Biologically Active Compounds Present in Tobacco Smoke: Potential Interactions Between Smoking and Mental Health

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Tobacco dependence remains one of the major preventable causes of premature morbidity and mortality worldwide. There are well over 8,000 compounds present in tobacco and tobacco smoke, but we do not know what effect, if any, many of them have on smokers. Major interest has been on nicotine, as well as on toxic and carcinogenic effects and several major and minor components of tobacco smoke responsible for the negative health effects of smoking have been elucidated. Smokers themselves report a variety of positive effects from smoking, including effects on depression, anxiety and mental acuity. Smoking has also been shown to have protective effects in Parkinson's Disease. Are the subjective reports of a positive effect of smoking due to nicotine, of some other components of tobacco smoke, or are they a manifestation of the relief from nicotine withdrawal symptoms that smoking provides? This mini-review summarises what is currently known about the components of tobacco smoke with potential to have positive effects on smokers.

Keywords: tobacco dependence, nicotine, tobacco smoke components, monoamine oxidase inhibition, mental health, Parkinson's Disease

INTRODUCTION

Smoking is a major cause of preventable premature death and disability, believed to cause six million deaths worldwide each year with smokers, on average, losing 10 years of their lives (West, 2017). The negative health effects of prolonged smoking are well established. Smoking primarily damages lung and cardiovascular health, as well as impacting negatively on every organ of the body (U.S. Department of Health and Human Services., 2004). The reason people keep smoking, despite knowing that this habit is likely to eventually kill them, is that smoking is addictive (Henningfield and Fant, 1999). The major addictive component of tobacco smoke is nicotine, but it is increasingly evident that tobacco dependence is multi-faceted (West and Cox, 2021). Nicotine in laboratory tests is much less addictive than the lived experience of smokers would suggest (Jain, 2003; Balfour, 2009). The explanations are varied, ranging from societal and behavioural influences (Moolchan et al., 2003), to strong cue...
association and cognitive effects (Sacco et al., 2004; Chiamulera, 2005) and to the influence of tobacco companies (Hoek et al., 2012).

Smokers themselves report smoking relieves stress and anxiety, and aids concentration (Benowitz, 2010) and smokers with a variety of mental health conditions report using smoking as a form of self-medicaiton (Leonard et al., 2001; Aubin et al., 2012). Unsurprisingly, the interpretation of these reports varies widely. However, smoking is proven to have protective effects in Parkinson's Disease (Castagnoli and Murugesan, 2004; Gigante et al., 2017), with smoking having a well-established neuroprotective effect against this disease (Veljkovic et al., 2018).

A strong theme coming through the literature is that components in tobacco smoke other than nicotine, possibly monoamine oxidase (MAO) inhibitors, may enhance tobacco dependence, or may have positive effects on mood (Fowler et al., 2003; Rose, 2006; Hogg, 2016; Harris et al., 2020). Thus, apart from the effects of nicotine, the observed difficulty that people have in stopping smoking could be influenced by other chemical components of the tobacco smoke. Such interpretations are controversial. For instance, the high rate of relapse in smokers attempting to stop smoking is attributed to relief from nicotine withdrawal, rather than thinking of a positive effect on mood or concentration from smoking as reinforcing nicotine dependence (Moylan et al., 2013).

It is completely accepted that tobacco smoking is a harmful habit, due to the many toxic and carcinogenic components of the smoke. However, this review aims to focus attention on areas of the literature suggesting pharmacological drivers behind tobacco dependence other than the immediate effects of nicotine in inducing dependence, and our current understanding of the short-term effects of tobacco smoke components on smokers. Are some of these effects positive, as in neuroprotection against Parkinson's Disease? Are there known tobacco components which could have a positive effect?

NICOTINE

Nicotine acts at the nicotinic acetylcholine receptors (nAChRs) to cause flow-on effects in specific areas of the brain. These nAChRs are believed to be important in coordinating brain responses, by action of the natural transmitter, acetylcholine (ACh). Nicotine binds to nAChRs mimicking the action of ACh and altering responses within the brain (Balfour, 2009).

The key addictive response is stimulation of nAChRs in the ventral tegmental area of the brain, which causes the release of dopamine in the nucleus accumbens, believed to be central to the development of all addictive responses (Di Chiara and Imperato, 1988). Nicotine's pharmacology has been well reviewed by others (Benowitz, 1996; Laviolette and van der Kooy, 2004) as have the complexities of the nicotinic receptors in the brain (Albuquerque et al., 2009; Wu and Lukas, 2011). The nAChRs belong to the ionotropic family of receptors, with five proteins forming a channel. Seventeen different proteins have been identified, leading to a wide diversity of nicotinic receptor subtypes. Thus the diversity of nicotinic receptor types and their varied localisation within the brain allows for much more nuanced and complex responses to nicotine than simple dopamine release.

Several milligrams of nicotine are present in each gram of tobacco, and nicotine reaches around 0.2 micromolar concentrations in the blood of smokers, sufficient to cause nAChR responses (Alkondon et al., 2000). Nicotine concentrations rise rapidly when tobacco smoke is inhaled, reaching the brain in under two minutes. It then dissipates slowly, having a half-life of around 2 h (Benowitz, 2009). As nicotine brain concentrations fall, in the addicted smoker, cravings for nicotine begin, leading to smokers repeating the experience. Many of the effects of smoking tobacco (dopamine release and dependence, and withdrawal effects and relapse back to smoking) can be related back to these key effects of nicotine on the brain.

As the major pharmacologically active component of tobacco smoke, nicotine has also been investigated to see whether it can cause other effects, reported from smoking, such as relief from anxiety and depression, improved concentration and symptom control in Schizophrenia, Parkinson's Disease, Attention Deficit Hyperactivity Disorder, and Alzheimer's Disease (Mihaiescu and Drucker-Colin, 2000; Newhouse et al., 2004a; Veljkovic et al., 2018).

Nicotine and Cognition/Concentration

In adults, however, nicotine is believed to have a positive effect on mental acuity (Conley et al., 2021; Nop et al., 2021). Again, this has been suggested by studies in animals and in humans (Kumari et al., 2003; Newhouse et al., 2004b), and trials of the effect of nicotine in older adults have produced some evidence of benefit. A recent meta-analysis by Majdi and coworkers suggested that nicotine has a moderate but positive effect on attentional ability in healthy non-smoking adults (Majdi et al., 2021) and it has been suggested that nicotine could be used to treat late-life depression, with part of this action being mediated by effects on cognition (Gandelman et al., 2018).

Smoking as a long-term enhancer of cognition is not recommended, however, since smoking is a risk factor for vascular dementia (Lopez-Arrieta et al., 2001) as well as for many other health conditions.

Nicotine and Anxiety/Depression

The suggestion that nicotine can relieve stress and help with anxiety and depression is controversial. The key argument is whether the positive effects noted by smokers are real and act as a reinforcer of tobacco dependence by improving mood (Pomerleau et al., 1984; Choi et al., 2015) or whether nicotine dependence causes depression and/or anxiety over time, with...
relief of cravings being misread by smokers as relief from the related mood disorders (Moylan et al., 2013; Molas et al., 2017). A systematic review by Fluharty and co-workers (Fluharty et al., 2017) found evidence for causation in both directions and suggested the need for more studies. More recent work tends to look at effects of transdermal nicotine, and has found potential for nicotine to be used in cases of major depressive disorder (Janes et al., 2018), and for relief of later life depression (Conley et al., 2021).

**Nicotine and Schizophrenia**

Schizophrenia is also strongly associated with tobacco smoking, however, the direction of causation is controversial. Scott and coworkers examined the evidence that smoking might cause schizophrenia and suggested nicotine as the causative agent (Scott et al., 2018). Others have suggested that nicotine's effects within the brain might give relief from the symptoms of schizophrenia, providing motivation to continue smoking (Postma et al., 2006). It is possible that nicotine's positive effect on cognition is the mediating mechanism for this (Waterhouse et al., 2018).

**Nicotine and Alzheimer's Disease**

The possibility that nicotine might be useful in treating early stages of Alzheimer's Disease has been studied for some years. As with Parkinson's Disease, smokers are under-represented among those with Alzheimer's Disease. Impaired cholinergic function is an early feature of Alzheimer's Disease and nAChR agonists have been suggested as potentially helpful in ameliorating symptoms (Albuquerque et al., 2001). Transdermal nicotine has been used in trials with positive effects on attention and other cognitive measures (Newhouse et al., 2004b). Nicotine has also been shown to be effective in reducing behavioural and synaptic plasticity deficits in a rat model of Alzheimer's disease (Estoves et al., 2017). In a separate line of enquiry nicotine is also thought to delay formation of amyloid plaque (Zhang et al., 2006), giving another mechanism by which smoking (nicotine) could be helpful in delaying Alzheimer's Disease onset.

**Nicotine and Parkinson's Disease**

While smoking is well known to be protective against Parkinson’s Disease (Gigante et al., 2017), the causative agents in this case are likely to include both nicotine, though its action on dopaminergic pathways (Thiriez et al., 2011), and monoamine oxidase inhibitors (Castagnoli and Murugesan, 2004). Additionally, Kardani et al. (2017) have demonstrated that nicotine can slow the formation of α-synuclein fibrils. However, although nicotine appeared to be neuroprotective in Parkinsonian animals it had no effect in restoring damage in the same experimental system (Huang et al., 2009) and was not significantly effective in Phase II clinical trials (Villafane et al., 2018).

**MINOR TOBACCO ALKALOIDS**

Like nicotine, the other nicotine analogues found in tobacco smoke have also been found to act on the nicotinic receptors. The “minor alkaloids” in tobacco smoke other than nicotine include nornicotine, myosmine, cotinine, anabasine and anatabine. Where nicotine is found at around 10 milligrams per cigarette in tobacco, these minor alkaloids are found in microgram per cigarette amounts (Smith et al., 2015).

Several groups have looked at the effect of these alkaloids in vivo (Harris et al., 2015; Marusich et al., 2017; Tan et al., 2022). Such studies have found that the minor tobacco alkaloids can partially substitute for nicotine in behavioural tests but have lower potency than nicotine itself. The different alkaloids are not equivalent in their behavioural effects. Differences in binding to different nAChR variants between the various nicotinic alkaloids would allow for differences in mode of action and potency even if all act through binding nAChRs. The combination of lower potency and lower concentration in tobacco make it unlikely that these alkaloids have significant effects on smoker behaviour.

Cotinine, a breakdown product of nicotine, has relatively long half-life in the blood [c.a. 16 h compared to c.a. 2 h for nicotine (Hukkanen et al., 2005)] and reaches low micromolar concentrations in a smoker's blood, several-fold higher than the concentration of nicotine. It may, therefore, be present in sufficient amounts to have a modulating effect on smoker behaviour, particularly for heavy smokers. Cotinine is a weak agonist of nAChRs, binding less strongly than nicotine (Tan et al., 2021) but has recently been shown to support self-administrative behaviour (Tan et al., 2022) although less robustly than nicotine. Cotinine upregulates the α7nAChRs and this activity is neuroprotective to glial cells (Jarkov et al., 2021). Cotinine also slows down the formation of α-synuclein fibrils, with a potency similar to that of nicotine, and may also have positive cognitive benefits.

**MONOAMINE OXIDASE INHIBITORS**

Monoamine oxidase inhibitors (MAOIs) in tobacco smoke have long been regarded as having potential significance in modulating the effects of smoking on the brain (Fowler et al., 2003; Dome et al., 2010; Hogg, 2016). Monoamine oxidase (MAO) enzymes in the brain are responsible for clearance of brain transmitters, notably dopamine, serotonin, adrenaline and noradrenaline (Lewis et al., 2007). Since inhibition of MAO activity will lead to reduced clearance of neurotransmitters such as dopamine it is suggested that MAOIs in the tobacco smoke might enhance the dopamine reward from nicotine and the addictiveness of smoking.

A variety of experimental evidence has been produced showing that MAO inhibition enhances behavioural responses to nicotine in rats (Guillen et al., 2005; Villegier et al., 2006, 2011; Smith et al., 2016b; Harris et al., 2020) but the extension to human smoking behaviour is less clear. MAO activity is well known to be reduced in smokers (Fowler et al., 1996a,b) with the inhibition being believed to be irreversible (Yu and Boulton, 1987). The time course of recovery of activity after smoking cessation extends...
TABLE 1 | Nicotine, cotinine, and monoamine oxidase inhibitors in tobacco and tobacco smoke: potential positive effects.

| Smoke component | Chemical structure | Mechanism | Disease state affected | References |
|-----------------|------------------|-----------|------------------------|------------|
| Nicotine        | ![Nicotine](image1) | nAChR activation | Cognitive improvement | Newhouse et al., 2004b; Majdi et al., 2021 |
| Nicotine        | ![Nicotine](image2) | ↓ α-synuclein fibril formation: (cognition ↑) | Parkinson’s disease | Kardani et al., 2017; Villafane et al., 2018 |
| Nicotine        | ![Nicotine](image3) | ↓ Amyloid β-peptide aggregation: (cognition↑) | Alzheimer’s disease | Zhang et al., 2006; Esteves et al., 2017 |
| Nicotine        | ![Nicotine](image4) | Cognition↑ | Schizophrenia | Waterhouse et al., 2018 |
| Nicotine        | ![Nicotine](image5) | Cognition↑ | Depression | Gandelman et al., 2018; Conley et al., 2021 |
| Cotinine        | ![Cotinine](image6) | ↓ α-synuclein fibril formation, neuroprotection | Parkinson’s disease, Alzheimer’s disease | Terry et al., 2005; Riveles et al., 2008; Iarkov et al., 2021 |
| Naphthoquinones | ![Naphthoquinones](image7) | MAO inhibition, neuroprotection (nitric oxide control) | Parkinson’s disease | Venkatakrishnan et al., 2009 |
| Harman/Norharman | ![Harman/Norharman](image8) | MAO inhibition | Antidepressant | Farzin and Mansouri, 2006 |
| Harman/Norharman | ![Harman/Norharman](image9) | MAO inhibition | Antianxiety | Smith et al., 2013 |
| 2,3,6-Trimethyl-1,4-naphthoquinone | ![2,3,6-Trimethyl-1,4-naphthoquinone](image10) | MAO inhibition | Neuroprotection, Parkinson’s disease | Sari and Khalil, 2015 |

over several weeks after smoking cessation, consistent with the equivalent recovery time from inhibition by known irreversible MAO inhibitory drugs (Fowler et al., 2003). Although epigenetic (Launay et al., 2009) and microRNA (Higuchi et al., 2018) mechanisms for MAO activity reduction have been suggested, direct inhibition by components of tobacco smoke as the major cause of the observed reduction remains a likely mechanism by which MAO inhibition is accomplished in smokers.

The known MAOIs in tobacco smoke include the β-carbolines harman and norharman, α-naphthylamine, farnesyl acetone and tetrahydroisoquinolines (TIQ’s) (Lewis et al., 2007; Hogg, 2016). Harman and norharman have been proposed as the major contributors to the observed MAOI activity in tobacco smoke, causing the observed decrease in MAO activity (Rommelspacher et al., 2002; Herraiz and Chaparro, 2005), with the discrepancy between the amounts of β-carboline measured in smokers, and the inhibition observed being ascribed to their accumulation in platelets (Rommelspacher et al., 2002) and in brain (Fekkes and Bode, 1993). However, harman and norharman make up less than 1/10th of the total direct MAO inhibitory activity in tobacco smoke (Truman et al., 2017) so the opportunity for other MAOIs in tobacco smoke to contribute substantially to the MAO activity reduction seen in smokers must exist. No irreversible MAO inhibitors have yet been reported from tobacco or tobacco smoke.

The question of whether MAO inhibitors in tobacco smoke can affect behaviour remains unresolved. As well as the potential for effects on addiction, MAO enzymes are drug targets for a variety of neurological disorders including depression, mood, anxiety, attention deficit hyperactivity, Tourette’s syndrome, Parkinson’s disease and Alzheimer’s disease (Sharma, 2016; Borroni et al., 2017). Of the known MAO inhibitors in tobacco smoke, high concentrations of harman and norharman can affect responses to nicotine (Harris et al., 2020) and may act as antidepressants (Farzin and Mansouri, 2006; Smith et al., 2013) in animals. However, when used in amounts relevant to smokers, they were not seen to affect rat self-administration of nicotine (Smith et al., 2015) and no pharmacological effects have been reported from the known tobacco MAO inhibitors at physiologically relevant concentrations. In contrast, use of tobacco smoke extracts in self-administration or intracranial self-stimulation experiments has been found to affect responses to nicotine (Harris et al., 2010; Costello et al., 2014; Brennan et al., 2015). This discrepancy may be in part because the full range of tobacco smoke MAO inhibitors has not yet been identified.

It has also been suggested that further MAO inhibitory activity is formed from smoke components in the body. Acetaldehyde is formed during tobacco combustion, from the sugars in the plant material, and from sugars added as humectants and flavour additives during tobacco and cigarette manufacture. Acetaldehyde is typically present in cigarette smoke amounts ranging from 0.6 to over 2 milligrams per cigarette (Seeman et al., 2002). Acetaldehyde yields locomotor stimulation and reinforcing effects (Queremont and Tambour, 2004) at concentrations higher than those seen in tobacco smoke and enhances self-administration of nicotine at concentrations similar to those in tobacco smoke (Belluzzi et al., 2005), although this finding was not confirmed by Smith and co-workers in similar trials (Smith et al., 2015).
It is suggested that acetaldehyde acts by reacting with chemicals naturally occurring in the brain to form MAO inhibitors (Talhout et al., 2007). A wide variety of TIQs are formed from acetaldehyde and catecholamines (dopamine, noradrenalin, adrenalin) (Naoi et al., 2004; Patsenka and Antkiewicz-Michaluk, 2004). Of particular note are the cyan derivatives, which inhibit MAO-A and -B with Ki values between 18 and 38 µM (Mendez-Alvarez et al., 1997). A group of tetrahydro-β-carbolines (THBCs) are also formed from condensation of acetaldehyde and indoleamines (serotonin, tryptamine, tryptophan) (Talhout et al., 2007) and are more potent, inhibiting with Ki values under 10 µM. It seems likely that concentrations of these MAOIs, formed in the body after smoking, are sufficient to have some effect on smoking addiction in humans, as well as in rats, since a randomized double-blind trial of a method to reduce the amount of acetaldehyde entering a smoker's body had some success in encouraging cessation (Syrijanen et al., 2017). Thus far, together with the β-carbolines, acetaldehyde is a leading candidate as a tobacco smoke component causing monoamine oxidase inhibition in smokers sufficient to modulate tobacco dependence, even though it does not directly cause MAO inhibition. The inhibitors formed from acetaldehyde appear to be reversible (Naoi et al., 2004).

The immediate effect on MAO activity of smoking a single cigarette could not be detected using PET methods (Fowler et al., 2003), whereas longer term effects of continued smoking results in an overall, apparently irreversible reduction of 30–40% in both MAO-A and -B activity, suggesting that long term exposure to tobacco smoke is required for this effect. Until we know which components of tobacco smoke cause the observed inhibition of MAO activity, and their mechanism of action, it will remain difficult to determine the extent, timing and overall effects of MAO inhibition in smokers.

Monoamine oxidase inhibitors in tobacco smoke may well have effects other than enhancing the addictive potential of nicotine. Smith and coworkers (Smith et al., 2016a) have examined the effects of MAO inhibitors and nicotine on brain function, measured using EEG techniques in non-smoking humans. They suggest that MAO inhibition alters brain function leading to lapses in cognition. Nicotine's effect in enhancing cognition is suggested to alleviate this.

Monoamine oxidase inhibitors are a major drug target, of interest for treatment of depression and anxiety, Parkinson's disease and Alzheimer's disease (Sharama, 2016). It has been suggested that the MAOI activity in tobacco smoke may enhance a smoker's mood (Pomerleau et al., 1984; Farzin and Mansouri, 2006; Smith et al., 2013; Choi et al., 2015) independently of their effects on dopamine reward from nicotine, and may contribute to smoker's dependence on tobacco by this indirect route (Arnold et al., 2014). However, rather than a positive effect on mood, it is possible that MAO inhibition serves to intensify withdrawal symptoms (Malin et al., 2013). Relief of withdrawal symptoms then serves to improve mood. This potential interplay of influences is not yet fully understood.

The potential role for MAO inhibition in tobacco dependence has led to suggestions that MAO inhibitors could be used for smoking cessation (Biberman et al., 2003; George and Weinberger, 2008). Current consensus is that, after some promising preliminary results, the known MAO inhibitory drugs are not particularly useful for smoking cessation (Howes et al., 2020).

The proposed effects of MAO inhibitors on smokers are listed in Table 1.

### OTHER COMPONENTS OF TOBACCO SMOKE

Additional components of tobacco and tobacco smoke which have been associated with beneficial effects are listed in Table 2, below. They have been identified as a result of investigations of the active components supporting herbal use, but from plants other than tobacco. These compounds are all known components of tobacco (Rodgman and Perfetti, 2013), however, while their biological activity is of interest, their contribution to the overall effects of tobacco smoking is unknown.

### CONCLUSION

It is accepted that nicotine has positive effects on cognition and attention in adults, even though having negative effects in infant and child development. For this reason, nicotine could be helpful in the management of a variety of mental health conditions and provide an explanation for high levels of smoking in disorders such as schizophrenia. Nicotine and its metabolite cotinine both stimulate cholinergic systems, which can be neuroprotective, and may also prevent β-amyloid fibre aggregation in Alzheimer's disease, and α-synuclein fibre formation in Parkinson's disease. While the long-term overall effect of smoking is deleterious, the same may not be true for nicotine and cotinine. The therapeutic use of cotinine deserves further work, since it is expected to have a lower abuse potential than nicotine but may provide similar neuroprotective and cognitive effects.

The other most significant and potentially beneficial biological activity in tobacco smoke is monoamine oxidase inhibitory activity. This is likely to have an impact on the response to nicotine, and to increase the addictiveness of smoking, both by increasing the dopamine rewards from the nicotine, and by its effects (if any) on mood. The extent to which these effects on mood from smoking are due to relief of nicotine cravings, or directly caused by the modulation of monoamine oxidase enzyme activity in the brain remains uncertain. In this respect it is important to recall that the first generation of antidepressant drugs were selective MAO inhibitors, with some still currently used for treatment of major depressive disorders. This question will only be resolved when the causative agents of the MAO inhibition seen in smokers have been identified, and their effects can be studied independently of the effects of nicotine. It may prove possible to separate immediate MAO inhibitory activity from the long-term irreversible inhibitory factors and assess
TABLE 2 | Additional tobacco smoke components with beneficial biological activity.

| Component     | Chemical structure | Observed effect                                      | Application               | References                        |
|---------------|--------------------|------------------------------------------------------|---------------------------|-----------------------------------|
| Catechol      | ↓ amyloid-β fibril formation | Alzheimer's disease                             | Huong et al., 2010       |
| Hydroquinone  | ↓ a-synuclein fibrillation           | Parkinson's disease                             | Hong et al., 2009         |
| Solanesol     | Neuroprotection                  | Stroke                                             | Rajdev et al., 2020       |
| Limonene      | Neuroprotection                  | Alzheimer's disease                             | Shin et al., 2020         |
| Cambranoids   | Nicotinic activation, neuroprotection | Alzheimer's disease | Ferechmin et al., 2009 |
| Quercetin     | ↓ cognitive function, ↓ amyloid-β fibril formation | Parkinson's disease | Paula et al., 2019; Khan et al., 2019 |
| Kaempferol    | ↑ striatal dopamine, SOD and GSH  | Parkinson's disease                             | Li and Pu, 2011          |
| Eugenol       | ↓ immobility in forced swim test  | Depression                                         | Irie et al., 2004         |
| β-Asarone     | ↑ efficacy of memantine, ↑ cognitive deficit | Alzheimer's disease | Han et al., 2020; Liu et al., 2016; Chang and Teng, 2018 |
| Vanillin      | ↓ oxidative stress response, ↓ behavioural impairment | Parkinson's disease | Dhanalakshmi et al., 2016 |
| Ferulic acid  | ↑ serotonin and norepinephrine, ↓ immobility in forced swim test, ↓ brain capillary constriction | Depression, Alzheimer's disease | Chen et al., 2015; Wang et al., 2021 |
| Caffeic acid  | ↓ mitochondrial disfunction, ↓ oxidative stress | Depression                                          | Takeda et al., 2002       |
| Chlorogenic acid | ↓ immobility in forced swim test | Parkinson's disease                             | Singh et al., 2020        |
| Rutin         | ↓ immobility in tail suspension test, ↓ effect of chronic induced stress | Depression, Anxiety | Parashar et al., 2017; Yusha’u et al., 2017 |
| Naringenin    | ↓ amyloid-β toxicity, ↓ immobility in tail suspension test, ↑ memory | Alzheimer's disease, Depression | Yi et al., 2012; Yang et al., 2014; Md et al., 2018 |
| Naringin      | ↓ mitochondrial disfunction, ↑ memory         | Alzheimer's disease                             | Sachdeva et al., 2014     |
| Scopoletin    | ↓ anxiety-like behaviour                       | Anxiety                                            | Luo et al., 2020          |
| Esculetin     | ↓ immobility in forced swim test              | Depression, Anxiety                               | Sulakhiya et al., 2016    |
| Quinic acid   | ↓ MAO-B, neuroprotection                      | Dementia                                           | Liu et al., 2020          |

SOD, superoxide dismutase; GSH, glutathione peroxidase.

their effects separately once any irreversible inhibitors have been identified.

It is likely that MAO-B inhibitory activity in tobacco smoke, perhaps together with the combined neuroprotective effect of a variety of smoke components may be useful for symptom control in Parkinson's disease, while MAO-A inhibitory activity is more likely to contribute to the alleviation of mood disorders.

While the contribution of biologically active molecules other than nicotine remains to be established it is clear that tobacco smoke contains many components which might have beneficial effects. While this in no way compensates for the overall deleterious effects of smoking, these effects may help explain the strength of tobacco dependence many smokers experience and are worthy of further study, so that we can disentangle the deleterious and beneficial effects, to better help smokers to stop smoking.

AUTHOR CONTRIBUTIONS

PT, RP, and PT-S guided the initial literature search, performed by SH. SH and PT wrote the manuscript. BE added expert advice. All authors contributed to the final shape of the manuscript.

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