Depressed mood induction in early cigarette withdrawal is unaffected by acute monoamine precursor supplementation

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Background: Cigarette smoking is the leading preventable cause of death; however, quitting is difficult and early relapse is common. Dysphoric mood during early cigarette withdrawal is associated with relapse, and with the exception of bupropion and nortriptyline, few interventions have been developed to prevent this. During early cigarette withdrawal there is an elevation in the levels of monoamine oxidase-A (MAO-A), which removes monoamines excessively and induces oxidative stress and is implicated in creating sad mood. Hence, we conducted a randomized, placebo-controlled, double-blind crossover trial of a dietary supplement designed to counter the effects of elevated MAO-A levels on vulnerability to depressed mood.

Methods: Twenty-one otherwise healthy cigarette smokers completed the protocol, receiving either active dietary supplement followed by washout and placebo or the same in reverse order. The dietary supplement was composed of monoamine precursors (2 g tryptophan, 10 g tyrosine) and blueberry antioxidants (blueberry juice with blueberry extract). Vulnerability to depressed mood was measured by the change in scores of depressed mood on the visual analog scale (VAS) following the sad mood induction paradigm (MIP).

Results: There was a significant increase in VAS depressed mood scores after the sad MIP during supplement and placebo, but no difference between active and placebo conditions. There was also a significant increase in urge-to-smoke scores after sad MIP during supplement and placebo but no difference between active and placebo conditions. Reliability of the increase in VAS after MIP was very good.

Conclusion: The dietary supplement had negligible effect on depressed mood, but sad MIP is a very reliable method that can be applied in future studies to assess other interventions for preventing dysphoric mood during early cigarette withdrawal.

Keywords: cigarette withdrawal, dysphoria, dietary supplement, mood induction, monoamines

Introduction
WHO ranks cigarette smoking as the leading preventable cause of death worldwide.1,2 Quitting is difficult and ~50% of people relapse within the first 3 days of cigarette cessation.3,4 The symptoms of early cigarette withdrawal are associated with resumption of cigarette smoking.5,6 Among these symptoms, dysphoric mood is important because it starts as early as 4 hours into cigarette cessation and is individually, strongly predictive of relapse.5,9

Most interventions, evaluated for their prevention of dysphoric mood in early cigarette withdrawal, were studied in subjects with a history of major depressive disorder (MDD). An advantage of this approach is that history of MDD is associated with more severe dysphoria during withdrawal10 but a disadvantage is that there is a paucity of...
investigations that include cases without past MDD. A double-blind randomized placebo controlled trial of sertraline found no effect of treatment on depressed mood or abstinence rates 1 week after smoking cessation or abstinence rates but found significantly less irritability, anxiety, craving and restlessness in those receiving sertraline compared to placebo. Another double-blind randomized controlled trial of placebo, 20 mg, or 40 mg fluoxetine in addition to cognitive behavioral therapy (6 weeks) and nicotine patch (10 weeks) found that fluoxetine had a moderate effect on lowering withdrawal symptoms as measured by Minnesota Tobacco Withdrawal Symptoms Checklist in both groups as compared to placebo, but no targeted effect on mood and did not significantly improve smoking cessation rates. Nortriptyline was shown to reduce the negative effect over the first 8 days of smoking cessation, as measured by the profile-of-mood-state total mood disturbance score (POMS-TMD), and improve abstinence rates over 64 weeks, which was predicted by the level of depressed mood in early withdrawal. N-acetylcyesteine administration in a small pilot study was associated with higher positive effect compared to placebo during early smoking abstinence, though another study found no effect on withdrawal symptoms. One report of oral tryptophan with a high carbohydrate diet was associated with lower withdrawal symptoms in subjects undergoing behaviorally oriented smoking cessation treatment in early smoking withdrawal, but depressive symptoms were unaffected. Bupropion was shown in several studies to alleviate depressive and other withdrawal symptoms. However, even with bupropion treatment, subjects often experience depressed mood during early withdrawal with effect sizes of change towards depressed mood being ~0.5–1. Overall, most pharmacotherapy intervention studies targeting depressed mood during cigarette cessation evaluated antidepressants, with only bupropion demonstrating benefit in a moderate proportion of cases.

The effects of cigarette smoking are complex, and there are multiple mechanisms implicated in the occurrence of depressed mood during cigarette withdrawal. Ascending mesolimbic dopamine pathways from the ventral tegmental area (VTA) are modulated by several systems including cholinergic and glutaminergic projections from the tegmental pedunculopontine nucleus as well as GABA releasing neurons of the VTA. With chronic nicotine use, the modulation of these systems may be altered, shifting the influence of nicotine from GABA releasing neurons towards DA releasing neurons, facilitating increased activity of the mesolimbic DA pathway, and leading to psychological symptoms of craving and withdrawal. There is also longstanding evidence that hypercholinergic states may lead to depression, such as through increased acetylcholine levels from organophosphate poisoning in humans, and greater nicotinic acetylcholine receptor affinity in Flinders Sensitive Line rats. Also, nicotinic antagonists may reduce depressed mood.

A relatively recent biological mechanism also implicated in the dysphoric mood of early cigarette withdrawal is the elevation in monoamine oxidase A (MAO-A) levels in those who smoke heavily. The B-carbolines harman and norharman are found in cigarette smoke and bind with high affinity to MAO-A, inhibiting it both in the brain and periphery during active smoking. During early withdrawal, lower levels of B-carbolines are associated with a rise in available MAO-A binding sites. Moreover, the increase in MAO-A binding sites in the prefrontal and anterior cingulate cortex was correlated with a shift to depressed mood between active smoking and withdrawal. Elevated MAO-A levels are consistently associated with depressed mood across a number of illnesses and conditions, including major depressive episodes (MDE), prior to recurrence of MDE, and perimenopause. MAO-A level is highly correlated with MAO-A activity in the brain, and MAO-A metabolizes monoamines such as serotonin, norepinephrine and dopamine, which support the functioning of neural structures underlying a healthy mood state.

The aim of this randomized double blind placebo controlled cross-over study was to determine if a new dietary supplement combination designed to counter effects of greater MAO-A levels during early cigarette withdrawal reduces vulnerability to depressed mood compared to placebo. The dietary supplement was composed of tyrosine, tryptophan and natural blueberry extract with blueberry juice. Tyrosine is an amino acid precursor to norepinephrine and dopamine, and tryptophan is a precursor amino acid for serotonin. Manipulations of tryptophan and tyrosine levels in plasma through dietary challenges may affect release of serotonin, dopamine and, to a lesser extent norepinephrine in rodent brain and may affect depressed mood in humans. Blueberry juice and extract contains anthocyanins, which are dietary antioxidants and were added to counter pro-oxidant effects of elevated MAO-A. Most investigations of blueberry anthocyanins in rodents and pigs suggest that they cross the blood brain barrier. It was hypothesized that administration of the dietary supplement would generate increased resiliency against induction of depressed mood in early cigarette withdrawal. Induction of depressed mood was chosen as a key outcome variable because it controls and incorporates environmental influences, is associated with vulnerability towards mood disorders, and the methods are well
standardized. Some studies have also shown that the effects of sad mood induction on increased smoking motivation is heightened in depressed smokers. In this study, mood induction procedure (MIP) was used to induce vulnerability to sadness in subjects in early cigarette withdrawal.

Methods
Study design and study participants
Healthy volunteers (aged 18–65 years), who reported smoking at least a half a regular pack (20 cigarettes per pack) of cigarettes per day regularly, were recruited through advertisement. All had reported a past history of smoking a pack a day. Sixty-six percent of subjects had tried to quit cigarette smoking in the past. Inclusion criteria included: report of stable, good, current physical health, a normal physical examination, and ability to give informed consent. Exclusion criteria were recent history of myocardial infarction and/or stroke, any current substance use disorder except for nicotine, and current diagnosis of MDD (based on Hamilton Depression Rating Scale [HDRS] > 7). This study was approved by the Research Ethics Board for Human Subjects at the Center for Addiction and Mental Health (CAMH), Toronto, Canada and the Natural Health Product Directorate and Drug Directorate of Health Canada. All subjects provided written informed consent after a thorough explanation of the study. Participation was voluntary and each participant was free to withdrawn anytime during the study. All experiments on human subjects were conducted in accordance with the Declaration of Helsinki and the International Conference on Harmonization Good Clinical Practice guidelines. (ClinicalTrials.gov, Identifier: NCT02042521).

Participants came in for three visits, which included a screening session and 2 active study days, at least 1 week apart. During the screening visit the subjects completed a standardized medical history, a urine drug screen to rule out current substance abuse, the Fagerstrom Test for Nicotine Dependence, the Nicotine Dependence Syndrome Scale, and the HDRS. A record of the brand of cigarettes smoked, number of cigarettes smoked per day in the past and currently, and the number of years they had been smoking cigarettes was obtained. Also, exhaled CO levels were measured at screening and a cutoff of 10 ppm was applied (Bedfont Scientific Ltd., Kent, England). For women, a pregnancy test was also done during the first visit.

Main study procedure
After screening, subjects were randomized for receiving dietary supplement, then washout then placebo or placebo then washout then dietary supplement. Randomization was done by computer generated fixed block model performed by an independent company (National Sanitation Foundation-Guelph Food Technology Center [NSF-GFTC]) and the placebo matched the active dietary supplement in taste and color. On each active study day, subjects were asked to refrain from smoking cigarettes for 8 hours (Figure 1). Exhaled CO levels were also measured within ~4.5 hours of the first measurement on each active study day (at 7:30 AM and 12:15 PM). After 5 hours of withdrawal, subjects underwent a sad MIP (see complete description below). Mood symptoms and symptoms of cigarette withdrawal obtained during and after the MIP were compared between the 2 different days.

Dietary supplement
The dietary supplement consisted of a morning and an evening serving. The key ingredients of the active evening dietary supplement were 2 g of tryptophan, 280 mL of blueberry juice.
and 1 g of blueberry extract. The key ingredients of the active morning dietary supplement were 10 g of tyrosine totaling 10 g, 280 mL of blueberry juice and 1 g of blueberry extract. Tryptophan was given in the evening before an active study day and tyrosine was given on the morning of an active study day, with a separation of at least 8 hours to avoid competition at the Large Neutral Amino Acids Transporters for transport though the blood–brain–barrier. The l-tryptophan capsules of 1 mg were obtained from Apotex (Apo-tryptophan by Apotex Incorporated, Toronto, Canada, Drug Identification Number [DIN]# 02248539). Tyrosine capsules of 500 mg were obtained from Trophic Canada (Trophic Canada, Richmond Hill, Canada, NPN#80010004). The blueberry extract powder called VitaBlue, was obtained from Future-Ceuticals (Future Ceuticals, Momence, IL, USA) and prepared and packaged by NSF-GFTC, in sachets containing 1 g. Blueberry juice was obtained from Milne Fruit Products and packaged by GFTC in 280 mL bottles. The blueberry juice and extract were designed to be mixed at the time of ingestion. Blueberry extracts were added to the juice, since the anthocyanins in blueberry juice are mostly damaged during pasteurization. Placebo blueberry juice and extract were designed and prepared by NSF-GFTC and placebo tryptophan and tyrosine were prepared by the local CAMH pharmacy. The placebo pills appeared identical to tryptophan and tyrosine pills. The placebo drink and extract were a natural drink and powder that were blue due to natural coloring, had a blueberry flavor through natural flavoring, had negligible antioxidant properties and had sugar levels similar to that of to the active drink and extract.

**Measures**

Mood symptoms and symptoms of cigarette withdrawal were compared between the 2 different days and were measured before and after the sad MIP on each visit (Figure 1). For measuring mood, a 10-cm visual analog scale (VAS) was used. VAS uses a 10-point scale for participants to indicate the extent to which each of the items indicated is consistent with how they feel at that moment, ranging from most happy to most depressed. In addition, there were VAS scales labeled from left to right from most energy to least energy, and least anxious to most anxious. Other questionnaires included were the short form 37-item of POMS scale, and the urge to smoke scale (USS), an analog scale, with 10 craving-related questions.

**Assessment of vulnerability to depressed mood induction**

During each active study day, the participants underwent a neutral MIP and a sad MIP. The first neutral MIP was applied to reduce environmental effects on mood. The MIP applied the Velten method in combination with the approach of Clark. The Velten method is the most widely used technique for studying affective influences upon behavior and has demonstrated effectiveness in altering subjective emotional states. Velten MIP is a series of 60 self-referent statements. Negative statements reflect pessimism, dissatisfaction, and lethargy; for example “life is a heavy burden.” Neutral statements reflect a lack of mood effect; for example, “an orange is a citrus fruit.” Subjects were asked to read each statement, printed individually, first to themselves and then aloud, and to “feel and experience each statement as it would apply to you personally.” For the method of Clark, to facilitate the MIP, participants were also presented with a piece of music while reading the statements. For sad MIP, subjects listened to Prokofiev’s “Russia under the Mongolian Yoke” and for neutral MIP, subjects listened to Mozart’s “Piano Concerto No 21 in C Major.”

MIP is well known to induce a temporary depressed mood state in healthy humans as well as in recovered states from mood disorders. MIP can act as an advanced form of mood rating where the environmental conditions are standardized. To obtain a maximal percentage of responders to mood induction in this study, a combination of the Velten and the musical MIP was used in this study. The Velten and Clark methods are common in mood disorders field although this is their first application together to assess mood during early cigarette withdrawal.

**Monitoring for compliance**

Compliance monitoring of the evening supplement was done with riboflavin fluorescence. Along with all the active or placebo dietary supplements, a pill of riboflavin (100 mg), which was identical in appearance to the tryptophan capsules, was given to the subjects. The subjects were notified that riboflavin was included in the dietary supplements, but they were not told which pill contained riboflavin. The urine sample taken on the day of the MIP was subjected to ultraviolet light to check for riboflavin fluorescence to verify compliance. The morning supplement was taken in the presence of the research study personnel.

**Data analytic plan**

First, a within subject t-test was done to assess whether there was an effect of the sad MIP on VAS depressed mood scores in both the active and placebo conditions by comparing the VAS depressed mood scores pre-MIP (VAS 4) and post sad MIP (VAS 5). Then, the change in VAS depressed mood scores, immediately before and after the sad MIP, was...
calculated (ie, the change in VAS scores from VAS 4 to 5) and applied in several subsequent analyses as follows. A repeated-measures ANOVA (rmANOVA) (considering the repeated visit as the repeated measure) was performed to compare the effect of the dietary supplement vs placebo and assess the effect of the order (supplement given with the first or second MIP). In the event that the primary outcome was negative, that is, no effect of the supplement on change in VAS scores of depressed mood, it was decided that test-retest reliability between the VAS depressed mood change scores on the first depressed MIP visit and the second depressed MIP visit would be assessed with the intraclass correlation coefficient (ICC).

In addition, a within subject t-test was done to assess whether there was an effect of the sad MIP on USS scores in both the active and placebo conditions by comparing the USS scores pre MIP (USS 3) and post sad MIP (USS 4). Then, the change in USS scores, immediately before and after the sad MIP, was calculated and applied in several subsequent analyses as follows. A rmANOVA was performed to compare the effect of the dietary supplement vs placebo and assess the effect of the order (supplement given with the first or second MIP).

Results
The mean age of the samples was 43.73±12.28 years and on average, participants smoked 19.81±5.06 cigarettes per day. The demographics and clinical characteristics of the subjects are presented in Table 1. Riboflavin was detected positive through visual urine fluorescence technique in all subjects.

Effect of dietary supplement on depressed mood after MIP
There was a significant increase in VAS depressed mood scores (VAS4 to VAS5) after the sad MIP during supplement (mean ± SD: 1.56±2.09 cm, t[20]=3.43, P=0.003) and placebo (mean ± SD: 1.12±1.81 cm, t[20]=2.84, P=0.01). Visual inspection of the plot of depressed mood over time indicated that the increase in depressed mood occurred primarily between time point four (corresponding to 55 minutes prior to MIP and 3 hours and 50 minutes into cigarette withdrawal) and (corresponding to 10 minutes post MIP and 4 hours and 55 minutes into cigarette withdrawal), with a decline in depressed mood after time point 5 (see Figure S1). rmANOVA showed that the change in VAS scores of depressed mood in the placebo and active conditions did not differ significantly (F[1,19]=0.56, P=0.47) (Figure 2), nor was there an effect of the order on the change in VAS scores of depressed mood (F[1,19]=0.02, P=0.89). There was also no interaction between the order and supplement condition (rmANOVA: F[1,19]=0.80, P=0.38).

Similar results was also observed for POMS depression/dejection and USS scores. There was a significant increase in POMS depression/dejection scores (POMS3 to POMS4) after sad MIP during supplement (mean ± SD: 2.95±4.57, t[20]=2.95, P=0.008) and placebo (mean ± SD: 2.19±3.37, t[20]=2.98, P=0.007) conditions (Figure S3). rmANOVA showed that the change in POMS depression/dejection in the active and placebo conditions did not differ significantly, (F[1,19]=0.002, P=0.97), nor was there an effect of the order on the change in POMS depression scores (F[1,19]=3.26, P=0.09). However, there was a significant interaction between the order and effect of supplement, (rmANOVA: F[1,19]=5.82, P=0.02) which was attributable to a greater effect of the MIP on the first visit with placebo as compared to the second visit with placebo as found on a post-hoc independent t-test (t[19]=4.23, P<0.005).

Table 1 Demographic and clinical characteristics

| Measures                        | Mean ± SD or % |
|--------------------------------|----------------|
| Age (years)                    | 43.73±12.28    |
| Sex (male)                     | 38.02%         |
| Duration of smoking (years)    | 23.63±12.56    |
| Average number of cigarettes smoked per day | 19.81±5.06 |
| HAM-D scores                  | 1.4±2.14       |
| Exhaled carbon monoxide levels | 22.19±14.79    |
| Fagerstrom test for nicotine dependence | 4.85±2.03 |
| Nicotine dependence syndrome scale (overall score) | -0.48±0.65 |
| Riboflavin positive urine detection | 100%          |

Note: These measures were obtained at screening visit (visit 1). Abbreviation: HAM-D, Hamilton depression rating scale.
There was a significant increase in USS scores (US$3 to US$4) after sad MIP during supplement (mean ± SD: 0.63±0.83, t[20]=3.50, P<0.005) and placebo (mean ± SD: 0.45±0.68, t[20]=3.04, P=0.006). The differential change in USS between the active and placebo conditions was not significant, (rmANOVA: F[1,19]=0.72, P=0.41) (Figure 3), nor was there an effect of the order on MIP (rmANOVA: F[1,19]=0.58, P=0.45). There was also no interaction between the order and supplement conditions (rmANOVA: F[1,19]=0.25, P=0.62). Review of USS scores over the individual time points showed a steady increase in scores over time (see Figure S2).

In order to assess whether the difference in VAS from one assessment to the next was not similar, a difference score for VAS was calculated between each subsequent assessment. rmANOVA assessing the difference scores found that the difference scores were unequal (F[2,91,100]=5.84, P=0.002). A similar approach was taken for the USS to assess whether there were sequential differences in USS change. A difference score for USS was calculated between each assessment time-point. Visual inspection of the data did not suggest that the change in USS between assessments differed much over time, with somewhat greater change across time-points 2–3 and 3–4. rmANOVA assessing the difference, including a factor for visit, was a trend only (F[3,60]=2.592, P=0.061).

Reliability of depressed mood induction
The application of MIP on VAS depressed mood scores between time point four and five (pre and post sad MIP) of depressed mood revealed significant ICC across the repeated measurement (ICC =0.72, P=0.003). The rise in VAS scores of depressed mood on the first and second active study days were 47% and 32%, respectively. There was also a strong, positive correlation between the change in VAS mood scores on both active study days, which was statistically significant (r=0.57, N=21, P=0.007, R²=0.32) (Figure 4).

Discussion
To our knowledge, this is the first study to evaluate a dietary supplement consisting of tryptophan, tyrosine and blueberry juice with blueberry extract for reducing sad mood in early cigarette withdrawal and the first to apply the MIP in early cigarette withdrawal. Although there was no effect of the supplement on vulnerability to depressed mood, the MIP robustly and reliably induced depressed mood during early withdrawal.

The lack of effect of the dietary supplement combination in reducing sad mood is unlikely to be accounted for by the study methodology. Increasing the sample size with the present protocol would have a low probability of affecting the outcome; the change in depressed mood was 0.44 cm greater in the active condition (it was worse in the active condition). The 95% CI for the differential effect of the condition was −0.32 to 1.2 cm; hence the probability of a true benefit from the supplement in reducing the MIP effect by more than 0.32 cm was less than 2.5%. This is a low value in the clinical setting, representing only 3% of the range of the scale. For example, differential effects between groups in other studies with the MIP procedure include a greater effect of 3.5 cm in early postpartum compared to women not recently pregnant, and a 4.3 cm effect in favor of active supplement over placebo in early postpartum. The intake of the supplement was adequate since the compliance rate was high at 100% for the supplement consumption, and both tryptophan and tyrosine supplements at this dose increase plasma levels approximately five to sevenfold. Similar tryptophan and tyrosine administration at doses influencing plasma levels similarly are known to strongly elevate tryptophan and tyrosine in the cerebrospinal fluid, as well as, elevate metabolites of serotonin and dopamine in the cerebrospinal fluid. The amount of blueberry anthocyanins in the extract given corresponded to high-end dietary intake of blueberries and blueberries have a wide range of anthocyanins which tend to be brain penetrating. For example, Janle et al reported that gavage administered [14C]labeled anthocyanins are present in rodent brain 24 hours later at ~2% dose per gram. Although key issues of sample size, compliance and supplement composition appear addressed
in the study design, an issue that could be considered for future studies is whether longer term administration of the supplement prior to or after withdrawal would reduce depressed mood. This supplement was strikingly impactful in an open trial in postpartum blues, another condition associated with elevated MAO-A level, with an effect size of 2.9; but there are several reasons why this supplement may not necessarily reduce the impact of the MIP in early cigarette withdrawal. Plausible reasons are that the elevation of the index of MAO-A level in the prefrontal and anterior cingulate cortex of early cigarette withdrawal is of lesser magnitude compared to early postpartum (25% vs 43%), and it is likely that there is a broader range of mechanisms that account for depressed mood in withdrawal from cigarette smoking, given that there are over 5,000 chemicals in cigarette smoke. Also, the magnitude of depressed mood increase after MIP is considerably greater in early postpartum, so there is a larger target for intervention. Hence, the matching of the supplement to the elevation of MAO-A level and its associated vulnerability to MIP, may be inferior for early cigarette withdrawal as compared to early postpartum.

This study demonstrates that sad MIP done by the combination of Velten and music was successful in creating a temporary sad mood state, during early cigarette withdrawal, that was similar over two different testing sessions. There is still a need for a range of rapidly impacting options to prevent sad mood during early cigarette withdrawal. Hence, the sad MIP, an easy, economical measure, that detects vulnerability to sadness in early cigarette withdrawal, could be considered as a tool to assess the effects of other promising interventions for reducing sad mood during early withdrawal, although, given that the effect of MIP on the second visit was lower in magnitude than the first, it is likely that there is a limit for its repeated use in the same subjects (first visit: 1.58±2.01, second visit: 1.1±1.90).

In conclusion, our study does not support use of acute oral monoamine precursors with blueberry juice and extract to reduce depressed mood during early cigarette withdrawal. The study design rules out acute benefits of this combination on depressed mood, although more chronic dosing could be considered. A likely explanation for lack of effect is inadequate match of the supplement to the underlying condition.

Figure 4 Significant correlation between the change in VAS depressed mood scores on both active study visits (Pearson $r$=0.57, $P$=0.007, $R^2$=0.32). B (degree of slope) in regression is 0.54.
Abbreviation: VAS, visual analog scale.
heterogeneous set of mechanisms implicated in the depressed mood induced by cigarette withdrawal. Nevertheless, MIP for depressed mood is a simple and practical method that may be straightforwardly applied to investigate interventions for preventing vulnerability to depressed mood during early cigarette withdrawal.

**Author contributions**

All authors contributed to data analysis, drafting and revising the article, gave final approval of the version to be published, and agree to be accountable for all aspects of the work.

**Disclosure**

Dr Yekta Dowlati is developing natural health products to overcome high MAO-A state in early postpartum.

Dr Jeffrey H Meyer has been a consultant to Mylan, Lundbeck, Takeda, Teva, and Trius for the past 5 years. He is an inventor with five patents (processed or filed) including blood markers to predict brain inflammation or to diagnose affective disorders, and a dietary supplement to reduce depressed mood post-partum. None of these companies participated in the funding, design or execution of this study or writing the manuscript. Dr Meyer is developing natural health products to treat high risk states for MDE and is the inventor on patent applications for this dietary supplement for preventing depressed mood states.

Dr Peter Selby reports grants from Pfizer Inc./Pfizer Canada Inc., Shoppers Drug Mart, Bhasin Consulting Fund, and Patient-Centered Outcomes Research Institute; has received honoraria for speaking engagements from Pfizer Canada Inc., ABBVie, and Bristol-Myers Squibb and has also received consulting fees from Pfizer Inc./Pfizer Canada Inc., Evidera Inc., Johnson & Johnson Group of Companies, Medcan Clinic, Miller Medical Communications, MedPlan Communications, NVision Insight Group, Inflexion Inc., V-CC Systems Inc., Kataka Medical Communications, Sun Life Financial, and Myelin & Associates. Through an open tender process, Johnson & Johnson, Novartis, and Pfizer Inc. are vendors of record for providing smoking cessation pharmacotherapy, free/discounted, for research studies. All the other authors report no conflicts of interest in this work.

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**Supplementary materials**

![Visual analog scale (VAS) mood scores](image1)

**Figure S1** Visual analog scale (VAS) mood scores in subjects receiving active dietary supplement and placebo. The sad mood induction procedure was done between time points 4 and 5. (Time points correspond to: 1=7:35 am, 2=8:40 am, 3=10:10 am, 4=11:20 am, 5=12.25 pm, 6=13:10 am, 7=14:30) (Data shown as mean and standard error).

![Urge to smoke scale (USS) scores](image2)

**Figure S2** Urge to smoke scale (USS) scores in subjects receiving active dietary supplement and placebo. The sad mood induction procedure was done between time points 3 and 4. (Time points correspond to: 1=7:35 am, 2=8:40 am, 3=11:20 am, 4=12.25 pm, 5=14:30 pm) (Data shown as mean and standard error).

![Profile-of-mood-state (POMS) scale depression/dejection score](image3)

**Figure S3** Significant increase in profile-of-mood-state (POMS) scale depression/dejection score after sad mood induction procedure (MIP) during supplement (mean ± SD: 2.95±4.37, \(t[20]=2.95, P=0.008\)) and placebo (mean ± SD: 2.19±3.37, \(t[20]=2.98, P=0.007\)), but no effect of dietary supplement in reducing vulnerability to sad MIP compared to placebo conditions (\(F[1,19]=0.002, P=0.97\)).
