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Celastrol and thymoquinone alleviate aluminum chloride-induced neurotoxicity: behavioral psychomotor performance, neurotransmitter level and oxidative-inflammatory burden in brain of rats

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Abstract:
The exposure to metal aluminum such as aluminum chloride (AlCl3) induces inflammatory-oxidative reactions with progressive neurodegeneration in different brain regions in animal models. The current study was designed to assess the role of celastrol or thymoquinone (TQ) in alleviating AlCl3 induced behavioral psychomotor changes and oxidative-inflammatory burden in albino male rats. Four groups were used in this study, (i) vehicle control group, (ii) AlCl3 control group: rats received intraperitoneal injection (i.p.) of AlCl3 (10 mg/kg), (iii) AlCl3+TQ (10 mg/kg, i.p.) group and (iv) AlCl3+celastrol (1 mg/kg, i.p.) group. In general, all injections remained for 6 weeks. Behavioral psychomotor evaluation (open field test, rotarod test and forced Swimming test) were done to assess locomotor, motor coordination, anxiety-like behavior and depressive-like behavioral. Markers of oxidative stress, malondialdehyde (MDA), total antioxidant capacity (TAC) and catalase enzyme activity (CAT) and the proinflammatory mediators, tumor necrosis factor-α (TNF-α) and interleukin-6 (IL-6) were measured in the rat brains. Neurotransmitters including acetylcholine (ACh), dopamine and serotonin in addition to acetylcholinesterase enzyme (AChE) level were measured in brain homogenates. Our results demonstrated that daily injection of TQ or celastrol significantly improved behavior psychomotor deficits, decreased AChE activity towards their normal levels. Tissue oxidative stress and proinflammatory markers were modulated by TQ and celastrol. These results concluded that TQ and celastrol have useful in alleviating AlCl3-induced neurotoxicity by their antioxidant and anti-inflammatory properties. Hence, they are looking promising for investigating their preventive effect in animal models of neurodegenerative diseases.

Keywords: AlCl3; psychomotor activity; celastrol, inflammation; neurotransmitters; oxidative stress; thymoquinone.
Introduction

Neurotoxicity usually occurs when the nervous system is exposed to a variety of toxic compounds, such as aluminum, and manifests as psychomotor and cognitive symptoms (Troshin 2009). Aluminum has no biological function in the human body and is known to cause neurotoxicity, which has been linked to the onset of dialysis encephalopathy. Some dialysis patients are exposed to Aluminum in the dialysis fluid, resulting in neurobehavioral symptoms, seizures, and death (Baydar et al. 2003). In animal models, exposure to metal aluminum, such as aluminum chloride, caused inflammatory reactions and neuronal abnormalities at the synapse level, as well as progressive neurodegeneration in various brain and spinal cord regions. Aluminum has the ability to cross the blood–brain barrier, causing central nervous system functioning and behavioral problems (Baydar et al. 2003, Bhalla et al. 2010, Yokel 2000). There are numerous evidences that suggest that aluminum exposure and accumulation in the brain is linked to chronic debilitating, incurable, and highly prevalent neurodegenerative disorders such as Parkinson's (PD) and Alzheimer's (AD) by altering neurotransmitter metabolism, such as cholinergic system function, which is highly susceptible to aluminum, accumulating iron, promoting iron-dependent lipid peroxidation, and producing reactive oxygen species. However, these conclusions are still debatable and unclear (Ghorbel et al. 2016, Gitler et al. 2017, Skalny et al. 2021, Yokel 2000). Aluminum exposure may cause alterations in the levels of several neurotransmitters that are tightly linked. In addition, Aluminum influences the metabolism of various neurotransmitters in the brain of animals, including acetylcholine (ACh), serotonin, and dopamine, which has an impact on behavioral responses (Bhalla et al. 2010, Kumar & toxicology 2002, Skalny et al. 2021). Many neurotransmitter levels that are intimately linked may vary as a result of aluminum exposure (Kumar & toxicology 2002). It is critical to investigate the mechanisms of its neurotoxicity, which remain controversial in animal models, and to suspect a useful therapeutic approach to aluminum, a highly toxic metal with widespread distribution in human environments (water, food, drugs, cookware, and industrial sources), as well as remaining living organisms, for the prevention and treatment of its neurotoxicity (Brough & Jouhara 2020, Skalny et al. 2021).

Recently, the use of antioxidant agents derived from plant extracts and diet has been shown to be a promising therapeutic approach for preventing metal toxicity and neurodegenerative disorders linked to oxidative stress and inflammation, which activate microglia, causing increased production of proinflammatory cytokines, reactive oxygen species, and lipid peroxidation, and eventually apoptosis (Al-Otaibi et al. 2018, Allison et al. 2000).

Thymoquinone (TQ), a redox-active quinone oil obtained from Nigella sativa seeds (black cumin), possesses polyphenol characteristics and is generated from Nigella sativa seeds (black cumin). Anti-inflammatory and antioxidant properties, as well as the ability to pass the blood-brain barrier, scavenge free radicals and prevent cell damage caused by oxidative chemicals (Ahmad et al. 2013, Elmaci et al. 2016, Hosseinzadeh et al. 2012). In addition, it exhibits anti-anxiety and anti-depression properties in rats (Elmaci et al. 2016). Furthermore, due to its significant antioxidant and anti-inflammatory qualities, it has a neuroprotective impact in numerous neurological disorders such as PD and epilepsy (Majdalawieh et al. 2010). Its neuroprotective impact against heavy metals and radiation exposure has been demonstrated in a number of investigations (Elmaci et al. 2016, Kassab et al. 2017, Majdalawieh et al. 2010).
Celastrol is an antioxidant herbal plant that chelates metals like calcium and cadmium while quenching reactive oxygen species (Zhang et al. 2017). It is a quinone and phenolic methide triterpenoid isolated from the Tripterygium wilfordii Hook F (TWHF) plant, and it's one of the most popular natural medicine compounds, especially in China, for its anti-inflammatory and antioxidant properties (Li & Hao 2019). Celastrol has been proposed as a potential preventative and treatment for neurodegenerative disorders due to its antioxidant and anti-inflammatory effects (Li & Hao 2019, Paris et al. 2010). Celastrol also protected animal models from locomotor dysfunction and social behavior impairment (Bove et al. 2020).

The current study was designed to assess the effects of TQ and Celastrol, two effective antioxidant and anti-inflammatory herbal plants, in aluminum chloride (AlCl₃) induced neurotoxicity in rats, with a focus on behavioral psychomotor performance, neurotransmitters changes, and oxidative-inflammatory burden in whole brain of albino rats.

Materials and Methods

Animals
Twenty-four adult male albino Wister rats weighing 175-187 g were used in this study. Animals were purchased from The Ophthalmology Research Institute (Giza, Egypt); they were housed in spacious wire mesh cages at room temperature and were kept with free access to standard rat chow diet and tap water. Rats were left for acclimatization for one week before the start of the study. Any animals with overt signs of illness were removed from the study. All experiments approved by the committee for animal care and ethics at the medical college of Suez Canal University with approval No. 4741.

Chemicals
AlCl₃, TQ and celastrol were purchased from Sigma-Aldrich Co. Weekly stock fresh solutions were prepared as follow: ALCL3 was dissolved in distilled water however, TQ and celastrol were dissolved in 1% dimethyl sulfoxide (DMSO).

Groups and design of the study
The rats were randomly divided into 4 groups of 6 rats. Group I (normal group): rats were given 1.0 ml normal saline once daily, Group II (AlCl₃ group): group of neurotoxicity, AlCl₃ (10 mg/kg/day) was given intra peritoneal (i.p.) once daily and this dosing regimen of AlCl₃ was selected based on a previous report because of its high rate of induction and low mortality (Khan et al. 2013), Group III (AlCl₃+TQ group): TQ was concomitantly administered (10 mg/kg/day, i.p.) once daily with AlCl₃ (10 mg/kg/day) (Al-Ali et al. 2008, Ince et al. 2013) and Group IV (AlCl₃+Celastrol group): Celastrol was concomitantly administered by i.p. injection (1 mg/kg/day); this is the dose recommended in models of neurodegenerative diseases once daily (Paris et al. 2010).

There was a week of acclimatization followed by a period of 6-weeks for the rat experiment. The drug preparation and injection were carried out under aseptic conditions and recorded the body weight changes by electronic animal weight scale of all the rats at the beginning of the experiment
as the initial record for rats, then finally before sacrificing them to calculate the final weight for each rat.

**Behavioral evaluation**

**The open-field test for evaluation of locomotor behavior**
The open-field arena (100×100×40 cm) was made of wood. The floor was painted with white lines that formed 25 squares (20 × 20 cm) pattern. Rats were introduced individually in the open field arena and rat behavioral parameters were observed and video-recorded for 5-min sessions; the test was performed under high-light conditions. Rats were observed for ambulation (the number of squares crossed; only when the animal enters a square with both its forelimb (one count is made). An experienced observer, who was blinded to the treatment groups, quantified the open-field locomotor behavior. In addition, number of rearing; i.e. standing on hind limbs (vertical movement) and leaning on the wall with forelegs, and looking around were also observed (Zghari et al. 2018).

**The rotarod test for evaluation of motor coordination**
Animals were monitored for motor coordination and balance with the help of rotarod apparatus (3 cm in diameter and rotating at a constant speed of 20rpm). Initially, animals were habituated to maintain posture on the rotarod by giving two training sessions of 5 min each with a gap of 10 min between the two sessions. After training, animals were allowed to move over the rotarod and their falling number was recorded per 5 min (Metz et al. 2005).

**The open-field test for anxiety-like behavior evaluation**
Another arena of 25 squares was divided into 9 central and 16 peripheral squares. At the beginning of the 5 min session, the rat was put in the center of the arena and its behavior was registered by mobile camera for manual analysis. The parameters of time spent by rats in the center of the arena (anxiolytics area) and in the peripheral area (anxiogenic area) were calculated and were used as a measure of anxiety level (Zghari et al. 2018).

**Forced swimming test for depressive-like behavioral**
The session of the forced swimming test was started by putting the rat in a cylindrical container (50 x 20 cm) filled with 30 cm of 22 °C water for 5 min). The time spent immobile (when rats stopped all active behaviors and became inactive floating) was considered as an index of depression-like behavior. High time of floating was analyzed as an increased depressive-like response (Rashwan et al. 2018).

**Preparation of brain homogenates**
The whole brains were dissected out from the rats after sacrificed them at the end of study duration and psychomotor evaluation by decapitation, cleared of the adhering tissues, weighed, washed with cold phosphate buffer saline and homogenized in normal saline (10% W/V). The homogenates were centrifuged at 3000 rpm for 10 min. The resulting supernatant was kept at-80°C for the following measurements (Al-Otaibi et al. 2018).
**Neurotransmitters evaluation**

ACh and its hydrolysis enzyme, AChE in the brain supernatant homogenate were measured by the using a choline/acetylcholine assay kit (Bio Vision Inc., Co., California, USA, catalog #: K615) as a quantification colorimetric method. Also, brain AChE level was evaluated calorimetrically by a kit from BioVision, (catalog #: k764). In addition, serotonin and dopamine were measured colorimetrically and their ELISA kits also from (BioVision Inc., co, California, USA, catalog #: E4294 and K4219) respectively.

**Measurement of brain oxidant-antioxidant markers**

Brain lipid peroxidation was measured as MDA (nmol/g tissue) in the supernatant of the homogenized brain by the thiobarbituric acid colorimetric method using the MDA assay kit to make a pink colored complex with absorbance level at 532 nm from Cell Biolabs (USA, Catalog #: STA-832). On the other hand, the TAC was evaluated by a colorimetric method based on the ability of brain antioxidants to suppress the production of thiobarbituric acid reactive substances (TBARS) by using ELISA kit (Cell Biolabs, USA, Catalog No.: sta-360). CAT was measured in the supernatant of the homogenized brain by ELISA kit from Cusabio Co. (Wuhan, China, Catalog No.: CSB- E13439r).

**Measurement of brain inflammatory markers**

TNF-α and IL-6 levels: these levels were measured in the brain homogenate using a rat TNF-α ELISA kit from Cusabio Co. (Wuhan, China, Catalog No.: CSB-E11987r) and a rat IL-6 ELISA kit from the same company, (Catalog No.: CSB- E04640r) use the quantities sandwich based immunoassay technique to measure them. We followed the steps and instructions of all previous kits that were used in the present study according to their manufacturers.

**Statistical analysis**

The data were presented as means ± standard deviation (SD). Statistical Analysis of data was done using the SPSS package (Inc., Chicago, USA) and the variance of one way (ANOVA) for non-parametric data to elucidate differences between all groups. Post hoc range test (Bonferroni’s) was used to test the difference between each pair of means. The level of significance was at p<0.05.

**Results**

**Animal body and brain weights**

The final animal's body weight showed a significant decrease in the AlCl3 group than the control group. AlCl3+TQ and AlCl3+celastrol groups showed a significant gain in final body weight than the AlCl3 group with significant superiority for TQ. The initial body weight and brain weight showed non-significant change in all groups (Table 1).
Table 1. Effect of AlCl3, TQ and Celastrol on body and brain weights.

| Groups               | Initial body weight (g) | Final body weight (g) | Brain weight (g) |
|----------------------|-------------------------|-----------------------|-----------------|
| Control              | 180.22 ±3.79            | 270.4± 4.19           | 2.11± 0.41      |
| AlCl3                | 178.99±3.34             | 239.2± 3.54a          | 1.8± 0.24       |
| AlCl3+TQ             | 177.23± 1.27            | 262.3± 5.44ab         | 1.9± 0.19       |
| AlCl3+Celastrol      | 179.42±3.01             | 250.7± 4.92 abc       | 1.92± 0.07      |

Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05. a, b and c Represents a statistically significant difference when compared to Control, AlCl3 and AlCl3+TQ, respectively.

Behavior and locomotor tests

Evaluation of locomotor function by the open field test

Behavioral locomotor performances of rats were evaluated by in the open field test as horizontal and vertical ambulation. There was a significant decrease in horizontal and vertical ambulation in the AlCl3 group as compared to the control group (34 ±7.29 vs. 90 ± 9.48). AlCl3+TQ and AlCl3+Celastrol groups significantly increased the results as compared to the AlCl3 group (78 ± 8.81 and 69 ± 8.24, respectively). There was no significant difference in horizontal and vertical ambulation between AlCl3+TQ and AlCl3+celastrol groups (Fig 1).

Rotarod test

The motor coordination and balance level of rats were evaluated by a rotarod test. There was a significant difference between all groups regarding the number of falling down in the rotarod test. AlCl3 produced disturbance in coordination of movement (number of falls = 14.53±.3.21 vs. 4.32±2.08 in the control group). AlCl3+TQ and AlCl3+celastrol groups showed better performance and lower number of falls with significant superiority for celastrol (9.56±1.86 and 6.01 ±1.47, respectively, Fig 2).

Anxiety-like behavior in the open field test

Anxiety level was evaluated in the open field as appeared in AlCl3 group that showed significant higher level of behavioral anxiety pattern than the control group [decreased spent time in central zone (anxiolytic area) and decreased grooming activity] (88.22± 7.41 vs. 145.31± 13.35 & 2.21±1.47 vs. 11.33±3.41, respectively). But all parameters of anxiety were significantly improved in AlCl3+TQ and AlCl3+Celastrol groups with significant superiority for TQ (Fig 3).

Depressive-like behaviors in the rat swimming test

The depression level of rats was measured by calculating the time of stability (immobility time) during physical activity in force swimming test. There was significantly increased immobility time in the AlCl3 group as compared to the normal group (16.23± 2.19 vs. 3.41±1.37). AlCl3+TQ and AlCl3+Celastrol significantly decreased this time (8.01±1.79 and 11.91±2.76, respectively) as compared to AlCl3 group with no significant difference between AlCl3+TQ and AlCl3+Celastrol groups (Fig 4).
**AChE enzyme and neurotransmitter evaluation**

There was a significant increase in acetylcholine esterase (AChE) level in rat brains in AlCl₃ group when compared with normal group. TQ and celastrol improved its level when compared with AlCl₃ group with no significant difference between them. There was a significant decrease in ACh level in rat brains in AlCl₃ group when compared with normal group. AlCl₃+TQ and AlCl₃+celastrol improved its level when compared with AlCl₃ group with no significant difference between them (Table 2).

| Groups       | AChE (nmol/min/mg of tissue) | ACh (nmol/mg of brain tissue) |
|--------------|------------------------------|------------------------------|
| Control      | 12.5 ± 1.95                  | 48.34 ± 2.79                 |
| AlCl₃        | 30.71 ± 2.87ᵃ                | 21.21 ± 1.76ᵃ                |
| AlCl₃+TQ     | 17.45 ± 2.89ᵇ                | 39.32 ± 1.73ᵇ                |
| AlCl₃+Celastrol | 15.82 ± 1.11ᵇ            | 37.41 ± 2.01ᵇ                |

Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05.ᵃ, b Represents a statistically significant difference when compared to control and AlCl₃, respectively.

There was significantly decreased in dopamine level in rat brains in AlCl₃ group when compared with normal group. AlCl₃+TQ and AlCl₃+Celastrol improved its level towards normal level when compared with AlCl₃ group with no significant difference between them. In addition, there was significantly decreased in serotonin level in brain rats of AlCl₃ group when compared with control group. AlCl₃+TQ and AlCl₃+Celastrol improved the dopamine and serotonin levels when compared with AlCl₃ group with significant superiority for TQ (Table 3).

| Groups       | Dopamine level (ng/g of brain tissue) | Serotonin level (ng/g of brain tissue) |
|--------------|--------------------------------------|----------------------------------------|
| Control      | 489.931 ± 4.47                       | 413.228 ± 1.81                         |
| AlCl₃        | 267.511 ± 7.62ᵃ                      | 215.772 ± 7.73ᵃ                         |
| AlCl₃+TQ     | 375.421 ± 14.04ᵇ                     | 352.881 ± 7.38ᵇ                         |
| AlCl₃+Celastrol | 363.322 ± 14.56ᵇ                  | 298.845 ± 11.74ᵇ                        |

Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05.ᵃ, b Represents a statistically significant difference when compared to control, AlCl₃ and AlCl₃+TQ, respectively.
Oxidative-antioxidative markers

Lipid peroxidation (LPO) as an oxidative marker in brain tissue was measured by its malondialdehyde (MDA) content. The level of MDA significantly increased in the AlCl₃ group when compared to the control group. AlCl₃+TQ and AlCl₃+Celastrol groups showed lower MDA level which is significantly different compared with AlCl₃ group with the superiority for TQ. The level of catalase (CAT) was significantly decreased in the AlCl₃ group when compared to the control group. TQ and celastrol improved the CAT level with significant differences with the AlCl₃ group and the superiority for celastrol. The total antioxidant capacity (TAC) was significantly decreased in the AlCl₃ group when compared to the control group. TQ and Celastrol improved its level with significant differences with the AlCl₃ group and the superiority for celastrol (Table 4).

Table 4. MDA, CAT and TAC levels in the brain tissue homogenates.

| Groups       | MDA (nmol/g of brain tissue) | CAT (µM of H₂O₂ decomposed/min/mg of brain tissue) | TAC (µmol/g) |
|--------------|------------------------------|--------------------------------------------------|--------------|
| Control      | 160.56±3.73                  | 45.34 ± 3.67                                     | 775.31 ± 20.28 |
| AlCl₃        | 270±8.36 a                   | 9.86 ± 2.56 a                                    | 219.42 ± 4.86 a |
| AlCl₃+TQ     | 230.32±8.78 ab               | 26.38 ± 2.46 ab                                  | 533.56 ± 17.86 b |
| AlCl₃+Celastrol | 200.74±8.82abc             | 39 ± 1.82 abc                                     | 570.26 ± 31.05 abc |

Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05. a, b and c Represents a statistically significant difference when compared to control, AlCl₃ and AlCl₃+TQ respectively.

Further, there was a significant increase in interleukin-6 (IL-6) and tumor necrosis factor-α (TNF-α) in the AlCl₃ group as compared to the control group. TQ and Celastrol significantly decreased the levels of these inflammatory markers as compared to the AlCl₃ group. AlCl₃+Celastrol group showed significant improvement in these inflammatory markers compared to the AlCl₃+TQ group (Table 5).

Table 5. Inflammatory markers in the brain homogenates.

| Groups       | IL-6 (pg/mg) | TNF-α (pg/mg) |
|--------------|--------------|---------------|
| Control      | 30.430 ± 3.94 | 20.102 ± 5.26 |
| AlCl₃        | 130.341±5.58 a | 100.163 ± 7.51 a |
| AlCl₃+TQ     | 70.025±6.87 ab | 55.902 ± 5.58 ab |
| AlCl₃+Celastrol | 57.301±2.88 abc | 40.040 ± 3.54 abc |

Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05. a, b and c Represents a statistically significant difference when compared to control, AlCl₃ and AlCl₃+TQ, respectively.
Discussion
Exposure to AlCl3 could alter blood brain permeability and induce neuronal structure changes and inflammatory-oxidative responses (Baydar et al. 2003). Molecules extracted from plants are now considered as complementary and/or alternative therapies against various neurodegenerative disorders that are associated with inflammatory-oxidative stress (Dai et al. 2019). The neurotoxicity effects of AlCl3 and neuroprotective effects of TQ and Celastrol are still not yet completely known. The current study was designed to investigate the psychomotor, some neurotransmitters and inflammatory-oxidative responses induced by AlCl3 and the neuro-preventive effects of TQ and Celastrol in albino rats.

Our findings showed that chronic exposure to AlCl3 produced a significant loss of final body weight when compared to the normal group. This result is in line with reports of other researchers who explained this loss on the basis of general toxicity of aluminum that affects energy metabolism through inhibition of glycolysis and Krebs cycle with promotion of lipid and protein oxidation (Han et al. 2013). The co-administration of TQ and Celastrol with AlCl3 significantly attenuated the loss in the final body weight that occurred by AlCl3 alone.

In the current study, the psychomotor behavioral responses in the AlCl3 group were assessed. In the open field test, the data from AlCl3 group revealed that the aluminum-toxicity significantly decreased the ambulation functions which are the number of squares crossed and the rearing movements compared to the control group. Similar observations were reported by Golub and Germann who noticed that the behavioral loco-motor activities significantly reduced after Aluminum administration by different doses (small, moderate and large) of aluminum (Golub et al. 2001). Moreover, there was a significant difference in the AlCl3 group regarding the number of falls in the Rotarod test when compared to the control group. Justin-Thenmozhi et al. reported significant decreases in the retention time of rotarod test with AlCl3 intoxication due to inflammatory markers such as (IL-6 and TNF-α) and oxidative stress (Justin-Thenmozhi et al. 2018).

The significant anxiogenic action of the AlCl3 toxicity was reported in the current study by parameters of the anxiety-like behavior of open field test when compared with the normal group and this finding is corroborated by several studies that investigated the aluminum effects on behavior in animals. A study on adult mice showed that the aluminum had anxiogenic action by decreasing the time spent in the central area of the open field (Bannon 2015). Depression was also shown in the AlCl3 group of our study as significantly increasing the stability time parameter in forced swimming test that reflects depressive-like behavior effect of aluminum toxicity. In a Wistar adult male rat model, the AlCl3 significantly increased immobility time parameter confirming its depressive like-behavioral action (Rashwan et al. 2018). The mechanism by which aluminum induces this behavioral change in animals is not yet known. Depression might link to hypo-function of the central serotonergic system and serotonin level reduction in brain regions (Ravi et al. 2000).

Regarding the inflammatory markers, in the present study, the AlCl3 group significantly increased the level of TNF-α and IL-6, the potent inflammatory cytokines in the homogenate brain of rats when compared with the control group. Milnerowicz et al. reported that the metal exposure such
as aluminum increased inflammatory cytokine TNF-α and some types of IL-6 and IL-1β which stimulate leukocytes recruitment and as the end result, this recruitment potentiates the inflammation stress by more releasing of pro-inflammatory cytokines (Milnerowicz et al. 2015). In addition, when the ACh is down regulated to a lower level in brain, the inflammatory system is activated by enhancing the expression of N-methyl-D-aspartate receptor, hence the secretion of inflammatory cytokines are increased that having anticholinergic activity forming vicious cycle augmenting the neurotoxicity of aluminum and more brain damage (Hori et al. 2013). This relation between inflammatory cytokines and lower levels of ACh with higher levels of its catalytic enzyme AChE is in line with our results. Aluminum neurotoxicity effects in previous studies produced imbalance between generation of ROS and antioxidants and induced oxidative stress in neurons. Regarding oxidative stress, we measured LPO as MDA level and quantified the endogenous antioxidant CAT enzyme in the homogenate brain of rats. Now, it is well documented that the brain has great risk to be injured by oxidative stress and LPO due to its high content of lipids (Belaid-Nouira et al. 2012). Our result showed a significant increase in MDA level in the homogenate brain of rats of the AlCl3 group when compared with the normal group and this finding reflects the potential role of aluminum in potentiation of lipid peroxidation, oxidative stress and brain damage on the basis of previous data.

In the present study, the AChE enzyme was significantly increased whereas the level of ACh decreased in brain rats of the AlCl3 group when compared to the normal group. These findings are in agreement with several studies, but also are in disagreement with another. These discrepancies between studies are explained by Peng et al. (Peng et al. 1992) who related them to the metal exposure duration, type of its salt and the difference in response of brain regions to aluminum toxicity. Aluminum might have caused lesions in the important regions of the brain like substantia nigra, cerebrum and hippocampus whereas little change was observed in other brain areas.

Dopamine neurotransmitter level was also measured and the rats of the AlCl3 group had significantly lower levels of dopamine in the brain when compared to the control group. Oxidative stress and hydrogen peroxide (H2O2) attenuates dopamine release and increases the loss of dopaminergic cells (Spanos et al. 2013). These previous observations formulate an attractive strategy to prevent or delay the progression of neurodegenerative diseases by controlling oxidative stress and the inflammatory response and this strategy is supported by Kato et al. (A Kato et al. 2013). Bowdler et al. (1979) (Bowdler et al. 1979) mentioned the fact that the serotonin has the important function in coordination between autonomic and neuroendocrine systems to control sensor- motor output, and this function explained the impairment of visuo-motor coordination performance by neural toxic inhibitory effect of aluminum on serotonin system. This fact may be partly explained the changes in psychomotor behavior tests in the present study such as the impairment in motor coordination in rotarod test and anxiety behavior in open field test in AlCl3 group. There is direct relationship between serotonin and dopamine and when the level of serotonin increased led to an increased in level of dopamine and vice versa (Yadid et al. 1994).

All neurotoxicity effects of AlCl3 were improved by TQ and celastrol in this study. TQ is present in Nigella sativa and has phenolic major bioactive components and potent antioxidant, Scavenge free radicals and anti-inflammatory activities against drug and chemicals induced multiple organ
toxicities (Ahmad et al. 2013). TQ improved the psychomotor performance, lipid peroxidation and CAT level in this study. TQ improved the muscle coordination and spontaneous locomotor activity, decreased AChE activity and the level of lipid peroxidation, and increased levels of antioxidant enzymes i.e., reduced glutathione, CAT, in a rat model pretreated with chlorpromazine toxicity (Safhi 2016). In accordance with our results some findings showed a significant decrease in IL-6, and TNF-α in brain rats of TQ group and attenuates depression-like behavior. These findings are also in agreement with previously obtained results (Kassab et al. 2017) who observed that TQ decreased the levels of MDA, TNF-α and increased the levels of CAT, in the cerebral cortex, cerebellum, and brain stem.

In the current study, TQ exerted significantly anticholinesterase activity and improved ACh level, and this finding is in line with Abulfadl et al. (2018) (Abulfadl et al. 2018). TQ significantly increased TAC and decreased MD, TNF-α level. These results indicate that TQ holds as a neuroprotector for the treatment of neurodegenerative disorders (Abulfadl et al. 2018). On the previous data, TQ may be a promising approach for the prevention and therapy of neurodegenerative disorders due to AlCl3 toxicity by modulating the levels of neurotransmitters and reducing inflammatory- oxidative stress and improving the psychomotor behavior that all impaired by AlCl3 toxicity in the present study.

Celastrol is a traditional Chinese herbal with potent anti-inflammatory and anti-oxidative effects and has beneficial effects in the treatment of neurodegenerative, and inflammatory diseases (Bian et al. 2016). Celastrol treatment significantly overcome the depletion in dopamine, attenuated the concentration of neuro-inflammation and improved the motor deficits. Celastrol also corrected the impairments in horizontal movements and vertical activity in electronic maze that were analyzed by Osaka University Computerized system and improved psychomotor coordination was evaluated in a rotarod apparatus of Female Sprague–Dawley rat model (Allison et al. 2000). Celastrol corrected the loco- motor behavior and decreased immobility time in the forced swimming test and tail suspension test in mouse model and increased allo-grooming with inhibition of pro-inflammatory (TNF-α, IL-6) in mice model (Bove et al. 2020, Zhu et al. 2021). Furthermore, Celastrol was found to be effective in another animal model of depression, such as the winter depression-like in fish by counterbalancing the increase in cortical ROS production, lipid peroxidation and inflammation (Nakayama et al. 2020). These previous benefits of Celastrol because of its phenol and carboxyl groups that scavenge free radicals and inhibit the peroxidation of the outer and inner mitochondrial membranes (Chen et al. 2014).

**Conclusions**

It is concluded that AlCl3 may be attributed to oxidative–inflammatory stress, disturbances of psycho – motor behaviors and some neurotransmitter levels. The results suggested that the TQ and celastrol were able to antagonize AlCl3 neurotoxicity at least partly, by their potent antioxidant and anti-inflammatory properties as well as by neurotransmitter modulation. Hence, TQ and celastrol are promising agents for investigating their preventive effect in animal models of neurodegenerative diseases.
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Conceptualization, Faten Abbas and Dalia Ibrahim; Data curation, Nadia Abozied, Amal Nabil and Shereen Mahmoud; Formal analysis, Faten Abbas and Dalia Ibrahim; Investigation, Faten Abbas and Dalia Ibrahim; Methodology, Faten Abbas and Dalia Ibrahim; Resources, Faten Abbas, Mohamed Ahmed Eladl, Mohamed El-Sherbiny, Nadia Abozied, Amal Nabil, Shereen Mahmoud, Sawsan Zaitone and Dalia Ibrahim; Software, Mohamed Ahmed Eladl and Sawsan Zaitone; Supervision, Faten Abbas; Validation, Mohamed El-Sherbiny; Visualization, Mohamed Ahmed Eladl, Mohamed El-Sherbiny, Nadia Abozied, Amal Nabil, Shereen Mahmoud, Sawsan Zaitone and Dalia Ibrahim; Writing – original draft, Faten Abbas, Nadia Abozied, Amal Nabil, Shereen Mahmoud and Dalia Ibrahim; Writing – review & editing, Mohamed Ahmed Eladl, Mohamed El-Sherbiny and Sawsan Zaitone.

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Fig 1. Motor activity of rats in the open field test. A) Horizontal ambulation and B) vertical ambulation. Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at $P<0.05$. $^{a,b}$ Represents a statistically significant difference when compared to control and AlCl$_3$, respectively.
Fig 2. Number of fallings in the rotarod test. **Results** were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05. a, b and c Represents a statistically significant difference when compared to control, AlCl3 and AlCl3+TQ, respectively.
Fig 3. Anxiety-like behavioral parameters in the open field test. A) Time spent in the central zone & B) Grooming frequency/5 min. Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05. a, b and c Represents a statistically significant difference when compared to control, AlCl₃ and AlCl₃+TQ, respectively.
Fig 4. Depressive-like behaviors in the rat forced swimming test. Results were expressed as mean ± SD and analyzed using one-way ANOVA followed by Bonferroni’s post-hoc test at P<0.05. a, b represents a statistically significant difference when compared to control and AlCl3, respectively.