decrease in the locomotion of the MPTP mice as evident in the significantly decreased number of lines crossed and frequency of walling when compared with other groups. Both were however prevented in the NS + MPTP group as seen in the significantly higher line crossings maintained, similar to the level recorded in the NS group.

3.6. Balance Beam

In this balance beam walk assay (Figure 5), we found the fastest turning interval and rod traversing time in the NS mice, while on the converse, MPTP and NS+MPTP groups took longer times than the control, with MPTP being the slowest to reach the safety point, even though it took a shorter time than the pre-treated NS+MPTP to turn 180 degrees.

3.7. Dopamine Assay

The following are the results obtained from the dopamine assay

As seen in Fig. 6, striatal dopamine level was significantly down-regulated in the MPTP mice, while the fronto-cortical and cerebellar dopamine levels in the MPTP mice were rather at par with or higher than in NS mice respectively. The NS + MPTP mice therefore had a higher striatal dopamine level than the parkinsonic (MPTP) mice.

3.8 Hematoxylin and Eosin Stains

Figure 7 above shows the striatal, fronto-cortical and cerebellar histology across the experimental mice, with striatal neurons and nigrostriatal bundles indicated in full and dashed arrows respectively. The neuronal density in the MPTP mice was the least while the NS+MPTP mice expressed significantly higher striatal neurons than in all other groups. No significant variation was observed across the fronto-cortical and cerebellar histological architectures.

4. Discussion

MPTP had been characterised as a neurotoxin in humans and rodents, with irreversible Parkinson's-like features such as tremor, rigidity, and slowness of movement (bradykinesia), postural instability, and freezing. The above motor symptoms were observed in the present study upon administration of MPTP in validation of earlier reports by Aarsland et al. (2009), Jellinger (1998), and Meredith & Rademacher (2011), in which rigidity, slowness of movement, postural instability, and freezing were reported in MPTP-administered humans, non-human primates, cats, rabbits, and some rodents strains. Furthermore, this study further shows that BALB/c mice are also sensitive to MPTP with respect to these Parkinson's-like features. However, these parkinsonic traits were absent in the mice pre-administered with Nigella sativa (NS + MPTP), indicating the prophylactic potential of Nigella sativa in preventing these Parkinson's-like symptoms.
Upon macroscopic examination, no detectable insult or injury was observed in the brains across all groups. However, idiopathic scoliosis (IS) was observed in one of the MPTP-administered mice. Described as a multifactorial disease involving many intrinsic factors such as genetics, imbalance of muscle structures, abnormal growth of vertebral bodies, asymmetrical growth of the neurocentral cartilage, length discrepancy between spine and spinal cord, abnormal platelet calmodulin, and abnormality in melatonin metabolism (Banala et al., 2018; Machida, 2018); the idiopathic scoliosis observed in one of the MPTP mice may however be an epiphenomenon rather than being consequential to the MPTP administered, as has been said of many hypothesised factors (Machida, 2018).

Human PD patients have been reported to experience low body weight, a phenotype hypothesised to be predisposed by many factors like dysphagia, chewing difficulty, impaired hand-mouth coordination, and hyposmia (Bachmann & Trenkwalder, 2006; Ma et al., 2018). Similar body weight trend has also been reported of non-human primates (Porras et al., 2012). Despite this trend however, no significant weight loss was recorded in C57BL/6 mice by Sundström et al. (1990) during 4 weeks after MPTP exposure. A further obvious contrast to the trend above was the observable body weight changes in the current study, in which the MPTP mice had the highest mean body weight increase compared with all other groups. While this may be explained by the increased food and water consumption rates observed in the mice, it is a further validation of the report in which 32.9% of PD patients studied by Cersosimo et al.(2018) showed increase in their body weights. This study thus shows MPTP as increasing MBW in BALB/c mice. The increased appetite and the consequent body weight gain also raises the question about the tendency of MPTP or perhaps some of its active constituents to to predispose exposed animals to obesity, as found in the BALB/c mice here studied. It may also be inferred that MPTP does not adversely affect the weight and that the growth rate is also normal in BALB/c’s mice, an effect that is yet to be understood.

However, the weight gain in the NS mice was significantly lower and in fact the least when compared to other groups. This is similarly explained by the feed consumption rate which was lower in the NS group than observed in all other groups .Reduced weight gain as recorded in this study, is a phenomenon that has been previously characterised to Nigella sativa exposure, in line with the findings of Bano et al. (2009) and Le et al. (2004), such that its significantly repressive effect on weight gain is obvious. This weight limiting effect of Nigella sativa was thus seen on the NS + MPTP mice which had a significantly lesser MBW than the MPTP group but higher than the NS group. Nigella sativa was thus able to limit the weight gain in this group, suggesting its ability to repress/counter the weight increasing effects of MPTP.

Relative brain weight (RBW), also known as brain-to-body weight ratio, is a hypothetical estimate of the brain functions of an animal, as it is believed to coincide much better with the observed cognitive abilities than absolute brain size (Ciáro, 2011). In this study, the highest RBW was recorded in the MPTP mice, possibly due to a progression of the brain weight along with the body weights of the MPTP animals as discussed above. The NS + MPTP mice however recorded significantly lower RBW than the MPTP mice, indicating the impact of NS pre-exposure in keeping the RBW at par with the level recorded in the NS mice.
Novel object recognition test (NORT) is a measure of recognition memory which harnesses the innate tendency of mice to explore novelty of their environment (SBFNL(b), 2019). The MPTP mice showed the highest exploration time with the novel object in the assay, when compared with other groups, thus accruing the highest recognition memory to the MPTP mice. This finding is inconsistent with the previous reports in which mild cognitive impairment and dementia are established as pathophysiological features of Parkinson’s disease at the early and later stages respectively (Weil et al., 2018). Recognition memory deficit particularly has been further documented by some studies in patients with Parkinson’s disease (Chiaravalloti et al., 2014). The improved recognition memory found in this study may be explained by the higher brain-body weight ratio found in the MPTP mice, since the later, as earlier explained is hypothesised as a measure of brain function.

MPTP is a known neurotoxin with neurodegenerative potentials in human and animals (Langston, 2017). However, the memory and brain weight findings in the BALB/c mice administered with MPTP raises questions about possible advantageous impacts of this neurotoxin on some brain functions, even if such impacts are species or strain dependent.

The novel exploration time of the MPTP mice was significantly higher than recorded in the NS + MPTP mice, with the latter value similar to that of the NS mice. This may suggest the impact of *Nigella sativa* in moderating the novel object exploration time in the pre-treated mice.

The NS group behaved best by spending the least time to turn and the least time to reach the end of the beam. The MP animals however spent the longest time to reach the end of the beam, similar to the significantly (p < 0.0001) longer time reported of the *Aphakia* mice (also known as *ak* mouse, which is a striatal denervation model of Parkinson’s disease exhibiting the cell specificity of neurodegeneration observed in humans) to complete the balance beam test (Singh et al., 2007). *Nigella sativa* was however able to reduce this traversing interval for NSMP but not the turning time. This suggests a therapeutic effect of *Nigella sativa* on motor balance and coordination in Parkinsonic mice.

The neurotransmitter assay revealed the lowest striatal dopamine in the MPTP mice, in validation of previous studies which characterised Parkinsonism with deficiency of dopamine and dopaminergic neurons in the midbrain, especially in the striatum (Scherman et al., 1989; Triarhou, 2013). The striatal dopamine levels in the NS and control group however were similar and both were significantly higher than in the MPTP mice. This may explain the up-regulated dopamine level recorded in the NS + MPTP mice in comparison with the parkinson’s-like MPTP mice. This indicates the prophylactic role of *Nigella sativa* in Parkinsonism. The cerebellar dopamine level in the MPTP mice was however higher than all other experimental groups, albeit insignificantly statistically, while in the frontal cortex, there was almost no difference in the dopamine levels across the mice. This corroborates the localisation of Parkinsonic dopamine depletion to the striatum (Folarin et al., 2019; Meredith & Rademacher, 2011; Scherman et al., 1989; Sedaghat et al., 2014; Singh et al., 2007).

The striatal neuronal density in the MPTP mice was the least, in validation of reports by Ross et al., (2004) where the mean neuronal densities in all striatal quadrants were significantly lower in the PD
group compared with the other groups. This thus confirms the depletion of striatal neurons in BALB/c strains of MPTP mice models of Parkinsonism, in slight contrast to its being referred to as MPTP-resistant (Meredith & Rademacher, 2011). However, when pre-treated with *Nigella sativa* oil, striatal neuronal density was markedly preserved than in mice administered with MPTP only and *Nigella sativa* oil only. The fact, however, that the NS + MPTP mice expressed more striatal neurons than the MPTP mice, corroborates the prophylactic potential of *Nigella sativa* against Parkinsonic endophenotypes as earlier described with other data above. Strains of mice are known to differ in their response to MPTP with respect to the degree of striatal DA depletion, amount of loss of midbrain DA neurons, and behavioural deficits. However, while the hallmark of sensitivity and resistance in the substantia nigra - pars compacta (SNpc) was put at “>50% SNpc neuron loss” and “<25% SNpc loss” respectively (Meredith & Rademacher, 2011), the neuronal loss recorded in the BALB/c MPTP mice was only significant statistically, but not up to 20% of the control's neuronal density. A “>50% loss” was however recorded in the dopamine assays in the MPTP mice.

In the cerebellum and frontal cortex however, no significant morphological or pathological difference was observed between the groups across the molecular, ganglionic and granular layers of their cerebellar cortices; and across the layers of the frontal cortex.

### 5. Conclusion

In this study, MPTP was validated with some known Parkinsonic features such as tremor, down-regulation of dopamine and reduced density of striatal neurons in the BALB/c mice. The BALB/c mice may thus be considered MPTP-sensitive as regards these Parkinsonic features. But MPTP also increased the appetite, body weight, brain-body weight ratio, and recognition memory in the MPTP-administered mice. *Nigella sativa* was found to ameliorate the negative Parkinson's-like features earlier mentioned, as induced by MPTP administration, and also moderated the up-regulations effected by MPTP, as shown in its prophylactic roles against striatal neuronal and dopamine down-regulation.

Conclusively, oral administration of *Nigella sativa* oil prevented functional loss of striatal dopaminergic neurons in Parkinson's-like mice as investigated through the histological and neurotransmitter assays of MPTP-induced BALB/c mice, even though insignificant difference was observed in the frontal cortex and cerebellum of the same mice. By and large, this study corroborates the possible candidacy of *Nigella sativa* in the treatment of Parkinson's disease.

### Declarations

#### Conflict of Interest

The authors declare no conflict of interest.

#### Acknowledgements
FRO shows immense gratitude to Prof. Marina Bentivoglio and Pof. Evelyne Sernagor of the IBRO/ISN Writing papers workshop, 2019, for their guidance.

**Funding**

The research was executed without any external funding support.

**Conflicts of interest/Competing interests**

The authors have no conflicting or competing interests to declare.

**Availability of data and material**

Data will be made available upon justifiable requests.

**Code availability**

Not applicable

**Authors' contributions**

The work was conceived, designed and drafted by RF, who also coordinated the acquisition, analysis, and interpretation of data by AO, PO and PF. IA revised it critically for important intellectual content while all authors approved the version to be published and agree to be accountable for all aspects of the work.

**Ethics approval**

Ethical clearance (numbered BS/18/VII/23-010) was obtained for the study from the Anatomical Research Ethics Committee (AREC), Olabisi Onabanjo University, Sagamu, Nigeria.

**Consent to participate**

Not applicable

**Consent for publication**

Not applicable

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**Figures**
Figure 1

Graphs showing the mean body weight gain (A) and Mean Growth Curve (B) of the animals in different groups during the experiment (* and ** indicate increasing levels of significance at p<0.05)

Figure 2

Graph Illustrating Relative Brain Weights (RBW) in Experimental Mice (* and ** indicate increasing levels of significance at p<0.05)
Figure 3

Graph showing Novel Object Recognition in Experimental Mice (* and ** indicate increasing levels of significance at p<0.05)
Figure 4

Showing Open Field Maze test in Experimental Mice. (* indicates significance at p<0.05)
Figure 5
Graph showing animals’ scores on the balance beam walk test

Figure 6
Graph illustrating Dopamine level in Striatum, Frontal cortex and Cerebellum of the Mice(* and ** indicate increasing levels of significance at p<0.05)

Figure 7
Photomicrographs of the Striatum stained with H & E stains across all mice groups. Magnification = (X400). Full arrows indicate striatal neurons, while dashed arrows indicate nigrostriatal bundles.

Supplementary Files
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- AppendixI.docx