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Smoking cessation in individuals who use vaping as compared to traditional nicotine replacement therapies; a systematic review and meta-analysis

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Smoking cessation in individuals who use vaping as compared to traditional nicotine replacement therapies; a systematic review and meta-analysis

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Contributors statements:

Dr Pound conceptualized and designed the study, carried out the analyses, interpreted the data, drafted the initial manuscript, reviewed, and revised the manuscript.

Mrs Zhang participated in the conceptualization and design of the study, carried out the analyses, interpreted the data, participated in drafting the initial manuscript, reviewed and revised the manuscript.

Ms Kodua participated in the conceptualization and design of the study, and reviewed the manuscript.

Dr Sampson participated in the design of the study, developed the search strategies, reviewed, and revised the manuscript.

All authors approved the final manuscript as submitted and agree to be accountable for all aspects of the work.

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ABSTRACT

Objectives: Despite the aggressive marketing of electronic nicotine device systems (ENDS) as smoking cessation tools, the evidence of their effectiveness is mixed. We conducted a systematic review of randomized controlled trials to determine the effect of ENDS on cigarette smoking cessation, as compared to other types of nicotine replacement therapies (NRT).

Methods: We included randomized controlled trials in which any type of ENDS was compared to any type of NRT, in traditional cigarette users. We searched MEDLINE, Embase, and the CENTRAL Trials Registry of the Cochrane Collaboration using the Ovid interface, as well as ClinicalTrials.gov and the World Health Organization International Clinical Trials Registry Platform trials registries regardless of study completion status. We used the Revised Cochrane risk-of-bias tool for randomized trials (RoB 2) for each outcome of interest. The primary outcome was smoking cessation. Secondary outcomes included smoking reduction, harms, withdrawal, and acceptance of therapy. A random-effect model was used, and data were pooled in meta-analyses where appropriate.

Results: Six studies were retained from an initial 270. Most outcomes were judged to be at high risk of bias. The overall quality of evidence was graded as 'low' or 'very low'. Pooled results showed no difference in smoking cessation (RR 1.42 [0.97, 2.09]), proportion of participants reducing smoking consumption (RR 1.25 [0.79, 1.98]), mean reduction in cigarettes smoked per day (MD 1.11 [-0.41, 2.63]), or harms (RR 0.96 [0.76, 1.20]), between groups.

Discussion: We found no difference in smoking cessation, harms, and smoking reduction between e-cigarette and NRT users. However, the quality of the evidence was low. Further research is needed before widespread recommendations can be made with regards to the use of ENDS. Research is also needed to investigate the long-term effects of ENDS, as well as optimal dosing.

Systematic review registration number: protocol registered with the International Prospective Register of Systematic Reviews (PROSPERO) on February 27th, 2020. Registration number pending.

Strengths and limitations of this study

- This study provides up to date meta-analyses of direct comparisons of vaping with nicotine replacement therapy for smoking cessation, studied through randomized controlled trials.
- We examined harms associated with vaping, which are becoming increasingly concerning.
- This study makes extensive efforts to obtain unreported data from investigators.
- Careful consideration is given to the potential impact of risk of bias and methodological heterogeneity.
- As we included only RCTs, many studies that used weaker study designs were ineligible for this review.

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SUMMARY OF FINDINGS TABLE

Nicotine-containing Electronic cigarettes (ENDS) vs Nicotine Replacement Therapies (NRT) for smoking cessation

Population: Current smoke Intervention: Nicotine-con				
Comparison: Nicotine-repl				
Outcomes ENDS as compared to NRT	Relative effect (95% Cl)	Number of participants (studies)	Quality of the evidence (GRADE)	Comments
Cessation	RR 1.42 [0.97, 2.09]	1800 (5 studies)	⊕⊕ OO ^{1,2} low	
Smoking reduction Proportion of people decreasing cigarette consumption by 50%	RR 1.25 [0.79, 1.98]	1460 (4 studies)	⊕⊕ OO ^{1,2} low	
Mean decrease in cigarettes per day	MD 1.11 [-0.41, 2.63]	633 (3 studies)	⊕⊕ 00 ^{1,2} low	
Adverse events (AEs)	RR 0.96 [0.76, 1.20]	758 (4 studies)	<pre> ① OOO^{1,2,3} Very low </pre>	No severe adverse events related to investigated products were reported
Withdrawal symptoms	Summary data not available	4 studies	<pre> ① OOO^{1,2,3} Very low </pre>	Withdrawal measures included Minnesota Nicotin Withdrawal Scale, QSU scores, frequency of urge and strength of urge score, and pre-specified symptoms of depressed mood, irritability, restlessness, and hunger

Acceptance of therapy	Summary data not available	4 studies	000 ^{1,2,3} Very low	Acceptance define as wanting to recommend product to friends, helpfulness, taste, satisfaction, psychological reward, enjoyment of sensation, aversion, and	
				ability to reduce craving depending on study	
GRADE Working Group grades High quality: Further research Moderate quality: Further res effect and may change the est	is very unlikely to chang earch is likely to have ar imate.	n important impact	on our confidence	in the estimate of	
Low quality: Further research and is likely to change the esti Very low quality: We are very	mate.		on our confidence i	n the estimate of effe	
¹ Downgraded one level becau ² Downgraded one level becau ³ Downgraded one level becau	se of heterogeneity se of imprecision of res				

INTRODUCTION

Despite a significant lack of rigorous pharmacological testing, the use of electronic nicotine device systems (ENDS), otherwise known as vaping devices, has been aggressively marketed as an effective method to quit smoking. In Canada, 32% of current and former smokers report having used ENDS as a smoking cessation aid. In addition to delivering nicotine to the user, ENDS are thought to replace some of the habitual behaviours and sensations associated with smoking, such as the action of bringing a cigarette to the mouth. By doing so, ENDS may provide coping mechanisms that other traditional nicotine replacement therapies (NRT) do not offer, and therefore may help with the behavioural component of smoking reduction and cessation.² While vaping is believed to be less harmful than cigarette smoking, a large number of emerging reports on the health impacts of vaping are worrisome. In addition, the evidence on the effectiveness of ENDS as a smoking cessation aid is mixed.

In 2016, a meta-analysis of 20 studies found that people using ENDS had a 28% reduction in the odds of stopping cigarette smoking as compared to those not using ENDS.³ However, in a 2019 recent randomized controlled trial (RCT), individuals randomized to nicotine-containing e-cigarettes were more likely to abstain from smoking at one year compared to individuals randomized to nicotine patches (18% compared to 9.9%, RR 1.83; 95% CI 1.30 to 2.58).⁴ A Cochrane review⁵ found that nicotine-containing e-cigarettes were more effective than non-nicotine containing e-cigarettes for smoking cessation, but was not able to compare ENDS products to traditional NRT.

Little information is known about the long-term health impacts of ENDS. Reports of acute toxicity have recently captured the public's attention. In late 2019 and early 2020, "e-cigarette, or vaping, product use-associated lung injury" (EVALI) caused 2807 illnesses and 68 deaths in the US,⁶ and 19 cases in Canada.⁷ Other short-term adverse events reported with the use of ENDS include cardiovascular changes such as increased heart rate and blood pressure, cough, wheeze,⁸ and mucus production.⁹ Burn injuries have also been reported, as well as fatalities from drinking or injecting the e-liquid.⁸

There is no long-term data available on the relationship between ENDS and oral, respiratory, and cardiovascular health, as well as cancer. There is however available data linking the chemicals present in e-liquids with cellular DNA damage and carcinogenicity.^{9,10} There is some evidence that the use of ENDS is associated with asthma exacerbations.¹¹ No human long-term data exist on the use of ENDS in pregnancy and their impact on the developing fetus.

Given the large number of smokers using ENDS as a potential smoking cessation tool, there is a need to review and synthesize the evidence of trials examining a head to head comparison of ENDS versus traditional NRT for smoking cessation.

Objective

The objective of this review is to systematically review the evidence found in RCTs to determine the effect of electronic nicotine delivery systems (ENDS) on cigarette smoking cessation in smokers, as compared to other types of nicotine replacement therapies (NRT).

METHODS

Protocol and registration

The protocol for this systematic review was submitted to International Prospective Register of Systematic Reviews (PROSPERO) on February 27th, 2020 (registration pending) and uploaded as a preprint on Open Science Framework (OSF) Preprints on May 12th 2020.¹²

Criteria for study inclusion

Study Characteristics:

RCTs in which ENDS were compared to non-electronic NRT in smokers were included. We restricted our inclusion to RCTs to minimize the risk of bias. No language limits were imposed. No date limits were imposed either, although we did not anticipate studies published prior to 2003, since this is when the first e-cigarette was invented.¹³ There was no geographical restriction of studies.

Study Population:

All traditional cigarette users were included, regardless of age, amount of traditional cigarette use, and motivation to quit.

Intervention of interest:

The intervention of interest comprised all types, models, and brands of ENDS.

<u>Comparators:</u>

All included studies compared ENDS with non-electronic NRT. NRT comprised, but were not limited to, nicotine patch, gum, lozenge, nasal spray, inhalator, mouth spray, mouth strips, microtabs, and combination of products.

Outcome measures:

The primary outcome measure is traditional cigarette smoking cessation defined as abstinence from traditional cigarette smoking for any time period, as reported in each included study, regardless of whether abstinence is self-reported or biochemically validated. Secondary outcomes include reduction in the number of traditional cigarettes smoked in any given time period, adverse events, withdrawal symptoms, and participants' acceptance of therapy. We had planned on collecting quit attempts information but none of the studies reported on this outcome.

Settings:

All health care and community settings were included.

Study Identification

The following databases were searched: MEDLINE (1946 to June 2020), Embase (1947 to June 2020) and the CENTRAL Trials Registry of the Cochrane Collaboration (May 2020 Issue) using the Ovid interface. The MEDLINE search was limited using the Cochrane Highly Sensitive Search Strategy and the Embase search was limited using the recommended limit for controlled

trials.¹⁴ Searches were developed by a librarian experienced in systematic reviews, using a method designed to optimize term selection.¹⁵ ClinicalTrials.gov and World Health Organization International Clinical Trials Registry Platform (WHO ICTRP) trials registries were searched for registered intervention studies, regardless of their completion status. Electronic search strategies are presented in Supplementary Material 1. The reference lists of included studies and any applicable review studies were searched.

Authors of protocols identified through registries were contacted electronically, to request data for the review. In addition, clinical experts in the field of vaping and smoking cessation were contacted to enquire about any unpublished research fulfilling our inclusion criteria.

Selection of Studies

Records retrieved by the electronic search were downloaded and imported into a Reference Manager database for duplicate removal, and then uploaded to Covidence. Throughout the review, newly identified records were integrated into the set for screening.

Each title and abstract was independently screened by two review authors (from CP, JZ, and ATK) against the eligibility criteria.¹⁴ Full text of all studies deemed potentially eligible was obtained and reviewed independently by two of the same review authors to determine eligibility. For screening, data extraction, and risk of bias assessment, disagreements were resolved by discussion, and with a third reviewer when needed.

Data extraction and management

For studies that fulfilled the inclusion criteria, two reviewers (CP, JZ) extracted the data into an electronic data collection form, which was piloted by both reviewers (Supplementary Material 2). The data collection was revised, based on feedback from the reviewers. Study authors were contacted electronically to obtain relevant but unavailable data.

Risk of bias assessment for included studies

Two reviewers (CP, JZ) independently conducted the risk of bias assessment for each study at the outcome level using the Revised Cochrane risk-of-bias tool for randomized trials (RoB 2).¹⁶

Measures of treatment effect

Dichotomous data was analyzed by calculating the risk ratio, using the longest follow-up time reported, as well as the 95% confidence interval. The risk ratio (RR) for smoking cessation was calculated as such:

 $RR = \frac{1}{N \text{ of subjects abstaining from smoking in control}/N \text{ of subjects in control}}$

Continuous data for the secondary outcomes were analyzed through mean differences between groups as the same scales were used. In the case of studies with multiple arms, we only extracted data for the groups relevant to this review.

Data synthesis

We provide a narrative synthesis of the included studies. Where appropriate, data have been pooled for meta-analyses, and random effects were used for all analyses in RevMan.¹⁴ The inverse-variance random-effects and the mean difference approach (using standard deviations and sample sizes) were used for dichotomous and continuous outcomes, respectively, to assign the weight given to each study. Participants with missing data were considered as still smoking.⁵ The proportion of adverse events reported was based on the number of people available for outcome assessment. For the reduction of the number of cigarettes smoked, missing values were assumed to be zero.

Assessment of heterogeneity

A p value of 0.10 for the chi-squared test (Cochrane Q) and an I² value of >50% were used as indicators of substantial heterogeneity. This however needs to be interpreted with caution given the small number of studies available for the meta-analysis. Clinical and methodological diversity was also explored.

We planned to assess reporting/publication bias using funnel plots of effect estimate against standard error, and testing for funnel plot asymmetry, however, the number of included studies was too low (<10).

We also planned on conducting a number of sensitivity analyses to determine the robustness of the results of the meta-analyses; subgroup analyses to investigate potentially modifying factors such as age and smoking intensity; as well as meta-regression to study the impact of covariates such as motivation to quit smoking, provision of training, and other factors,¹⁷ but minimum data thresholds were not met.

We present a 'Summary of Findings' table for all outcomes. We used the five GRADE considerations (study limitations, consistency of effect, imprecision, indirectness, and publication bias)¹⁴ to assess the quality of evidence for each outcome and to draw conclusions about the robustness of evidence within this review.

RESULTS

Our initial bibliographic search yielded 270 records, and after screening and full-text review, we retained 6 RCTs. An updated search conducted in June 2020 yielded an additional 116 records (for a total of 386 records), none of which were included after screening (Figure 1).

We identified six RCTs (Bullen 2013,¹⁸ Eisenhofer 2015,¹⁹ Hajek 2019,⁴ Hatsukami 2019,²⁰ Lee SH 2019,²¹ Lee SM 2018²²). Of these, five contributed data to our primary outcome of smoking cessation.^{4,18,20-22} Four studies^{4,18,21,22} examined cessation at 6 months or longer, while one²⁰ examined short term cessation (< 6 months). Table 1 includes the salient features of the

included studies. A more detailed description of included studies can be found in Supplementary Material Table 3.

Author and year of publication	Design	Country	Number of participants	Main eligibility criteria	Intervention	Comparator	Main outcome of interest
Bullen, 2013 ¹⁷	3-group, parallel, single center	Australia	657 total, 584 included in this review (2 of 3 groups)	\geq 18 years, smoked \geq 10 cigarettes per day in the past year, motivated to quit	First- generation e- cigarette x 12 weeks	Nicotine patch x 12 weeks	Continuous abstinence 6 months after quit day
Hajek 2019 ⁴	2-group, parallel, multi- centre	United Kingdom	884	Adults with no strong preference towards e- cigarette or NRT	Any type of e-cigarette	Any nicotine- replacement therapy	Continuous abstinence 52 weeks after quit day
Lee SH, 2019 ²⁰	2-group, parallel, single center	Republic of Korea	150	\geq 18 years, smoked \geq 10 cigarettes per day in the past year, motivated to quit	e-cigarette x 24 weeks	Nicotine gum x 24 weeks	Continuous abstinence 24 weeks after quit day
Lee SM, 2018 ²¹	2 group, parallel, single center	USA	30	Adults, smoked \geq 2 cigarettes per day in the past year, smoked at least once in last 7 days	e-cigarette x 6 weeks	Nicotine patch x 5 weeks, then placebo patch x 1 week	7-day point prevalence abstinence at 6 months
Characteris Hatsukami, 2019 ¹⁹	4 group, parallel, multi- center	T measuri USA	ng smoking c 264 total, 152 included in this review (2 of 4 groups)	Example 2 18 years, smoked \geq 5 cigarettes per day	rlier than 6 m e-cigarettes	Nicotine gum or nicotine lozenge	7-day point prevalence abstinence at 8 months

Eisenhofer, 2015 ¹⁸	2-group, parallel, single center	USA	11	Veterans who met criteria for tobacco	e-cigarettes x 3 weeks	Nicotine patch x 3 weeks	Reduction in number of cigarettes smoked per
				disorder			day at 3
							weeks

Risk of bias in included studies

We assessed risk of bias for each included study. A detailed report of the risk of bias assessment can be found in Supplementary Material Table 4.

Figures 2a, 2b, 2c, 2d, and 2e illustrate the risk of bias for each outcome.

Effect of Interventions

Smoking cessation

Five of the six studies reported on smoking cessation.^{4,18,20-22} When comparing e-cigarettes to NRT in the context of smoking cessation, there was no significant difference between groups in verified self-reported continuous abstinence at 6 months (21/289 vs 17/295, RR 1.26 [0.68, 2.34], p=0.46) in the Bullen 2013¹⁸ study, and in continuous abstinence from 9 to 24 weeks (16/75 vs 21/75, RR 0.76 [0.43, 1.34], p = 0.344) in the Lee SH 2019²¹ study. In addition, the Lee SM 2018²² study showed no difference between groups for the 7-day point prevalence abstinence at 6 months in the context of perioperative smoking cessation (5/20 vs 1/10, RR 2.50 [0.34, 18.63], p = 0.63).

In the Hajek 2019⁴ study, self-reported, verified continuous abstinence at 1 year was found to be higher in the e-cigarette group (79/438 vs 44/446, RR 1.83 [1.30, 2.58], P<0.001), and smoking cessation assessed by 7-day point prevalence at 8 weeks in the Hatsukami 2019²⁰ trial was also higher in the e-cigarette group (25/76 vs 13/76, RR 1.92 [1.07, 4.37], p = 0.039).

We combined data from all 5 studies comparing smoking cessation between e-cigarettes and NRT and obtained a pooled RR of 1.42 [0.97, 2.09].

Smoking reduction

All six studies^{4,18-22} assessed smoking reduction. Bullen 2013,¹⁸, Eisenhofer,¹⁹ Hajek 2019,⁴ and Lee SM 2018²² reported the proportion of participants reducing smoking by at least 50%. While Lee SH 2019²¹ also reported on this outcome, the size of the reduction was not specified. Bullen 2013¹⁸ and Lee SH 2019²¹ reported an absolute reduction, and Hatsukami 2019²⁰ reported a relative reduction in cigarettes per day from baseline.

In the Bullen 2013 study,¹⁸ mean cigarette consumption at 6 months decreased by 9.7 (SE 0.4) in the e-cigarette group, and by 7.7 (SE 0.4) in the NRT group. Mean difference between groups was 1.9 (SE 0.6) (p = 0.002). After excluding people who successfully quit smoking, the RR of decreasing cigarette smoking by at least 50% when comparing the e-cigarette to the NRT groups was 1.61 [1.31, 1.99].

Eisenhofer 2015¹⁹ compared week 3 to week 1, and showed that both e-cigarettes (t = 5.3, p = 0.013) and NRT (t = 3.4, p = 0.015) significantly reduced (\sim 50%) self-reports of cigarettes smoked in the previous 24 hours. This was confirmed by significant reductions of breath CO levels in both groups No additional information could be obtained from the abstract and none of the authors could be reached.

In the Hajek 2019⁴ study, 44 of 345 participants in the e-cigarette group, and 29 of 393 participants in the NRT group experienced a carbon monoxide-validated reduction in smoking of \geq 50% in participants without abstinence between weeks 26 and 52, yielding a relative risk of smoking reduction of 1.73 (1.11-2.70).

Hatsukami 2019²⁰ defined smoking reduction by the estimated ratio of cigarettes smoked at 8 weeks as compared to baseline, with a result of 0.25 (0.17, 0.37) in the e-cigarette group, and 0.29 (0.21, 0.39) in the NRT group (p = 0.185). Additional data obtained from the author showed that 19 participants in the e-cigarette group and 22 participants in the NRT group reduced smoking consumption by 50% (RR 0.86 [0.51, 1.46]) at 8 weeks, and that mean cigarette consumption decreased by 9.22 (SD 7.95) in the e-cigarette group, and by 7.61 (SD 8.27) in the NRT group. The mean difference between groups was 1.61 [-0.97, 4.19].

In the Lee SH 2019²¹ study, mean cigarette consumption decreased at 24 weeks by 6.5 +/- 2.87 (SD) in the e-cigarette group, and by 6.60 +/- 3.75 (SD) in the NRT group (p = 0.974). In addition, 31 out of 75 participants (41.3%) in the e-cigarette group and 19 out of 75 participants (25.3%) in the NRT group reduced their daily cigarette consumption (p = 0.038), but no information on size of smoking reduction is provided. After excluding abstainers, a RR of 1.49 [0.97, 2.31] was obtained for decrease in daily cigarette consumption.

Lastly, in the Lee SM 2018,²² 1 participant in the END group and 4 participants in the NRT group reduced their cigarette consumption by at least half, resulting in a RR 0.15 [0.02, 1.14].

We combined data from the Bullen 2013,¹⁸ Hajek 2019,⁴ Hatsukami 2019²⁰ and Lee SM 2018²² studies comparing smoking reduction of at least 50% between e-cigarettes and NRT, as they used similar measures. Pooled results comparing the difference in smoking reduction between the e-cigarette and the NRT groups produced a RR of 1.25, with the line of equivalence falling within the confidence interval [0.79, 1.98].

We also combined data from the Bullen 2013,¹⁸, Hatsukami 2019,²⁰ and Lee SH 2019²¹ comparing mean reduction of cigarettes per day from baseline for ENDs and NRT (Figure 3c). Meta-analysis yielded a MD of 1.11, with the line of equivalence falling within the confidence interval [-0.41, 2.63].

<u>Harms</u>

Five studies reported on harms (Bullen 2013,¹⁸ Hajek 2019,⁴ Hatsukami 2019,²⁰ Lee SH 2019,²¹ Lee SM 2018²¹). None of the included studies reported serious adverse events (SAEs) related to e-cigarettes or NRT.

In the Bullen 2013¹⁸ study, 107 participants in the e-cigarette group reported 137 adverse events, while 96 participants in the NRT group (patches) reported 119 events, and, using the number of participants available for analysis at 6 months, there was no difference in the incidence of adverse events between groups (RR 0.99, [0.81, 1,22]). No difference between groups was also observed in the Hatsukami 2019²⁰ study, where additional data provided by the author showed that 51 of 69 participants in the e-cigarette group and 53 of 72 participants in the NRT group reported adverse events (1.00 [0.82, 1.22]), and in the Lee SM 2018²² study, where no significant difference in the incidence of adverse events between at 8 weeks (RR 1.24 [0.54, 2.84]).

Hajek 2019⁴ defined adverse events of interest as nausea, sleep disturbances, and throat and mouth irritation. There were 27 SAEs in the e-cigarette group and 22 in the NRT group, none felt to be related to the intervention or control products. Based on the number of participants available at the 12 month follow-up, e-cigarettes were found to be less likely associated with nausea (RR 0.78 [0.66, 0.92]) and sleep disturbances (RR 0.88 [0.83, 0.95]), but more likely associated with throat/mouth irritation (RR 1.24 [1.13, 1.37]). These numbers however should be interpreted with caution as it was not possible to determine with certainty the denominator from the data.

In the Lee SH 2019 study, ²¹ 5 participants in the e-cigarette group and 13 participants in the nicotine gum group reported adverse events. There were no SAEs. Based on the number of participants who completed the study, e-cigarettes were less likely to be associated with adverse events (RR 0.13 [0.12, 0.87]).

We combined data from the Bullen 2013,¹⁸ Hatsukami 2019,²⁰ Lee SH 2019,²¹ Lee SM 2018²² studies comparing harms between e-cigarettes and NRT. Hajek 2019⁴ was excluded as they did not clearly report the number of participants that experienced any adverse events and reported only on specific adverse events. Pooled results comparing ENDS to NRT yielded a RR of 0.96 [0.76, 1.20].

<u>Withdrawal symptoms</u>

Four studies reported on the results of withdrawal symptoms (Eisenhofer 2015,¹⁹ Hajek 2019,⁴ Hatsukami 2019,²⁰ and Lee SM 2018²²) and all used different scales. Eisenhofer 2015¹⁹ assessed withdrawal with the Questionnaire on Smoking Urges (QSU), Hajek⁴ used a composite urge score (frequency and strength of urge to smoke), Hatsukami 2019²⁰ measured the severity of withdrawal using the Minnesota Nicotine Withdrawal Scale, and Lee SM 2018²¹ assessed withdrawal symptoms as part of their adverse event assessment. In light of the differences in outcome assessment measures, the data were not pooled.

In Eisenhofer 2015,¹⁹ urges and cravings to smoke were significantly reduced in the e-cigarette group (t=3.8, p = 0.03), but not in the NRT group (t=2.1, p = 0.08).

In Hajek 2019,⁴ urges for e-cigarette users decreased more than for NRT users at 1 week (MD: - 0.4 (-0.6 to -0.2)) and at 4 weeks (MD: -0.3 (-0.5 to -0.1)). E-cigarette users also reported a smaller increase from baseline in irritability, restlessness, inability to concentrate, hunger, and depression. The withdrawal symptoms disappeared mostly for both groups by week 4.

In Hatsukami 2019,²⁰ participants in the e-cigarette group reported lower median [min/max] changes from baseline on the severity scale compared to participants in the NRT group at all measurement points, with week 1 (3.0 [-9.0/25.0] vs 3.5 [-20.0/32.0]), week 2 (1.0 [-13.0/25.0] vs 3.0 [-13.0/39.0]), and week 4 (1.0 [-17.0/30.0] vs 2.5 [-28.0/29.0]). The planned pairwise comparisons were significant with p <0.017. As well, fewer participants (5.3%) withdrew from the complete substitution e-cigarettes group than from the NRT group (15.8%) for product related reasons (disliking product or experiencing withdrawal symptoms; p value not reported).

Lee SM 2018²² only reported on withdrawal symptoms for the NRT group, and did not report on withdrawal symptoms for the e-cigarette group.

Acceptance of therapy

Four studies reported on acceptance of therapy (Bullen 2013,¹⁸ Hajek 2019,⁴ Hatsukami 2019,²⁰ and Lee SM 2018²²), and all used different scales. In light of the difference in outcome assessment measures, the data were not pooled.

In the Bullen 2013¹study,¹⁸ 230 out of 260 participants (88%) in the e-cigarettes group said they would recommend their allocated product to a friend at 1 month, as compared to 130 out of 232 participants (56%) in the NRT group (RR 1.58 [1.40, 1.78]). At 6 months, 205 out of 241 participants (85%) in the e-cigarettes group said they would recommend their allocated product as compared to 107 out of 215 participants (50%) in the NRT group (RR 1.71 [1.48, 1.97]).

In the Hajek 2019 study,⁴ acceptance of therapy was measured with a Likert scale (1 to 5, with a higher score associated with higher acceptance). At 4 weeks post quit date, helpfulness of e-cigarettes was rated 4.3 (SD 0.9) while that of NRT was 3.7 (SD 0.9) (mean difference 0.6 (0.4, 0.7)). Taste was scored at 3.5 (SD 1.3) for the e-cigarette group and 3.1 (SD 1.5) (mean difference 0.4 (0.2,0.6)), and satisfaction was rated at 2.7 (SD 1.1) and 2.3 (SD 1.2), respectively, for the e-cigarette and NRT groups (mean difference 0.5 (0.3, 0.6)).

In the Hatsukami 2019 study²⁰, acceptance of therapy was defined as satisfaction with the product, psychological reward, enjoyment of sensation, aversion, and ability to reduce craving. Results are reported for the NRT group as an estimated mean difference and 95% CI in product evaluation sub-scales using the e-cigarette group as a reference. The following results are reported; satisfaction: -0.6 (-1.0, -0.1), psychological reward: -0.4 (-0.8, 0.01), enjoyment of sensation: -0.6 (-1.1, -0.1), aversion: 0.1 (-0.2, 0.4), and ability to reduce craving: -0.3 (-0.8, 0.2).

Lastly, the Lee SM 2018 trial²² defined acceptance of therapy as satisfaction with the assigned product, measured with a Likert scale (1 to 7, with a higher score associated with higher satisfaction). Median scores and IQR are reported. Participants randomized to the e-cigarette

group reported scores of 6 [4-7], 5.5 [2.5-7], and 6 [5-7], respectively, while participants randomized to the NRT group reported scores of 5 [3-7], 5 [3-6], and 7 [6-7], respectively for the following questions. "The product is helpful for quitting smoking", "I was satisfied with the product to help with quitting", "I would recommend the product to someone interested in quitting smoking".

Risk of bias across studies

The review process we used was thorough, and we took every precaution to minimize the risk of bias due to publication bias or selective reporting. We reached out to clinical experts to enquire about unpublished reports, examined protocol registries, and contacted the authors of identified protocols to request unpublished results. Given the low number of retained studies, we did not include a funnel plot.

Sensitivity, subgroup and meta-regression analyses

We performed a sensitivity analysis for the smoking cessation outcome by removing the Lee SM 2018 study²². While the other 4 studies aimed to assess smoking cessation in general, Lee et al were targeting a peri-operative population, who may have had different motivations to quit smoking. The pooled data, once Lee SM 2018²² is removed, yield a RR of smoking abstinence of 1.39 [0.92, 2.11] when comparing ENDS to NRT (Figure 4a).

We had planned on undertaking multiple subgroup analyses. We were unable to perform the subgroup analyses based on age (all participants were adults), smoking intensity (no study enrolled smokers \geq 25 cigarettes per day), or biochemically validated smoking cessation (all studies used biochemical validation). We also could not perform a subgroup analysis of studies with ties to industry as only Bullen 2013¹⁸ was found to have ties to the vaping industry.

We did, however, perform the following subgroup analyses: limiting comparator to nicotine patches (Bullen 2013¹⁸ and Lee SM 2018²¹), and including only studies assessing continuous/sustained smoking abstinence ≥ 6 months given that smoking cessation is defined as sustained abstinence for at least 6 months;²³ (Bullen 2013,¹⁸ Hajek 2019,⁴ Lee SH 2019²¹) (Figures 4b and 4c, respectively).

Metaregression analyses were not performed as our threshold of 10 eligible studies was not met.

DISCUSSION

In our review, there was no significant difference in smoking cessation, smoking reduction, or harms between e-cigarette and NRT users. However, we report on results from a limited number of RCTs, and the level of evidence is low. Our efficacy results are similar to those described in a 2016 Cochrane review,⁵ which also showed no difference between abstinence rates between the nicotine e-cigarette group and NRT group. Their review only included one study¹⁸, also included in our review for this particular outcome. Similar to the evidence we are

presenting, none of the studies examined in the Cochrane review reported serious adverse events considered to be related to e-cigarette use.

Although our meta-analysis of the 5 trials that examined *smoking cessation* showed no significant difference between e-cigarette and nicotine replacement therapy, there was a trend towards favoring e-cigarettes. Interestingly, our sensitivity analysis limiting inclusion to studies reporting smoking cessation of 6 months or greater yielded a smaller point estimate than the one obtained from the main analysis, although still with no difference between groups. It could be hypothesized that additional benefits that may be attributed to e-cigarette early on in smoking cessation may be attenuated as time progresses. This again should be interpreted with caution given the small number of studies^{4,18,20} and the very significant heterogeneity.

In all comparisons, our results need to be interpreted carefully. There was significant clinical heterogeneity between studies in terms of the population enrolled, smoking intensity at baseline, type and nicotine concentration of e-cigarettes, type and dose of NRT, as well as methodological heterogeneity in terms of study conduct, and intervention and control protocols. For instance, one of the included studies¹⁸ used first-generation e-cigarettes, with nicotine delivery about 20% of that obtained from cigarette smoking. While e-cigarette users were couriered the supplies needed, NRT users had to redeem vouchers from community pharmacies to obtain their patches. The low nicotine content of the e-cigarettes, the extra step in obtaining NRT supplies, and the low intensity of additional co-interventions likely contributed to the low rate of smoking abstinence at 6 months in both groups, limiting the generalizability of the results. Another included study⁴ allowed for multiple types and concentrations of ENDS, as well as upwards of 10 NRT products and doses, complicating the interpretation of the results. Nicotine concentrations reported in the trials ranged from 0.01 to 48 mg/mL,^{4,18,20-22} making comparisons between studies difficult.

Given that the risk of bias was assessed as high in 5 of 6 included studies^{4,18-21}, our smoking cessation outcome results need to be interpreted with caution. In addition, it is interesting to note that all studies verified self-reported smoking cessation with an exhaled carbon monoxide test, however different cut-off values were used. Additionally, there are limitations to using carbon monoxide (CO) as a way to verify smoking cessation. CO has a relatively short half-life and is eliminated from the body within 24 hours; it can, therefore, lead to false negative results. However, this issue is somewhat mitigated by the fact that smoking cessation study participants tend to be daily smokers.

All studies included in this review examined *smoking reduction*. There was no difference between groups in the mean reduction of cigarettes from baseline in the studies that measured that outcome, or in the proportion of participants successfully reducing their smoking consumption.

None of the included studies reported severe *adverse events* related to ENDS or NRT, and, for the four studies with data that could be pooled, there was no difference between groups in terms of harms related to either therapy. However, in addition to the clinical heterogeneity

mentioned above, there was significant methodological heterogeneity in how adverse events were collected. We evaluated the quality of the evidence as very low, given the high risk of bias of included studies, the significant heterogeneity, and the inability to accurately determine the number of subjects involved in this outcome, thus leading to result imprecision.

Since the included trials were powered to detect a difference in the primary outcome, it is possible that rare or unexpected harms were not detected due to a lack of power for this specific outcome. Also, it is important to acknowledge that these studies are limited by their short time-frame. Data on long-term side effects of ENDS are lacking. The recent e-cigarette, or vaping product use-associated lung injury (EVALI) epidemic, is a reminder that further research is needed before widespread recommendations can be made with regards to the use of ENDS. In addition, there are now emerging concerns that respiratory disease caused by the novel coronavirus SARS-CoV-2, the virus responsible for the COVID-19 pandemic, could be exacerbated by exposure to ENDS.²⁴⁻²⁶

Finally, although there seemed to be *increased acceptance of therapy* towards e-cigarettes in the four studies that considered it,^{4,18,20,22} high risk of bias, significant heterogeneity, and the small number of studies using widely different scales leading to imprecise measures, mean that the results should be interpreted with extreme caution. In addition, given that the trials were unblinded, participants who were disappointed with their treatment allocation may have reported less acceptability than their counterparts.

Limitations at review level

We restricted our search to RCTs to try to minimize the risk of bias, however, this considerably limited the number of available studies for this review. It is surprising that, given the widespread availability of e-cigarettes and how aggressively they have been marketed as smoking cessation agents, there are so few head-to-head trials comparing ENDS and traditional NRT. While there may be some unpublished studies that our review did not capture, our literature search was thorough and included personal communications to multiple experts in the field.

Our review identified 7 ongoing trials²⁷⁻³³ that potentially met our inclusion criteria, totaling over 1500 targeted participants. None of the investigators had any data ready to be shared, however it is hoped that this ongoing research can shed light on the effectiveness of ENDS as smoking cessation tools, as compared to traditional NRTs. Long-term research is also needed to investigate the long-term effects of ENDS, as well as the optimal dosing and method of delivery.

Conclusion

We found no difference in smoking cessation, harms, and smoking reduction between ecigarette and NRT users. However, the quality of the evidence was low. Further research is needed before widespread recommendations can be made with regards to the use of ENDS. Research is also needed to investigate the long-term effects of ENDS, as well as optimal dosing.

Acknowledgements

We thank Katie O'Hearn, MSc, (Children's Hospital of Eastern Ontario Research Institute), Dr Matthew McInnes, and Dr Dean Fergusson (University of Ottawa), for methodological assistance.

Data sharing

Data collection forms and all raw data can be requested through the corresponding author.

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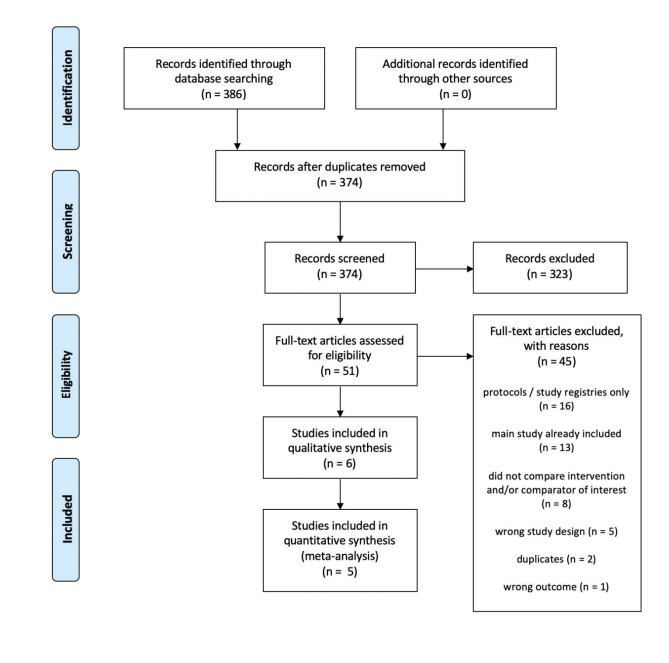
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Figure 1. Study flow diagram



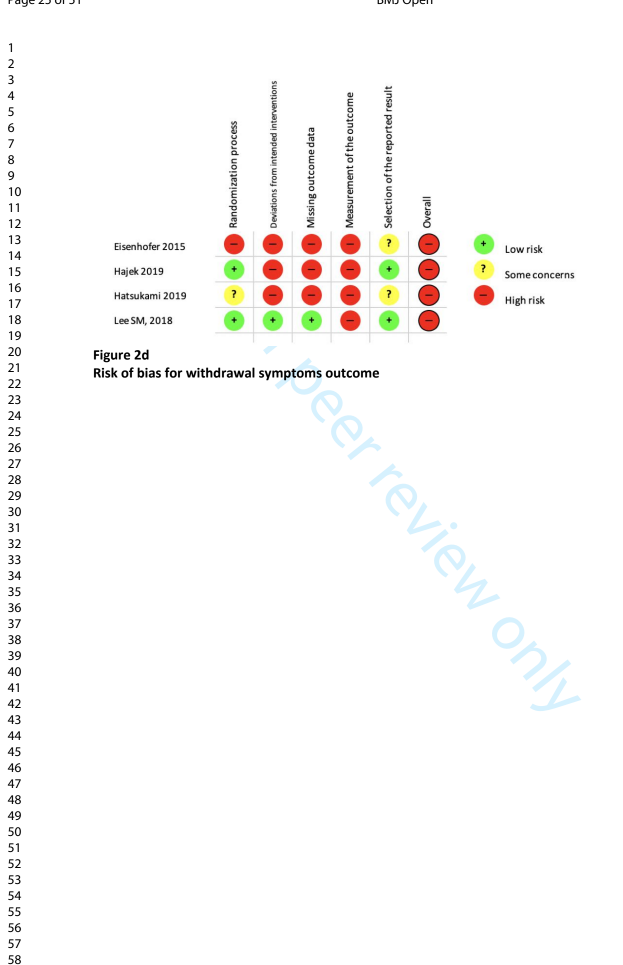
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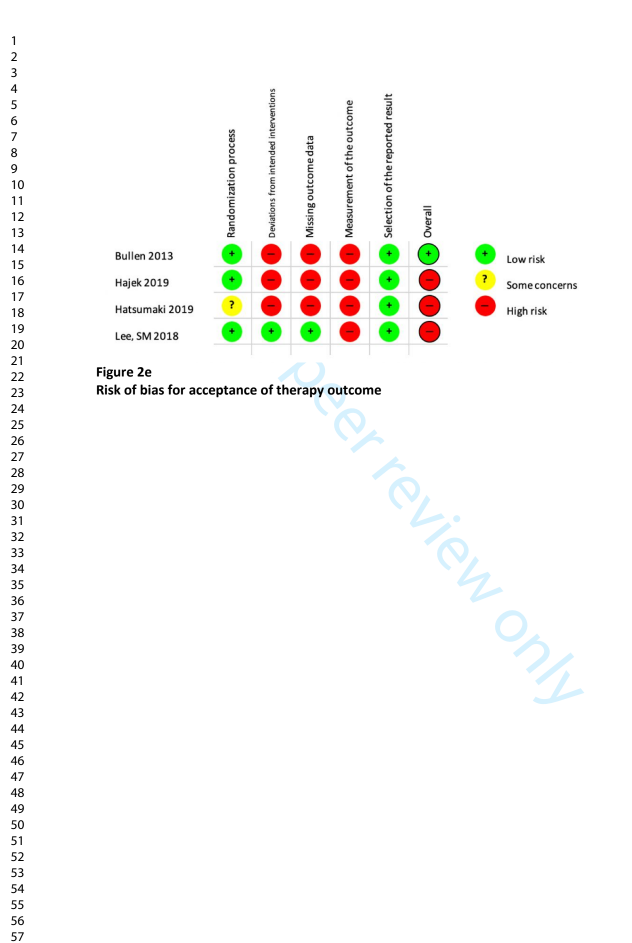
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3 4	Study or Subgroup	ENDS Events		NRT Events		Weight	Risk Ratio IV, Random, 95% CI	Risk Ratio IV, Random, 95% Cl
5	Bullen	21	289	17	295	20.5%	1.26 [0.68, 2.34]	
6	Hajek	79	438	44	446	32.2%	1.83 [1.30, 2.58]	
7	Hatsukami Lee SH	25 16	76 75	13 21	76 75	21.5% 22.4%	1.92 [1.07, 3.47] 0.76 [0.43, 1.34]	
8	Lee SM	5	20	1	10	3.4%	2.50 [0.34, 18.63]	
9								
9 10	Total (95% CI)	140	898	0.0	902	100.0%	1.42 [0.97, 2.09]	•
10	Total events Heterogeneity: Tau ² =	146 0.09 [.] Chi	$i^2 = 8.0$	96 4 = 10 00	4 (P =	0 09)· 12	= 50%	
12	Test for overall effect:				. (. –	0.05), 1	_ 50%	0.01 0.1 1 10 100 Favours [NRT] Favours [E-cigarette]
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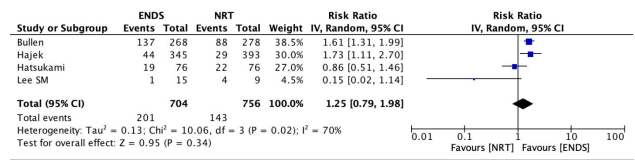


Figure 3b Proportion of participants successfully reducing smoking consumption by 50%

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Study on Subarrow	ENDS Moon SE		NRT Moon S		Weicht	Mean Difference	Mean Difference
Study or Subgroup Bullen	Mean SE 9.7 5.37	Total 7 180	7.7 5.			IV, Random, 95% CI 2.00 [0.89, 3.11]	
Hatsukami	9.22 7.95	76	7.61 8.2	7 76	20.6%	1.61 [-0.97, 4.19]	+
Lee SH	6.55 2.87		6.6 3.7	5 61	39.3%	-0.05 [-1.20, 1.10]	T
Total (95% CI)	1 10 01 2	327	<pre>c > /p </pre>		100.0%	1.11 [-0.41, 2.63]	
Heterogeneity: Tau ² = Test for overall effect:			f = 2 (P = 0)	.04); 1- =	= 69%		-100 -50 0 Favours [NRT] Favours
							Favours [INKT] Favours
Figure 3c Mean re	eduction d	or cigai	rettes fro	m bas	eline		

	END	s	NR	Г		Risk Ratio	Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Bullen	107	241	96	215	43.5%	0.99 [0.81, 1.22]	*
Hatsukami	51	69	53	72	44.7%	1.00 [0.82, 1.22]	+
Lee SH	5	71	13	61	5.0%	0.33 [0.12, 0.87]	a <u></u>
Lee SM	11	20	4	9	6.7%	1.24 [0.54, 2.84]	
Total (95% CI)		401		357	100.0%	0.96 [0.76, 1.20]	•
Total events	174		166				
Heterogeneity: Tau ² =	= 0.02; Cł	$ni^2 = 5.$	= 42%	0.01 0.1 1 10 100			
Test for overall effect:	Z = 0.36	6 (P = C)).72)				Favours [NRT] Favours [ENDS]

Figure 3d Proportion of participants experiencing adverse events

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Study or Subgroup	F	5	NRT	T		Risk Ratio	Risk Ratio
						IV, Random, 95% CI	
Bullen	21	289	17	295	21.6%	1.26 [0.68, 2.34]	
Hajek	79	438	44	446	32.3%	1.83 [1.30, 2.58]	
Hatsukami	25	76	13	76	22.6%	1.92 [1.07, 3.47]	
Lee SH	16	75	21	75	23.4%	0.76 [0.43, 1.34]	
Total (95% CI)		878		892	100.0%	1.39 [0.92, 2.11]	•
Total events	141		95				
Heterogeneity: Tau ² = Test for overall effect:	= 0.11; Ch	$i^2 = 7.3$	74, df = 3	8 (P =	0.05); l ²	= 61%	0.01 0.1 1 10
							Favours [NRT] Favours [ENDS]
Figure 4a Sensitivitige general population		sis—s	Smoking	g ces	sation,	for studies exami	ining smoking cessation in the
general population	n						

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3		ENDS	NRT	Ris	k Ratio	Risk Ratio	
4	Study or Subgroup			Weight IV, Ran		IV, Random, 95% CI	
5	Bullen Lee SM	21 289 5 20			[0.68, 2.34] [0.34, 18.63]		
6		200	205				
7 8	Total (95% CI) Total events	309 26	18	100.0% 1.34	4 [0.74, 2.42]		
9	Heterogeneity: Tau ² =	$= 0.00; Chi^2 = 0$.41, df = 1 (P =	0.52 ; $I^2 = 0\%$	0.0		100
10	Test for overall effect	Z = 0.97 (P =	0.33)		0.0	Favours [NRT] Favours [ENDS	
11	Figure 4b Subgrou	up Analysis-	-Smoking ce	essation. com	paring e-cigare	ettes to nicotine patches o	nlv
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Bullen 21 289 17 295 29.2% 1.26 $[0.68, 2.34]$ Hajek 79 438 44 446 39.7% 1.83 $[1.30, 2.58]$ Lee SH 16 75 21 75 31.1% 0.76 $[0.43, 1.34]$ Total (95% Cl) 802 816 100.0% 1.25 $[0.73, 2.14]$ Heterogeneity: Tau ² = 0.16; Chi ² = 6.85, df = 2 (P = 0.03); l ² = 71% 0.01 0.1 1	Study or Subgroup	ENDS Events Total Ev	NRT vents Total	Weight	Risk Ratio IV, Random, 95% CI	Risk Ratio IV, Random, 95% CI
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	Figure 4c Subgrou	o Analysis— Cor	ntinuous/sı	ustained	l abstinence, 6 m	onths and greater only

Supplementary Material 1 Search strategies

MEDLINE, Embase, CENTRAL

Note: Searches were conducted using an Ovid multi-database search and duplicate records were removed online giving preference to MEDLINE, then Embase, with no field preference. Lines 1-3 are optimized for MEDLINE and the main question constructs are broken out in separate lines for clarity. Lines 4-7 are optimized for Embase and lines 8-10 are optimized for CENTRAL. The next lines isolate the records to the database the search was designed for, combine those sets and then remove duplicate records and final isolate the records from each database again so each can be downloaded and imported into the citation manager using a database-specific import filter.

 Electronic Nicotine Delivery Systems/ or (e cig* or electr* cigar* or electronic nicotine).mp. or (vape or vaper or vapers or vaping or non-combustible nicotine-containing product).ti,ab,kf.
 exp "Tobacco Use Cessation Devices"/ or NRT.ti,ab,kf. or (nicotine adj2 (patch* or gum or nasal spray or mouth spray or mouth strips or lozenge* or tablet* or microtab* or sublingual or replac*)).mp. or (nicotine adj3 therapy).mp.

3. (1 and 2 and ((randomized controlled trial or controlled clinical trial).pt. or randomized.ab. or placebo.ab. or drug therapy.fs. or randomly.ab. or trial.ab. or groups.ab.)) not exp animals/ not humans.sh.

4. Electronic Cigarette/ or (e cig* or electr* cigar* or electronic nicotine).mp. or (vape or vaper or vapers or vaping or non-combustible nicotine-containing product).ti,ab,kw.

5. Nicotine Replacement Therapy/ or NRT.ti,ab,kw. or (nicotine adj2 (patch* or gum or nasal spray or mouth spray or mouth strips or lozenge* or tablet* or microtab* or sublingual or replac*)).mp. or (nicotine adj3 therapy).mp.

6. 4 and 5 and (Crossover-Procedure/ or Double-Blind Procedure/ or Randomized Controlled Trial/ or Single-Blind Procedure/ or (random* or factorial* or crossover* or cross over* or cross-over* or placebo* or (doubl* adj blind*) or (singl* adj blind*) or assign* or allocat* or volunteer*).ti,ab,kw.)

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8. (e cig* or electr* cigar* or electronic nicotine).mp. or (vape or vaper or vapers or vaping or non-combustible nicotine-containing product).ti,ab,kw.

9. NRT.ti,ab,kw. or (nicotine adj2 (patch* or gum or nasal spray or mouth spray or mouth strips or lozenge* or tablet* or microtab* or sublingual or replac*)).mp. or (nicotine adj3 therapy).mp.

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- 13. 10 use cctr
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10	OR microtab OR microtablet OR sublingual) Interventional Studies
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21	Note: As the ICTRP registry has limited search capabilities ³⁵ , only terms related to the
22	intervention were used and protocols with a NCT number were removed from the retrieval, as
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Supplementary Material 2 Abstracted data

The abstracted data included the following:

1- study characteristics:

author names, year of publication, ties with tobacco industry, funding of study, country of study, study setting, study design, number of participating sites, recruitment procedures, enrolment dates, length of study period, random sequence generation, allocation sequence concealment, blinding, methods for preventing and controlling confounding, selection bias, information bias and missing bias, unit of analysis, covariates inclusion, funding, financial and conflict of interest disclosure including ties with industry, inclusion and exclusion criteria, sample size, number of participants that were analyzed, number of participants lost to follow up for each outcome and for the whole study, number of participants at study onset and randomized to each group, and type of analysis (intention to treat vs per protocol)

2- participant characteristics:

age, gender, comorbidities, ethnicities, socio-economic status, income, education, cigarettes smoked per day, Fagerström test for cigarette dependence

3- intervention characteristics:

type, model, brand and generation of ENDS, type and flavor of e-liquid, nicotine content, intervention protocol, length of time ENDS were provided free of charge, frequency of use, duration of intervention, integrity of intervention, description of co-interventions

4- comparator characteristics:

type of nicotine replacement therapy used, dose, frequency of use, nicotine content, control protocol, frequency of use, length of time supplies were provided free of charge, combination of products, frequency of use duration of control, integrity of control, description of co-interventions

5- outcomes:

smoking cessation, method of assessment for smoking cessation used (self-report vs biochemical), smoking abstinence definition, longest time point of smoking cessation, harms assessment, methods of harms assessment, definition of harms, withdrawal symptoms, method of assessment for withdrawal symptoms, reduction in cigarettes smoked, method of assessment of reduction in cigarettes smoked, number of quit attempts, method of quit attempt measurement, acceptance of ENDS/NRT, method of acceptance assessment, method of aggregation used for each outcome, timing of measurement for each outcome, summary data for each outcome, method of aggregation used for each outcome.

Supplementary Table 1a. Characteristics of randomized controlled trials measuring smok cessation at 6 months or later Characteristics of randomized controlled trials measuring				
	n at 6 months or later			
Bullen, 2013				
Methods	Design: 3 parallel groups RCT			
	Recruitment: Participants were recruited via community newspapers			
	inviting people to call the study centre for eligibility pre-screening			
	Setting: one single center in Auckland Australia			
	Inclusion criteria: 18 years of age or older, smoked 10 or more cigare			
	per day for the past year, and wanted to quit smoking.			
	Exclusion criteria: Pregnant or breastfeeding women, people using			
	smoking cessation drugs, those reporting heart attack, stroke, severe			
	angina in the previous 2 weeks, and people with poorly controlled me			
	disorders allergies, or other chemical dependence were excluded			
Participants	Total N: 657 smokers were included in this study, but we only extract			
	584 participants for our review (2 of the 3 groups) as the e-cigarette			
	placebo group did not fit our eligibility criteria.			
	Most participants were women (62%), of a mean age > 40. Approxim			
	one third were of Maori descent, and a little over half had completed			
	grade 12 or above education level. The average daily number of ciga			
	smoked at study onset was around 18, and mean Fagerström test res			
	to 10 scale) for cigarette dependence was > 5.			
Interventions	Randomization: 4:4:1 ratio to nicotine e-cigarettes, nicotine patches			
	placebo e-cigarette group			
	Nicotine e-cigarette group			
	Participants were couriered a first-generation e-cigarette, spare batte			
	and charger, as well as cartridges containing 10 to 16mg of nicotine p			
	mL (although labelled to contain 16 mg), plus simple instructions to u			
	the e-cigarettes as desired from 1 week before until 12 weeks after the			
	chosen quit day. Participants received on average around 20% of the			
	nicotine obtained from cigarette smoking.			
	Nicotine patch group			
	Participants were sent exchange cards in the mail redeemable for nic			
	patches 21 mg from community pharmacies, with instructions to use			
	patches daily, from 1 week before until 12 weeks after their chosen of			
	day. Vouchers were also supplied to participants to cover dispensing			
	costs.			

	Participants in all groups were also referred to telephone-based behavioural support			
Outcomes	Continuous abstinence at 6 months after quit day, defined as self-reported abstinence over the whole follow-up period allowing for 5 or less			
	cigarettes in total, was self-reported, and verified with exhaled breath			
	carbon monoxide of <10 ppm. Harms were both clinically assessed and self-reported, throughout the study period. Withdrawal symptoms were			
	assessed at 1, 3, and 6 months. Reduction in daily cigarettes smoked was			
	measured at 6 months, and acceptance of therapy was measured at 1 and 6 months.			
Notes	Some of this study's authors reported ties to e-cigarette manufacturers,			
	and smoking cessation drug companies			
Hajek, 2019				
Methods	Design: 2 parallel groups RCT			
	Recruitment: Participants were recruited through stop smoking services,			
	which included trial information in their advertising. Participants were			
	also recruited through social media, and leaflets advertising the trial were			
	delivered to local households.			
	Setting: 3 sites in the United Kingdom			
	Inclusion criteria: Adults, with no strong preference towards e-cigarette			
	or NRT, who were not using either type of product at the time of study			
	enrolment			
	Exclusion criteria: Pregnant women or breastfeeding women			
Participants	Total N: 884 participants were included in this study			
	Median age for both groups was 41, and women comprised 48% of			
	participants. Most participants were White British, and the majority had			
	post-secondary education. Median daily number of cigarettes smoked at			
	study onset was 15, and mean Fagerström test result for cigarette			
	dependence was 4.5 in the e-cigarette group and 4.6 in the NRT group.			
Interventions	Randomization: nicotine-containing e-cigarettes of varying doses, and an			
	choice of a list of NRT, in a 1:1 ratio			
	E-cigarette group			
	Participants were provided with a starter pack called One Kit, which			
	included an atomizer, a battery, and one 30 mL bottle of Tobacco Royale			
	flavor e-liquid. Participants were asked to purchase their future e-liquid			
	online or from local vape shops and to buy a different e-cigarette device i			
	the one supplied did not meet their needs. They were encouraged to			
	experiment with e-liquids of different strengths and flavors. Those who			
	were unable to obtain their own supply were provided with one further			
	10-ml bottle, but this was not offered proactively. Participants received			
	oral and written information on how to operate the e-cigarette.			

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	NRT group
	Participants were informed about the range of nicotine-replacement
	products (patch, gum, lozenge, nasal spray, inhalator, mouth spray, mouth
	strip, and microtabs) and selected their preferred product. Use of
	combinations was encouraged, typically the patch and a faster-acting oral
	product. Participants were also free to switch products.ps
	Both groups
	Participants in both groups were offered multisession behavioral support
	as per UK stop smoking service practice, involving weekly one on one session with local clinicians.
	Participants were also asked to sign a commitment to not use the
	unassigned treatment for 4 weeks
Outcomes	Continuous abstinence at 52 weeks after guit day, defined as self-reported
	abstinence over the whole follow-up period allowing for 5 or less
	cigarettes in total, was self-reported, and verified with exhaled breath
	carbon monoxide of <8 ppm. Harms were self-reported throughout the
	study period. Withdrawal symptoms were assessed at 1 and 4 weeks in
	abstainers. Reduction in daily cigarettes smoked was also measured at 52
	weeks, as well as acceptance of e-cigarettes and NRT
Notes	Some of this study's authors reported ties to smoking cessation drug
	companies.
Lee SH, 2019	
	Design: 2 parallel groups RCT
Methods	Recruitment: Participants were recruited from a motor company in the
	Republic of Korea.
	Setting: One site in Cheonan, Republic of Korea
	Inclusion criteria: Participants were adults 18 years and above, male, who
	smoked at least 10 cigarettes per day in the preceding year, and who were
	smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette
	smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption
	smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical
	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the
	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT.
Participants	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study
Participants	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study Mean age was 42 years and all participants were men. Almost 40% had
Participants	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study Mean age was 42 years and all participants were men. Almost 40% had post-secondary education. Median daily number of cigarettes smoked at
Participants	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study Mean age was 42 years and all participants were men. Almost 40% had post-secondary education. Median daily number of cigarettes smoked at study onset was 1 pack per day, and mean Fagerström test result for
	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study Mean age was 42 years and all participants were men. Almost 40% had post-secondary education. Median daily number of cigarettes smoked at study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4.
Participants Interventions	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study Mean age was 42 years and all participants were men. Almost 40% had post-secondary education. Median daily number of cigarettes smoked at study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4. Randomization: nicotine-containing e-cigarettes, and nicotine gum in a
	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study Mean age was 42 years and all participants were men. Almost 40% had post-secondary education. Median daily number of cigarettes smoked at study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4.
	 smoked at least 10 cigarettes per day in the preceding year, and who were motivated to stop smoking entirely or to reduce their cigarette consumption Exclusion criteria: Participants were excluded if they had a past medical history of serious clinical diseases or had attempted to stop smoking in the last 12 months by using other NRT. Total N: 150 participants were included in the study Mean age was 42 years and all participants were men. Almost 40% had post-secondary education. Median daily number of cigarettes smoked at study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4. Randomization: nicotine-containing e-cigarettes, and nicotine gum in a

	Participants received a 24-week supply of e-cigarettes eGo-C Ovale, Janty-Korea Co., Janty-Asia Co., Seoul, Republic of Korea, nicotine 0.01 mg/mL.
	Nicotine gum group Participants received a 24-week supply of nicotine gum Nicoman, Daewoog Pharmaceutical, Seongnam, Republic of Korea, 2 mg/tablet
	Both groups Participants in both groups were offered 55-minute education sessions or smoking cessation aids
Outcomes	Continuous abstinence was defined as abstinence from smoking from 9 to 24 weeks, validated with end-expiratory carbon monoxide (<10 ppm) and a negative urine cotinine result. Harms were self-reported throughout the study period. Reduction in daily cigarettes smoked was also measured at 24 weeks.
Notes	None of the study authors were found to have ties to industry.
Lee SM, 2018	
Participants	 Design: 2 parallel groups RCT Recruitment: Participants were recruited from an anesthesia preoperative clinic for elective surgery. Setting: San Francisco Veterans' Affairs Medical Center, affiliated with the University of California in San Francisco United States of America Inclusion criteria: Participants were eligible if they presented to the clinic 3 or more days prior to elective surgery, smoked more than two cigarette per day, and had smoked at least once in the last 7 days Exclusion criteria: Participants were excluded if they exclusively used other forms of tobacco (e.g. pipe tobacco) or marijuana only, were pregnant or breastfeeding, had an unstable condition, were using smoking cessation trial, or currently used e-cigarettes daily. Total N: 30 participants were included in this study
	Most participants were men (90%) in their 50's. Some had comorbidities including diabetes, hypertension, heart disease, and chronic obstructive pulmonary disease. Most were Caucasians. The average daily number of cigarettes smoked at study onset was 15.3 in the e-cigarette group, and 10.8 in the NRT group, and the mean Fagerström test result for cigarette dependence was 3.7 in the e-cigarette group and 2.5 in the NRT group.
Interventions	 Randomization: e-cigarettes and nicotine patches in a 2:1 ratio E-cigarette group Participants received a 6-week supply of NJOY e-cigarettes (Scottsdale, AZ USA), a disposable first-generation e-cigarette that is available in shops and online. They were issued a number of e-cigarettes corresponding to

	 the reported baseline cigarettes smoked per day, calculated assuming or NJOY e-cigarette was equivalent to 10 cigarettes. Participants were instructed to smoke bold (4.5%) e-cigarettes ad libitum for 3 weeks, then the Gold (2.4%) e-cigarettes ad libitum for 2 weeks, and then the Study (0%) e-cigarettes ad libitum for the final week. Nicotine patch group Participants randomized to the nicotine patches group were given a 6-week supply of Nicoderm CQ patches (5 weeks) and placebo patches (1 week) appropriate to baseline nicotine consumption. Those smoking an
	average of ten or more cigarettes per day were given a 21 mg/day patch for 3 weeks, a 14 mg/day patch for 1 week, a 7 mg/day patch for 1 week, and a 0 mg/day patch for 1 week. Participants who reported smoking an average of fewer than 10 cigarettes per day at baseline were given a 14 mg/day patch for 3 weeks, a 7 mg/day patch for 2 weeks, and a 0 mg/day patch for 1 week.
	Both groups Participants in both groups were given referral California Smokers' Helpline and were asked to refrain from the use of cigarettes during the study period.
Outcomes	Smoking cessation at 6 months was self-reported through 7-day point- prevalence abstinence and verified with exhaled breath carbon monoxid of <10 ppm. Harms and withdrawal symptoms were systematically collected at 8 weeks. Reduction in daily cigarettes smoked was also measured at 6 months, as well as acceptance of e-cigarettes and NRT.
Notes	None of the study authors were found to have ties to industry.

Supplementary Table 1b. Characteristics of randomized controlled trial measuring smoking cessation earlier than 6 months

Hatsukami, 2019	
Methods	Design: 4 parallel groups RCT
	Recruitment: Participants were culled from two sets of studies, one of
	which also included two groups randomized to snus (spitless smokeless
	tobacco); one was complete substitution with snus, and the other was ad
	libitum use. Due to recruitment challenges, the two snus groups were
	dropped midway through the study, resulting in four experimental groups:
	ad libitum use of e-cigarettes (participants may smoke as many cigarettes
	as they like), complete substitution with e-cigarettes (aiming for smoking
	as they like), complete substitution with e-cigarettes (aiming for smoking

	NRT group
	Participants randomized to this group used Vuse Solo, manufactured by Reynolds Inc as the primary e-cigarette. Early in the study, Blu e-cigarett (cartridge-based system) and Fin (prefilled tanks system) were used, bu Vuse attained the highest market share early on so the study switched exclusively to Vuse. E-cigarettes with a 4.8% nicotine concentration were provided to participants free of charge for 8 weeks, as well as 7 cartridg weekly, with the option of returning to the clinic to obtain additional cartridges if needed. Tobacco, menthol, mint, and berry flavors were available.
	E-cigarette group
Interventions	3. Randomization: e-cigarettes and nicotine gum or lozenges
	was 15, and median Fagerström test result for cigarette dependence wa
	Most participants were White, and the majority had post-secondary education. The median daily number of cigarettes smoked at study onse
	Median age was 47 years, and women comprised 49% of participants.
	participants), as the other two groups did not fit our eligibility criteria.
	cigarette group, and complete substitution with NRT group (152
raiticipalits	review were only extracted from the complete substitution with e-
Participants	become pregnant, or breastfeeding Total N: 264 participants were included in the study, but data for this
	NRT or other cessation medication, or if they were pregnant or planning
	conditions affecting results of biomarker analyses, were currently using
	planning to quit smoking in the next 3 months, suffered from chronic
	problems, regular use of other nicotine or tobacco products, were
	attempt in the past 3 months, recent (<3 months) alcohol or drug abuse
	Exclusion criteria: Participants were excluded if they had a serious quit
	of at least 10 ppm or a NicAlert test = level 6, and in stable physical and mental health.
	smoked at least 5 cigarettes per day with a breath carbon monoxide tes
	Inclusion criteria: Participants were adults at least 18 years of age,
	United States of America
	State University, Columbus, OH; Roswell Park Cancer Center, Buffalo, NY
	Settings: 3 sites, University of Minnesota, Twin Cities (lead site); The Oh
	exposure to harmful tobacco smoke.
	smokers who were interested in trying a product that may reduce
	institutions. The advertisements stated that a study was recruiting
	Participants were recruited through various media outlets across three
	cessation), complete substitution with NRT, continued smoking with usu brand of cigarettes.

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	Participants could choose between mint, cinnamon or fruit-flavore
	nicotine gum or nicotine lozenge, at a dose of 4 mg. If adverse eff
	were recorded, the dose was decreased to 2 mg.
	Both groups
	After randomization, participants were asked to complete daily di interactive voice recording to chart the number of cigarettes smol as well as document assigned product use for the duration of the Participants received a monetary bonus if they complied with the protocol; this included keeping an accurate record of product use,
	completing the daily diaries, and returning unused products. They a bonus payment if they had a carbon monoxide level \leq 4 ppm at a visit. Participants also received a brief counseling session on how smoking.
Outcomes	Smoking cessation was determined by 7-day point prevalence at 8 mainly through biochemical verification but also by self-report Rec in daily cigarettes smoked was also measured at 8 weeks, as well a acceptance of e-cigarettes and NRT.
	Harms were assessed systematically at 20 weeks, 12 weeks after t of the study period. Withdrawal symptoms were assessed at week
	4, 6, and 8.
Notes	One of the study authors is a member of the FDA Tobacco Product
	Scientific Advisory Committee and another one has served as an e
	witness in tobacco company litigation.
Eisenhofer,	witness in tobacco company litigation.
outcomes Eisenhofer, 2015	witness in tobacco company litigation. Table 1c. Characteristics of randomized controlled trial measuring ot
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outcomes Eisenhofer, 2015	witness in tobacco company litigation. Table 1c. Characteristics of randomized controlled trial measuring ot Design: 2 parallel groups RCT Recruitment: Not specified Setting: Not specified Inclusion criteria: Veterans who met criteria for tobacco disorder a the DSM Exclusion criteria: Not specified Total N: 11 participants were included
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Dutcomes Eisenhofer, 2015 Methods	witness in tobacco company litigation. Table 1c. Characteristics of randomized controlled trial measuring ot Design: 2 parallel groups RCT Recruitment: Not specified Setting: Not specified Inclusion criteria: Veterans who met criteria for tobacco disorder a the DSM Exclusion criteria: Not specified Total N: 11 participants were included Mean age was 52, and 82% were males. The vast majority of parti were African American. The average daily number of cigarettes sn
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	Participants received nicotine-containing e-cigarettes with 16 mg of nicotine per cartridge
	NRT group Participants received nicotine patch 16 mg daily
	Both groups All participants were instructed to smoke ad libitum during week 1, and to smoke as little as possible during week 3.
Outcomes	Reduction in cigarettes smoked per day was self-reported at 3 weeks and compared to week 1. Withdrawal symptoms were compared between week 1 and week 3.
Notes	This study was available as an abstract only therefore limited details are available.

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Supplementary Material 4 Details on Risk of Bias Assessment for each outcome of interest

Supplementary Table 2. Detailed description of concerns for each domain marked identified as "some concerns" or "high risk" on Risk of Bias Assessment

	Randomization Process	Deviations from intended intervention	Missing of outcome data	Measurement of the outcome	Selection of the reported result
Bullen 2013	Low risk	Adherence higher in the ENDS group compared to NRT group at all timepoints. At 6 months, 29% of ENDS group vs 8% of NRT group still using assigned treatment.	Low risk	Low risk	Low risk
Hajek 2019	Low risk	At 52 weeks among participants with 1-year abstinence, 80% were using e- cigarettes in the ENDS group vs 9% in the NRT group. Also, 6% of participants in the ENDS group reported using non-allocated NRT for at least five consecutive days in the past six months compared to 22% in the NRT group that reported using non-allocated product	Low risk	Low risk	Low risk
Hatsukami 2019	No information provided with regards to randomization process and allocation concealment. However, there were no	The NRT group had the highest dropout rates compared to the other groups in the study. At 8 weeks, 24% dropped out in the ENDS group compared to 30% in the NRT group.	Large number of dropouts; participants who did not stop smoking could be less motivated to continue with study follow up	Low risk	Low risk

	significant baseline differences between groups				
Lee, SH 2019	The use of constant block sizes of 2 makes it easy to determine order of randomization.	No participants discontinued the intervention. However, 4 and 14 participants in the ENDS and NRT group dropped out before treatment, respectively.	Although data was missing for 12% of randomized individuals, all dropouts occurred prior to the start of treatment. Missingness in this case less likely to be due to the value of the outcome as it happened prior to onset of therapy	Low risk	Low risk
Lee, SM 2018	Low risk	Low risk	Low risk	Low risk	Low risk
	luction outcome				
Smoking red Bullen 2013	Low risk	Refer to smoking cessation outcome	Sensitivity analyses conducted for the smoking cessation outcome were not performed for the smoking reduction outcome	Low risk	Low risk
	1	Refer to smoking cessation outcome Not enough information available in abstract	conducted for the smoking cessation outcome were not performed for the smoking reduction	Low risk	Low risk Not enough informatic available in abstract
Bullen 2013 Eisenhofer	Low risk Not enough information available in	Not enough information available in	conducted for the smoking cessation outcome were not performed for the smoking reduction outcome Not enough information		Not enough information
Bullen 2013 Eisenhofer 2015 Hajek 2019 Hatsukami 2019	Low risk Not enough information available in abstract	Not enough information available in abstract	conducted for the smoking cessation outcome were not performed for the smoking reduction outcome Not enough information available in abstract	Low risk	Not enough informatio available in abstract
Bullen 2013 Eisenhofer 2015 Hajek 2019 Hatsukami	Low risk Not enough information available in abstract Low risk Refer to smoking	Not enough information available in abstract Refer to smoking cessation outcome	conducted for the smoking cessation outcome were not performed for the smoking reduction outcome Not enough information available in abstract	Low risk Low risk	Not enough informatio available in abstract

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Bullen 2013	Low risk	Differences in treatment adherence	No information on the	high likelihood that	Low risk
		could potentially lead to	proportion of	participants who were	
		discrepancies in harm reporting	participants on whom	unhappy with their	
			adverse events were	treatment allocation	
			collected; it is likely that	would report side	
			people who experienced	effects more often	
			more severe side effects	than their	
			did not continue with	counterparts.	
			study follow-up activities		
Hajek 2019	Low risk	Differences in treatment adherence	The authors reported	High likelihood that	Low risk
		could potentially lead to	harm data based on	participants who were	
		discrepancies in harm reporting	number of participants	unhappy with their	
			at randomization,	treatment allocation	
			however significant	would report side	
		Deer.	dropout seen at 4-week	effects more often	
			follow up, raising	than their counterparts	
			concerns that adverse		
			event data not collected		
			on all participants		
Hatsukami	Refer to smoking	Differences in treatment adherence	No information on the	High likelihood that	Low risk
2019	cessation outcome	could potentially lead to	proportion of	participants who were	
		discrepancies in harm reporting	participants on whom	unhappy with their	
			adverse events were	treatment allocation	
			collected; it is likely that	would report side	
			people who experienced	effects more often	
			more severe side effects	than their counterparts	
			did not continue with		
			study follow-up activities		
Lee, SH 2019	Refer to smoking	Differences in treatment adherence	Low risk	High likelihood that	Low risk
	cessation outcome	could potentially lead to		participants who were	
		discrepancies in harm reporting		unhappy with their	
		however non-adherence happened		treatment allocation	
		prior to onset of treatment,		would report side	
		therefore less likely to have an		effects more often	
		impact		than their counterparts	
Lee, SM 2018	Low risk	Low risk	Low risk	High likelihood that	Low risk
				participants who were	
				unhappy with their	

Withdrawol	symptoms outco			treatment allocation would report side effects more often than their counterparts	
Eisenhofer 2015	Not enough information available in abstract	Not enough information available in abstract	Not enough information available in abstract	Not enough information available in abstract	Not enough information available in abstract
Hajek 2019	Low risk	Differences in treatment adherence could potentially lead to discrepancies in withdrawal symptoms reporting	Outcome not available for all randomized participants; likely that people who experienced more nicotine withdrawal symptoms did not continue with study follow-up activities	Given that the withdrawal measurements were self-reported, there is a high likelihood that participants who were unhappy with treatment allocation reported more withdrawal symptoms than their counterparts	Low risk
Hatsukami 2019	Refer to smoking cessation outcome	Differences in treatment adherence could potentially lead to discrepancies in withdrawal symptoms reporting	Outcome not available for all randomized participants; likely that people who experienced more nicotine withdrawal symptoms did not continue with study follow-up activities	Given that the withdrawal measurements were self-reported, there is a high likelihood that participants who were unhappy with treatment allocation reported more withdrawal symptoms than their counterparts	No information on how withdrawal symptom assessment was performed
Lee, SM 2018	Low risk	Low risk	Low risk	Given that the withdrawal measurements were self-reported, there is a high likelihood that participants who were	Low risk

Accentance	of therapy out	come		than their counterparts	
Bullen 2013	Low risk	Differences in treatment adherence could potentially lead to discrepancies in acceptance of therapy outcome	Participants unhappy with their assigned therapy likely did not continue with study follow-up activities	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk
Hajek 2019	Low risk	Differences in treatment adherence could potentially lead to discrepancies in acceptance of therapy outcome	Participants unhappy with their assigned therapy likely did not continue with study follow-up activities	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk
Hatsukami 2019	Not enough information available in abstract	Differences in treatment adherence could potentially lead to discrepancies in acceptance of therapy outcome	Participants unhappy with their assigned therapy likely did not continue with study follow-up activities	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk
Lee, SM 2018	Low risk	Low risk	Low risk	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk





PRISMA 2009 Checklist

Section/topic	#	Checklist item	Reported on page #
TITLE		·	
Title	1	Identify the report as a systematic review, meta-analysis, or both.	0
ABSTRACT		·	
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.	1
INTRODUCTION			
Rationale	3	Describe the rationale for the review in the context of what is already known.	4
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).	4
METHODS			
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.	5
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.	5
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.	5,6
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.	5,6, Supp material 1
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).	6
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.	6
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.	6, Supp material 2
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.	6
Summary measures	13	State the principal summary measures (ajgperist ration difference in omeans) es. xhtml	6

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PRISMA 2009 Checklist

4 Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I ²) for each meta-analysis.	7
6 7	-	Page 1 of 2	
8 9 Section/topic	#	Checklist item	Reported on page #
1 Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).	6
13 Additional analyses 14	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.	7
16 RESULTS			
17 Study selection 18	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.	7, Figure 1
20 Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.	7,8, Supp material 3
22 23 Risk of bias within studies 24 25	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).	9, Figures 2a,b,c,d,e
 ²⁶ Results of individual studies 27 28 29 	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.	9-13, Figures 3a,b,c,d,
30 Synthesis of results 31 32 33	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.	9-12, Figures 3a,b,c,d
34 Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).	13
35 36 37 37 38	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).	13, Figures 4a,b,c
41 Summary of evidence 42	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).	13-15
⁴³ Limitations 44 45	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias), identified research, reporting bias), For peer review only - http://bmjopen.bmj.com/site/about/guidelines.xhtml	13-15
46 47			



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4 (5 (Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.	13-15
6	FUNDING		·	
8	Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.	0
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28 29 30 31 32 33 34 35 36 37 38 39				
40 41 42 43 44 45 46 47			For peer review only - http://bmjopen.bmj.com/site/about/guidelines.xhtml	

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Smoking cessation in individuals who use vaping as compared to traditional nicotine replacement therapies; a systematic review and meta-analysis

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Smoking cessation in individuals who use vaping as compared to traditional nicotine replacement therapies; a systematic review and meta-analysis

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Contributors statements:

Dr Pound conceptualized and designed the study, carried out the analyses, interpreted the data, drafted the initial manuscript, reviewed, and revised the manuscript.

Mrs Zhang participated in the conceptualization and design of the study, carried out the analyses, interpreted the data, participated in drafting the initial manuscript, reviewed and revised the manuscript.

Ms Kodua participated in the conceptualization and design of the study, and reviewed the manuscript.

Dr Sampson participated in the design of the study, developed the search strategies, reviewed, and revised the manuscript.

All authors approved the final manuscript as submitted and agree to be accountable for all aspects of the work.

ABSTRACT

Objectives: Despite the aggressive marketing of electronic nicotine device systems (ENDS) as smoking cessation tools, the evidence of their effectiveness is mixed. We conducted a systematic review of randomized controlled trials to determine the effect of ENDS on cigarette smoking cessation, as compared to other types of nicotine replacement therapies (NRT).

Design: Systematic review and meta-analysis using the Grading of Recommendations Assessment, Development, and Evaluation (GRADE) approach.

Data sources: MEDLINE, Embase, the CENTRAL Trials Registry of the Cochrane Collaboration using the Ovid interface, ClinicalTrials.gov and the World Health Organization International Clinical Trials Registry Platform trials registries were searched through June 17th 2020.

Eligibility criteria for studies: Randomized controlled trials in which any type of ENDS was compared to any type of NRT, in traditional cigarette users.

Data extraction and synthesis: The primary outcome was smoking cessation, defined as abstinence from traditional cigarette smoking for any time period, as reported in each included study, regardless of whether abstinence is self-reported or biochemically validated. Secondary outcomes included smoking reduction, harms, withdrawal, and acceptance of therapy. A random-effect model was used, and data were pooled in meta-analyses where appropriate.

Results: Six studies were retained from 270. Most outcomes were judged to be at high risk of bias. The overall quality of evidence was graded as 'low' or 'very low'. Pooled results showed no difference in smoking cessation (RR 1.42 [0.97, 2.09]), proportion of participants reducing smoking consumption (RR 1.25 [0.79, 1.98]), mean reduction in cigarettes smoked per day (MD 1.11 [-0.41, 2.63]), or harms (RR 0.96 [0.76, 1.20]), between groups.

Conclusion: We found no difference in smoking cessation, harms, and smoking reduction between e-cigarette and NRT users. However, the quality of the evidence was low. Further research is needed before widespread recommendations are made with regards to the use of ENDS.

Systematic review registration number: protocol registered with the International Prospective Register of Systematic Reviews (PROSPERO) on February 27th, 2020. Registration number pending.

Strengths and limitations of this study

- This study provides up to date meta-analyses of direct comparisons of vaping with nicotine replacement therapy for smoking cessation, studied through randomized controlled trials.
- We examined harms associated with vaping, which are becoming increasingly concerning.

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- This study makes extensive efforts to obtain unreported data from investigators. •
- Careful consideration is given to the potential impact of risk of bias and methodological heterogeneity.
- As we included only RCTs, many studies that used weaker study designs were ineligible

Abstract word count: 298 words Text word count: 4917 words

INTRODUCTION

Background

Despite a significant lack of rigorous pharmacological testing, the use of electronic nicotine device systems (ENDS), otherwise known as vaping devices, has been aggressively marketed as an effective method to quit smoking. In Canada, 32% of current and former smokers report having used ENDS as a smoking cessation aid.¹ In addition to delivering nicotine to the user, ENDS are thought to replace some of the habitual behaviours and sensations associated with smoking, such as the action of bringing a cigarette to the mouth. By doing so, ENDS may provide coping mechanisms that other traditional nicotine replacement therapies (NRT) do not offer, and therefore may help with the behavioural component of smoking reduction and cessation.² While vaping is believed to be less harmful than cigarette smoking, a large number of emerging reports on the health impacts of vaping are worrisome. In addition, the evidence on the effectiveness of ENDS as a smoking cessation aid is mixed.

In 2016, a meta-analysis of 20 studies found that people using ENDS had a 28% reduction in the odds of stopping cigarette smoking as compared to those not using ENDS.³ However, in a 2019 recent randomized controlled trial (RCT), individuals randomized to nicotine-containing e-cigarettes were more likely to abstain from smoking at one year compared to individuals randomized to nicotine patches (18% compared to 9.9%, RR 1.83; 95% CI 1.30 to 2.58).⁴ A Cochrane review⁵ found that nicotine-containing e-cigarettes were more effective than non-nicotine containing e-cigarettes for smoking cessation, but was not able to compare ENDS products to traditional NRT.

Little information is known about the long-term health impacts of ENDS. Reports of acute toxicity have recently captured the public's attention. In late 2019 and early 2020, "e-cigarette, or vaping, product use-associated lung injury" (EVALI) caused 2807 illnesses and 68 deaths in the US,⁶ and 19 cases in Canada.⁷ Other short-term adverse events reported with the use of ENDS include cardiovascular changes such as increased heart rate and blood pressure, cough, wheeze,⁸ and mucus production.⁹ Burn injuries have also been reported, as well as fatalities from drinking or injecting the e-liquid.⁸

There is no long-term data available on the relationship between ENDS and oral, respiratory, and cardiovascular health, as well as cancer. There is however available data linking the chemicals present in e-liquids with cellular DNA damage and carcinogenicity.^{9,10} There is some evidence that the use of ENDS is associated with asthma exacerbations.¹¹ No human long-term data exist on the use of ENDS in pregnancy and their impact on the developing fetus.

Given the large number of smokers using ENDS as a potential smoking cessation tool, there is a need to review and synthesize the evidence of trials examining a head to head comparison of ENDS versus traditional NRT for smoking cessation.

Objective

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The objective of this review is to systematically review the evidence found in RCTs to determine the effect of electronic nicotine delivery systems (ENDS) on cigarette smoking cessation in smokers, as compared to other types of nicotine replacement therapies (NRT).

METHODS

Protocol and registration

The protocol for this systematic review was submitted to International Prospective Register of Systematic Reviews (PROSPERO) on February 27th, 2020 (registration pending) and uploaded as a preprint on Open Science Framework (OSF) Preprints on May 12th 2020.¹²

Patient and public involvement

No patient involved

Criteria for study inclusion

Study Characteristics:

RCTs in which ENDS were compared to non-electronic NRT in smokers were included. We restricted our inclusion to RCTs to minimize the risk of bias. No language limits were imposed. No date limits were imposed either, although we did not anticipate studies published prior to 2003, since this is when the first e-cigarette was invented.¹³ There was no geographical restriction of studies.

Study Population:

All traditional cigarette users were included, regardless of age, amount of traditional cigarette use, and motivation to quit.

Intervention of interest:

The intervention of interest comprised all types, models, and brands of ENDS.

<u>Comparators:</u>

All included studies compared ENDS with non-electronic NRT. NRT comprised, but were not limited to, nicotine patch, gum, lozenge, nasal spray, inhalator, mouth spray, mouth strips, microtabs, and combination of products.

Outcome measures:

The primary outcome measure is traditional cigarette smoking cessation defined as abstinence from traditional cigarette smoking for any time period, as reported in each included study, regardless of whether abstinence is self-reported or biochemically validated. Secondary outcomes include reduction in the number of traditional cigarettes smoked in any given time period, adverse events, withdrawal symptoms, and participants' acceptance of therapy. We had planned on collecting quit attempts information but none of the studies reported on this outcome.

Settings:

All health care and community settings were included.

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Study Identification

The following databases were searched through June 17th 2020: MEDLINE (1946 to June 2020), Embase (1947 to June 2020) and the CENTRAL Trials Registry of the Cochrane Collaboration (May 2020 Issue) using the Ovid interface. The MEDLINE search was limited using the Cochrane Highly Sensitive Search Strategy and the Embase search was limited using the recommended limit for controlled trials.¹⁴ Searches were developed by a librarian experienced in systematic reviews, using a method designed to optimize term selection.¹⁵ ClinicalTrials.gov and World Health Organization International Clinical Trials Registry Platform (WHO ICTRP) trials registries were searched for registered intervention studies, regardless of their completion status. Electronic search strategies are presented in Supplementary Material 1. The reference lists of included studies and any applicable review studies were searched.

Authors of protocols identified through registries were contacted electronically, to request data for the review. In addition, clinical experts in the field of vaping and smoking cessation were contacted to enquire about any unpublished research fulfilling our inclusion criteria.

Selection of Studies

Records retrieved by the electronic search were downloaded and imported into a Reference Manager database for duplicate removal, and then uploaded to Covidence. Throughout the review, newly identified records were integrated into the set for screening.

Each title and abstract was independently screened by two review authors (from CP, JZ, and ATK) against the eligibility criteria.¹⁴ Full text of all studies deemed potentially eligible was obtained and reviewed independently by two of the same review authors to determine eligibility. For screening, data extraction, and risk of bias assessment, disagreements were resolved by discussion, and with a third reviewer when needed.

Data extraction and management

For studies that fulfilled the inclusion criteria, two reviewers (CP, JZ) extracted the data into an electronic data collection form, which was piloted by both reviewers (Supplementary Material 2). The data collection was revised, based on feedback from the reviewers. Study authors were contacted electronically to obtain relevant but unavailable data.

Risk of bias assessment for included studies

Two reviewers (CP, JZ) independently conducted the risk of bias assessment for each study at the outcome level using the Revised Cochrane risk-of-bias tool for randomized trials (RoB 2).¹⁶

Measures of treatment effect

Dichotomous data was analyzed by calculating the prevalence rate ratio, using the longest follow-up time reported, as well as the 95% confidence interval. The prevalence rate ratio (RR) for smoking cessation was calculated as such:

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 $RR = \frac{N \text{ of subjects abstaining from smoking in intervention}}{N \text{ of subjects in intervention}}$

N of subjects abstaining from smoking in control
$$/N$$
 of subjects in control

Continuous data for the secondary outcomes were analyzed through mean differences between groups as the same scales were used. In the case of studies with multiple arms, we only extracted data for the groups relevant to this review.

Data synthesis

We provide a synthesis of the included studies (Table 1). Where appropriate, data have been pooled for meta-analyses, and random effects were used for all analyses in RevMan.¹⁴ The inverse-variance random-effects and the mean difference approach (using standard deviations and sample sizes) were used for dichotomous and continuous outcomes, respectively, to assign the weight given to each study. Participants with missing data were considered as still smoking.⁵ The proportion of adverse events reported was based on the number of people available for outcome assessment. For the reduction of the number of cigarettes smoked, missing values were assumed to be zero.

Assessment of heterogeneity

A p value of 0.10 for the chi-squared test (Cochrane Q) and an I² value of >50% were used as indicators of substantial heterogeneity. This however needs to be interpreted with caution given the small number of studies available for the meta-analysis. Clinical and methodological diversity was also explored.

We planned to assess reporting/publication bias using funnel plots of effect estimate against standard error, and testing for funnel plot asymmetry, however, the number of included studies was too low (<10).

We also planned on conducting a number of sensitivity analyses to determine the robustness of the results of the meta-analyses; subgroup analyses to investigate potentially modifying factors such as age and smoking intensity; as well as meta-regression to study the impact of covariates such as motivation to quit smoking, provision of training, and other factors,¹⁷ but minimum data thresholds were not met.

We present a 'Summary of Findings' table (Table 2) for all outcomes. We used the five GRADE considerations (study limitations, consistency of effect, imprecision, indirectness, and publication bias)¹⁴ to assess the quality of evidence for each outcome and to draw conclusions about the robustness of evidence within this review.

RESULTS

Our initial bibliographic search yielded 270 records, and after screening and full-text review, we retained 6 RCTs. An updated search conducted in June 2020 yielded an additional 116 records (for a total of 386 records), none of which were included after screening (Figure 1).

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We identified six RCTs (Bullen 2013,¹⁸ Eisenhofer 2015,¹⁹ Hajek 2019,⁴ Hatsukami 2019,²⁰ Lee SH 2019,²¹ Lee SM 2018²²). Of these, five contributed data to our primary outcome of smoking cessation.^{4,18,20-22} Four studies^{4,18,21,22} examined cessation at 6 months or longer, while one²⁰ examined short term cessation (< 6 months). Table 1 includes the salient features of the included studies. A more detailed description of included studies can be found in Supplementary Material 3.

Author and	Design	Country	Number of	Main	Intervention	Comparator	Main
year of publication			participants	eligibility criteria			outcome of interest
Bullen,	3-group,	Australia	657 total,	\geq 18 years,	First-	Nicotine	Continuous
2013 ¹⁷	parallel,		584 included	smoked \geq	generation e-	patch x 12	abstinence
	single		in this	10	cigarette x 12	weeks	6 months
	center		review (2 of	cigarettes	weeks		after quit
			3 groups)	per day in			day
				the past			
				year,			
				motivated			
				to quit			
Hajek	2-group,	United	884	Adults	Any type of	Any	Continuous
2019 ⁴	parallel,	Kingdom		with no	e-cigarette	nicotine-	abstinence
	multi-	_		strong	_	replacement	52 weeks
	centre			preference		therapy	after quit
				towards e-			day
				cigarette or			
				NRT			
Lee SH,	2-group,	Republic	150	\geq 18 years,	e-cigarette x	Nicotine gum	Continuous
2019 ²⁰	parallel,	of Korea		smoked \geq (24 weeks	x 24 weeks	abstinence
	single			10			24 weeks
	center			cigarettes			after quit
				per day in			day
				the past			
				year,			
				motivated			
				to quit	·		
Lee SM,	2 group,	USA	30	Adults,	e-cigarette x	Nicotine	7-day poin
2018 ²¹	parallel,			smoked \geq	6 weeks	patch x 5	prevalence
	single			2		weeks, then	abstinence
	center			cigarettes		placebo	at 6 month
				per day in		patch x 1	
				the past		week	
				year,			
				smoked at			
				least once			
				in last 7			
				days			

Table 1. Characteristics of included studies

Characteristics of RCT measuring smoking cessation earlier than 6 months

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Hatsukami, 2019 ¹⁹	4 group, parallel, multi- center	USA	264 total, 152 included in this review (2 of 4 groups)	\geq 18 years, smoked \geq 5 cigarettes per day	e-cigarettes	Nicotine gum or nicotine lozenge	7-day point prevalence abstinence at 8 months
<u>a</u>		_					
Characteris Eisenhofer,	2-group,	T measur USA	ing other out	comes Veterans	e-cigarettes x	Nicotine	Reduction

Risk of bias in included studies

We assessed risk of bias for each included study. A detailed report of the risk of bias assessment can be found in Supplementary Material 4.

Figure 2 illustrates the risk of bias for each outcome.

Effect of Interventions

Smoking cessation

Five of the six studies reported on smoking cessation.^{4,18,20-22} When comparing e-cigarettes to NRT in the context of smoking cessation, there was no significant difference between groups in verified self-reported continuous abstinence at 6 months (21/289 vs 17/295, RR 1.26 [0.68, 2.34], p=0.46) in the Bullen 2013¹⁸ study, and in continuous abstinence from 9 to 24 weeks (16/75 vs 21/75, RR 0.76 [0.43, 1.34], p = 0.344) in the Lee SH 2019²¹ study. In addition, the Lee SM 2018²² study showed no difference between groups for the 7-day point prevalence abstinence at 6 months in the context of perioperative smoking cessation (5/20 vs 1/10, RR 2.50 [0.34, 18.63], p = 0.63).

In the Hajek 2019⁴ study, self-reported, verified continuous abstinence at 1 year was found to be higher in the e-cigarette group (79/438 vs 44/446, RR 1.83 [1.30, 2.58], P<0.001), and smoking cessation assessed by 7-day point prevalence at 8 weeks in the Hatsukami 2019²⁰ trial was also higher in the e-cigarette group (25/76 vs 13/76, RR 1.92 [1.07, 4.37], p = 0.039).

We combined data from all 5 studies comparing smoking cessation between e-cigarettes and NRT and obtained a pooled RR of 1.42 [0.97, 2.09] (Figure 3).

Smoking reduction

All six studies^{4,18-22} assessed smoking reduction. Bullen 2013,¹⁸, Eisenhofer,¹⁹ Hajek 2019,⁴ and Lee SM 2018²² reported the proportion of participants reducing smoking by at least 50%. While Lee SH 2019²¹ also reported on this outcome, the size of the reduction was not specified. Bullen 2013¹⁸ and Lee SH 2019²¹ reported an absolute reduction, and Hatsukami 2019²⁰ reported a relative reduction in cigarettes per day from baseline.

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In the Bullen 2013 study,¹⁸ mean cigarette consumption at 6 months decreased by 9.7 (SE 0.4) in the e-cigarette group, and by 7.7 (SE 0.4) in the NRT group. Mean difference between groups was 1.9 (SE 0.6) (p = 0.002). After excluding people who successfully quit smoking, the RR of decreasing cigarette smoking by at least 50% when comparing the e-cigarette to the NRT groups was 1.61 [1.31, 1.99].

Eisenhofer 2015¹⁹ compared week 3 to week 1, and showed that both e-cigarettes (t = 5.3, p = 0.013) and NRT (t = 3.4, p = 0.015) significantly reduced (\sim 50%) self-reports of cigarettes smoked in the previous 24 hours. This was confirmed by significant reductions of breath CO levels in both groups No additional information could be obtained from the abstract and none of the authors could be reached.

In the Hajek 2019⁴ study, 44 of 345 participants in the e-cigarette group, and 29 of 393 participants in the NRT group experienced a carbon monoxide-validated reduction in smoking of \geq 50% in participants without abstinence between weeks 26 and 52, yielding a relative risk of smoking reduction of 1.73 (1.11-2.70).

Hatsukami 2019²⁰ defined smoking reduction by the estimated ratio of cigarettes smoked at 8 weeks as compared to baseline, with a result of 0.25 (0.17, 0.37) in the e-cigarette group, and 0.29 (0.21, 0.39) in the NRT group (p = 0.185). Additional data obtained from the author showed that 19 participants in the e-cigarette group and 22 participants in the NRT group reduced smoking consumption by 50% (RR 0.86 [0.51, 1.46]) at 8 weeks, and that mean cigarette consumption decreased by 9.22 (SD 7.95) in the e-cigarette group, and by 7.61 (SD 8.27) in the NRT group. The mean difference between groups was 1.61 [-0.97, 4.19].

In the Lee SH 2019²¹ study, mean cigarette consumption decreased at 24 weeks by 6.5 +/- 2.87 (SD) in the e-cigarette group, and by 6.60 +/- 3.75 (SD) in the NRT group (p = 0.974). In addition, 31 out of 75 participants (41.3%) in the e-cigarette group and 19 out of 75 participants (25.3%) in the NRT group reduced their daily cigarette consumption (p = 0.038), but no information on size of smoking reduction is provided. After excluding abstainers, a RR of 1.49 [0.97, 2.31] was obtained for decrease in daily cigarette consumption.

Lastly, in the Lee SM 2018,²² 1 participant in the END group and 4 participants in the NRT group reduced their cigarette consumption by at least half, resulting in a RR 0.15 [0.02, 1.14].

We combined data from the Bullen 2013,¹⁸ Hajek 2019,⁴ Hatsukami 2019²⁰ and Lee SM 2018²² studies comparing smoking reduction of at least 50% between e-cigarettes and NRT, as they used similar measures. Pooled results comparing the difference in smoking reduction between the e-cigarette and the NRT groups produced a RR of 1.25, with the line of equivalence falling within the confidence interval [0.79, 1.98] (Figure 3).

We also combined data from the Bullen 2013,¹⁸, Hatsukami 2019,²⁰ and Lee SH 2019²¹ comparing mean reduction of cigarettes per day from baseline for ENDs and NRT. Meta-analysis

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yielded a MD of 1.11, with the line of equivalence falling within the confidence interval [-0.41, 2.63] (Figure 3).

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<u>Harms</u>

Five studies reported on harms (Bullen 2013,¹⁸ Hajek 2019,⁴ Hatsukami 2019,²⁰ Lee SH 2019,²¹ Lee SM 2018²¹). None of the included studies reported serious adverse events (SAEs) related to e-cigarettes or NRT.

In the Bullen 2013¹⁸ study, 107 participants in the e-cigarette group reported 137 adverse events, while 96 participants in the NRT group (patches) reported 119 events, and, using the number of participants available for analysis at 6 months, there was no difference in the incidence of adverse events between groups (RR 0.99, [0.81, 1,22]). No difference between groups was also observed in the Hatsukami 2019²⁰ study, where additional data provided by the author showed that 51 of 69 participants in the e-cigarette group and 53 of 72 participants in the NRT group reported adverse events (1.00 [0.82, 1.22]), and in the Lee SM 2018²² study, where no significant difference in the incidence of adverse events between at 8 weeks (RR 1.24 [0.54, 2.84]).

Hajek 2019⁴ defined adverse events of interest as nausea, sleep disturbances, and throat and mouth irritation. There were 27 SAEs in the e-cigarette group and 22 in the NRT group, none felt to be related to the intervention or control products. Based on the number of participants available at the 12 month follow-up, e-cigarettes were found to be less likely associated with nausea (RR 0.78 [0.66, 0.92]) and sleep disturbances (RR 0.88 [0.83, 0.95]), but more likely associated with throat/mouth irritation (RR 1.24 [1.13, 1.37]). These numbers however should be interpreted with caution as it was not possible to determine with certainty the denominator from the data.

In the Lee SH 2019 study, ²¹ 5 participants in the e-cigarette group and 13 participants in the nicotine gum group reported adverse events. There were no SAEs. Based on the number of participants who completed the study, e-cigarettes were less likely to be associated with adverse events (RR 0.13 [0.12, 0.87]).

We combined data from the Bullen 2013,¹⁸ Hatsukami 2019,²⁰ Lee SH 2019,²¹ Lee SM 2018²² studies comparing harms between e-cigarettes and NRT. Hajek 2019⁴ was excluded as they did not clearly report the number of participants that experienced any adverse events and reported only on specific adverse events. Pooled results comparing ENDS to NRT yielded a RR of 0.96 [0.76, 1.20] (Figure 3).

Withdrawal symptoms

Four studies reported on the results of withdrawal symptoms (Eisenhofer 2015,¹⁹ Hajek 2019,⁴ Hatsukami 2019,²⁰ and Lee SM 2018²²) and all used different scales. Eisenhofer 2015¹⁹ assessed withdrawal with the Questionnaire on Smoking Urges (QSU), Hajek⁴ used a composite urge score (frequency and strength of urge to smoke), Hatsukami 2019²⁰ measured the severity of withdrawal using the Minnesota Nicotine Withdrawal Scale, and Lee SM 2018²¹ assessed withdrawal symptoms as part of their adverse event assessment. In light of the differences in outcome assessment measures, the data were not pooled. In Eisenhofer 2015,¹⁹ urges and cravings to smoke were significantly reduced in the e-cigarette group (t=3.8, p = 0.03), but not in the NRT group (t=2.1, p = 0.08).

In Hajek 2019,⁴ urges for e-cigarette users decreased more than for NRT users at 1 week (MD: -0.4 (-0.6 to -0.2)) and at 4 weeks (MD: -0.3 (-0.5 to -0.1)). E-cigarette users also reported a smaller increase from baseline in irritability, restlessness, inability to concentrate, hunger, and depression. The withdrawal symptoms disappeared mostly for both groups by week 4.

In Hatsukami 2019,²⁰ participants in the e-cigarette group reported lower median [min/max] changes from baseline on the severity scale compared to participants in the NRT group at all measurement points, with week 1 (3.0 [-9.0/25.0] vs 3.5 [-20.0/32.0]), week 2 (1.0 [-13.0/25.0] vs 3.0 [-13.0/39.0]), and week 4 (1.0 [-17.0/30.0] vs 2.5 [-28.0/29.0]). The planned pairwise comparisons were significant with p <0.017. As well, fewer participants (5.3%) withdrew from the complete substitution e-cigarettes group than from the NRT group (15.8%) for product related reasons (disliking product or experiencing withdrawal symptoms; p value not reported).

Lee SM 2018²² only reported on withdrawal symptoms for the NRT group, and did not report on withdrawal symptoms for the e-cigarette group.

Acceptance of therapy

Four studies reported on acceptance of therapy (Bullen 2013,¹⁸ Hajek 2019,⁴ Hatsukami 2019,²⁰ and Lee SM 2018²²), and all used different scales. In light of the difference in outcome assessment measures, the data were not pooled.

In the Bullen 2013¹study,¹⁸ 230 out of 260 participants (88%) in the e-cigarettes group said they would recommend their allocated product to a friend at 1 month, as compared to 130 out of 232 participants (56%) in the NRT group (RR 1.58 [1.40, 1.78]). At 6 months, 205 out of 241 participants (85%) in the e-cigarettes group said they would recommend their allocated product as compared to 107 out of 215 participants (50%) in the NRT group (RR 1.71 [1.48, 1.97]).

In the Hajek 2019 study,⁴ acceptance of therapy was measured with a Likert scale (1 to 5, with a higher score associated with higher acceptance). At 4 weeks post quit date, helpfulness of e-cigarettes was rated 4.3 (SD 0.9) while that of NRT was 3.7 (SD 0.9) (mean difference 0.6 (0.4, 0.7)). Taste was scored at 3.5 (SD 1.3) for the e-cigarette group and 3.1 (SD 1.5) (mean difference 0.4 (0.2,0.6)), and satisfaction was rated at 2.7 (SD 1.1) and 2.3 (SD 1.2), respectively, for the e-cigarette and NRT groups (mean difference 0.5 (0.3, 0.6)).

In the Hatsukami 2019 study²⁰, acceptance of therapy was defined as satisfaction with the product, psychological reward, enjoyment of sensation, aversion, and ability to reduce craving. Results are reported for the NRT group as an estimated mean difference and 95% CI in product evaluation sub-scales using the e-cigarette group as a reference. The following results are reported; satisfaction: -0.6 (-1.0, -0.1), psychological reward: -0.4 (-0.8, 0.01), enjoyment of sensation: -0.6 (-1.1, -0.1), aversion: 0.1 (-0.2, 0.4), and ability to reduce craving: -0.3 (-0.8, 0.2).

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Lastly, the Lee SM 2018 trial²² defined acceptance of therapy as satisfaction with the assigned product, measured with a Likert scale (1 to 7, with a higher score associated with higher satisfaction). Median scores and IQR are reported. Participants randomized to the e-cigarette group reported scores of 6 [4-7], 5.5 [2.5-7], and 6 [5-7], respectively, while participants randomized to the NRT group reported scores of 5 [3-7], 5 [3-6], and 7 [6-7], respectively for the following questions. "The product is helpful for quitting smoking", "I was satisfied with the product to help with quitting", "I would recommend the product to someone interested in quitting smoking".

Risk of bias across studies

The review process we used was thorough, and we took every precaution to minimize the risk of bias due to publication bias or selective reporting. We reached out to clinical experts to enquire about unpublished reports, examined protocol registries, and contacted the authors of identified protocols to request unpublished results. Given the low number of retained studies, we did not include a funnel plot.

Sensitivity, subgroup and meta-regression analyses

We performed a sensitivity analysis for the smoking cessation outcome by removing the Lee SM 2018 study²². While the other 4 studies aimed to assess smoking cessation in general, Lee et al were targeting a peri-operative population, who may have had different motivations to quit smoking. The pooled data, once Lee SM 2018²² is removed, yield a RR of smoking abstinence of 1.39 [0.92, 2.11] when comparing ENDS to NRT (Figure 4).

We had planned on undertaking multiple subgroup analyses. We were unable to perform the subgroup analyses based on age (all participants were adults), smoking intensity (no study enrolled smokers \geq 25 cigarettes per day), or biochemically validated smoking cessation (all studies used biochemical validation). We also could not perform a subgroup analysis of studies with ties to industry as only Bullen 2013¹⁸ was found to have ties to the vaping industry.

We did, however, perform the following subgroup analyses: limiting comparator to nicotine patches (Bullen 2013¹⁸ and Lee SM 2018²¹), and including only studies assessing continuous/sustained smoking abstinence ≥ 6 months given that smoking cessation is defined as sustained abstinence for at least 6 months;²³ (Bullen 2013,¹⁸ Hajek 2019,⁴ Lee SH 2019²¹) (Figure 4).

Metaregression analyses were not performed as our threshold of 10 eligible studies was not met.

DISCUSSION

In our review, there was no significant difference in smoking cessation, smoking reduction, or harms between e-cigarette and NRT users. However, we report on results from a limited number of RCTs, and the level of evidence is low. Our efficacy results are similar to those described in a 2016 Cochrane review,⁵ which also showed no difference between abstinence rates between the nicotine e-cigarette group and NRT group. Their review only included one Smoking cessation in individuals who use vaping, as compared to traditional nicotine replacement therapies; a 2 systematic review and meta-analysis

study¹⁸, also included in our review for this particular outcome. Similar to the evidence we are presenting, none of the studies examined in the Cochrane review reported serious adverse events considered to be related to e-cigarette use.

Although our meta-analysis of the 5 trials that examined *smoking cessation* showed no significant difference between e-cigarette and nicotine replacement therapy, there was a trend towards favoring e-cigarettes. Interestingly, our sensitivity analysis limiting inclusion to studies reporting smoking cessation of 6 months or greater yielded a smaller point estimate than the one obtained from the main analysis, although still with no difference between groups. It could be hypothesized that additional benefits that may be attributed to e-cigarette early on in smoking cessation may be attenuated as time progresses. This again should be interpreted with caution given the small number of studies^{4,18,20} and the very significant heterogeneity.

In all comparisons, our results need to be interpreted carefully. There was significant clinical heterogeneity between studies in terms of the population enrolled, smoking intensity at baseline, type and nicotine concentration of e-cigarettes, type and dose of NRT, as well as methodological heterogeneity in terms of study conduct, and intervention and control protocols. For instance, one of the included studies¹⁸ used first-generation e-cigarettes, with nicotine delivery about 20% of that obtained from cigarette smoking. While e-cigarette users were couriered the supplies needed, NRT users had to redeem vouchers from community pharmacies to obtain their patches. The low nicotine content of the e-cigarettes, the extra step in obtaining NRT supplies, and the low intensity of additional co-interventions likely contributed to the low rate of smoking abstinence at 6 months in both groups, limiting the generalizability of the results. Another included study⁴ allowed for multiple types and concentrations of ENDS, as well as upwards of 10 NRT products and doses, complicating the interpretation of the results. Nicotine concentrations reported in the trials ranged from 0.01 to 48 mg/mL,^{4,18,20-22} making comparisons between studies difficult.

Given that the risk of bias was assessed as high in 5 of 6 included studies^{4,18-21}, our smoking cessation outcome results need to be interpreted with caution. In addition, it is interesting to note that all studies verified self-reported smoking cessation with an exhaled carbon monoxide test, however different cut-off values were used. Additionally, there are limitations to using carbon monoxide (CO) as a way to verify smoking cessation. CO has a relatively short half-life and is eliminated from the body within 24 hours; it can, therefore, lead to false negative results. However, this issue is somewhat mitigated by the fact that smoking cessation study participants tend to be daily smokers.

All studies included in this review examined *smoking reduction*. There was no difference between groups in the mean reduction of cigarettes from baseline in the studies that measured that outcome, or in the proportion of participants successfully reducing their smoking consumption.

None of the included studies reported severe *adverse events* related to ENDS or NRT, and, for the four studies with data that could be pooled, there was no difference between groups in Smoking cessation in individuals who use vaping, as compared to traditional nicotine replacement therapies; a 3 systematic review and meta-analysis

terms of harms related to either therapy. However, in addition to the clinical heterogeneity mentioned above, there was significant methodological heterogeneity in how adverse events were collected. We evaluated the quality of the evidence as very low, given the high risk of bias of included studies, the significant heterogeneity, and the inability to accurately determine the number of subjects involved in this outcome, thus leading to result imprecision.

Since the included trials were powered to detect a difference in the primary outcome, it is possible that rare or unexpected harms were not detected due to a lack of power for this specific outcome. Also, it is important to acknowledge that these studies are limited by their short time-frame. Data on long-term side effects of ENDS are lacking. The recent e-cigarette, or vaping product use-associated lung injury (EVALI) epidemic, is a reminder that further research is needed before widespread recommendations can be made with regards to the use of ENDS. In addition, there are now emerging concerns that respiratory disease caused by the novel coronavirus SARS-CoV-2, the virus responsible for the COVID-19 pandemic, could be exacerbated by exposure to ENDS.²⁴⁻²⁶

Finally, although there seemed to be *increased acceptance of therapy* towards e-cigarettes in the four studies that considered it,^{4,18,20,22} high risk of bias, significant heterogeneity, and the small number of studies using widely different scales leading to imprecise measures, mean that the results should be interpreted with extreme caution. In addition, given that the trials were unblinded, participants who were disappointed with their treatment allocation may have reported less acceptability than their counterparts.

Limitations at review level

We restricted our search to RCTs to try to minimize the risk of bias, however, this considerably limited the number of available studies for this review. It is surprising that, given the widespread availability of e-cigarettes and how aggressively they have been marketed as smoking cessation agents, there are so few head-to-head trials comparing ENDS and traditional NRT. While there may be some unpublished studies that our review did not capture, our literature search was thorough and included personal communications to multiple experts in the field.

Our review identified 7 ongoing trials²⁷⁻³³ that potentially met our inclusion criteria, totaling over 1500 targeted participants. None of the investigators had any data ready to be shared, however it is hoped that this ongoing research can shed light on the effectiveness of ENDS as smoking cessation tools, as compared to traditional NRTs. Long-term research is also needed to investigate the long-term effects of ENDS, as well as the optimal dosing and method of delivery.

Conclusion

We found no difference in smoking cessation, harms, and smoking reduction between ecigarette and NRT users. However, the quality of the evidence was low. Further research is needed before widespread recommendations can be made with regards to the use of ENDS. Research is also needed to investigate the long-term effects of ENDS, as well as optimal dosing. Smoking cessation in individuals who use vaping, as compared to traditional nicotine replacement therapies; a 4 systematic review and meta-analysis

Table 2- Summary of Findings Table

Nicotine-containing Electronic cigarettes (ENDS) vs Nicotine Replacement Therapies (NRT) for smoking cessation

Population: Current smoke Intervention: Nicotine-con				
Comparison: Nicotine-repl				
Outcomes ENDS as compared to NRT	Relative effect (95% Cl)	Number of participants (studies)	Quality of the evidence (GRADE)	Comments
Cessation	RR 1.42 [0.97, 2.09]	1800 (5 studies)	⊕⊕ 00 ^{1,2} low	
Smoking reduction Proportion of people decreasing cigarette consumption by 50%	RR 1.25 [0.79, 1.98]	1460 (4 studies)	⊕⊕ OO ^{1,2} low	
Mean decrease in cigarettes per day	MD 1.11 [-0.41, 2.63]	633 (3 studies)	⊕⊕ 00 ^{1,2} low	
Adverse events (AEs)	RR 0.96 [0.76, 1.20]	758 (4 studies)	<pre> ① OOO^{1,2,3} Very low </pre>	No severe adverse events related to investigated products were reported
Withdrawal symptoms	Summary data not available	4 studies	+ OOO ^{1,2,3} Very low	Withdrawal measures included Minnesota Nicotin Withdrawal Scale, QSU scores, frequency of urge and strength of urge score, and pre-specified symptoms of depressed mood, irritability, restlessness, and hunger

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Acceptance of therapy	Summary data not available	4 studies	① 000 ^{1,2,3} Very low	Acceptance defin as wanting to recommend		
				product to friends, helpfulness, taste, satisfaction, psychological reward, enjoyment		
				of sensation, aversion, and ability to reduce craving dependin on study		
				on study		
GRADE Working Group grade						
High quality: Further research		-				
Moderate quality: Further re	-	n important impact	on our confidence	in the estimate of		
effect and may change the es Low quality: Further research		important impact o	n our confidence i	n the estimate of eff		
and is likely to change the es						
Very low quality: We are very		imate.				
¹ Downgraded one level beca						
² Downgraded one level beca						
³ Downgraded one level beca	use of imprecision of res	ults				
Data availability						
Data collection forms an	d all raw data can be	e requested thro	ugh the corresp	onding author		
Acknowledgements		-				
We thank Katie O'Hearn	, MSc, (Children's Ho	spital of Eastern	i Ontario Resear	rch Institute) Dr		
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assistance. Figure captions Figure 1: Study Flow Dia Figure 2: Risk of bias for Figure 3: Pooled results	gram each outcome per outcome Subgroup Analyses als who use vaping, as c		awa), for metho	odological		
assistance. Figure captions Figure 1: Study Flow Dia Figure 2: Risk of bias for Figure 3: Pooled results Figure 4: Sensitivity and Smoking cessation in individu systematic review and meta-	gram each outcome per outcome Subgroup Analyses als who use vaping, as c	ompared to traditio	awa), for metho	ement therapies; a		

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Smoking cessation in individuals who use vaping, as compared to traditional nicotine replacement therapies; a 8 systematic review and meta-analysis

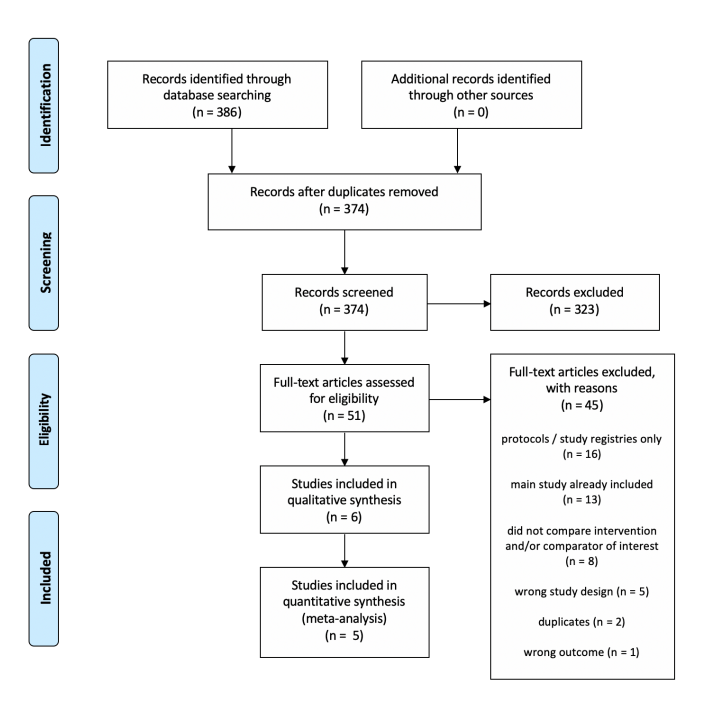
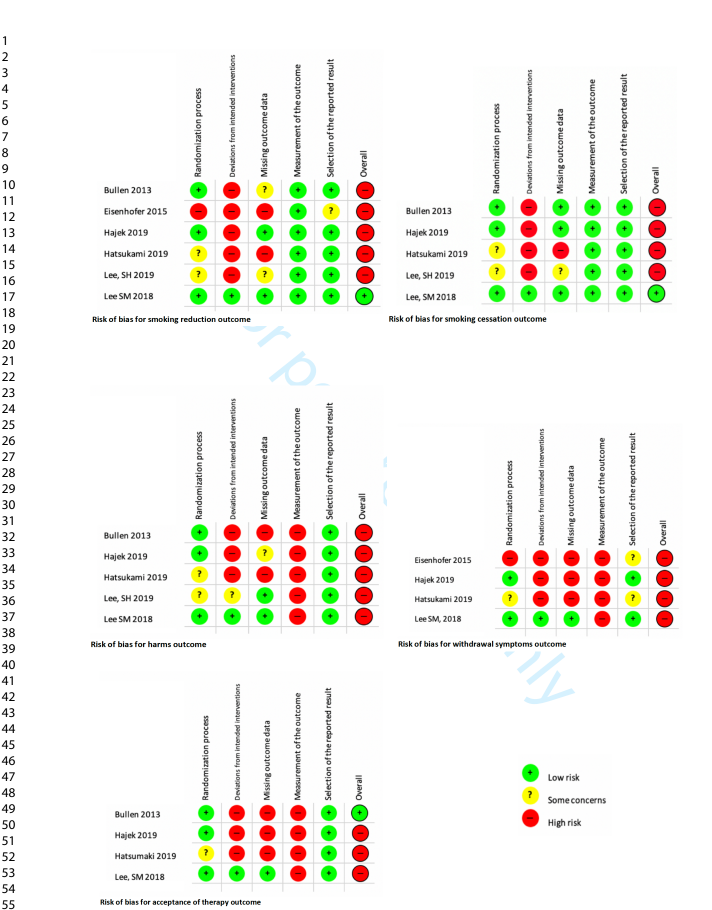


Figure 1. Study flow diagram



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Figure 2. Risk of bias for each outcome

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	END	S	NR	Г		Risk Ratio		Risk Ratio
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI		IV, Random, 95% CI
Bullen	21	289	17	295	20.5%	1.26 [0.68, 2.34]		
Hajek	79	438	44	446	32.2%	1.83 [1.30, 2.58]		
Hatsukami	25	76	13	76	21.5%	1.92 [1.07, 3.47]		
Lee SH	16	75	21	75	22.4%	0.76 [0.43, 1.34]		
Lee SM	5	20	1	10	3.4%	2.50 [0.34, 18.63]		
Total (95% CI)		898		902	100.0%	1.42 [0.97, 2.09]		◆
Total events	146		96					
Heterogeneity: Tau ² =	= 0.09; Cł	$ni^2 = 8.$	00, df =	4 (P =	0.09); I ²	= 50%		
Test for overall effect							0.01	0.1 1 10 10 Favours [NRT] Favours [E-cigarette]

Smoking cessation outcome

	END	s	NR	г		Risk Ratio	Risk Ratio
Study or Subgroup	Events Total		Events	Total	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Bullen	137	268	88	278	38.5%	1.61 [1.31, 1.99]	•
Hajek	44	345	29	393	30.0%	1.73 [1.11, 2.70]	
Hatsukami	19	76	22	76	27.0%	0.86 [0.51, 1.46]	_
Lee SM	1	15	4	9	4.5%	0.15 [0.02, 1.14]	
Total (95% CI)		704		756	100.0%	1.25 [0.79, 1.98]	•
Total events	201		143				
Heterogeneity: Tau ² =	= 0.13; Cł	$ni^2 = 10$).06, df =	= 3 (P =	= 0.02); I	$^{2} = 70\%$	
Test for overall effect	Z = 0.95	5 (P = C)	.34)				0.01 0.1 1 10 100 Favours [NRT] Favours [ENDS]

Proportion of participants successfully reducing smoking consumption by 50%

		ENDS			NRT			Mean Difference		Mea	າ Differen	ce	
Study or Subgroup	Mean	SD	Total	Mean	SD	Total	Weight	IV, Random, 95% CI		IV, Ra	ndom, 95%	6 CI	
Bullen	9.7	5.37	180	7.7	5.2	169	40.0%	2.00 [0.89, 3.11]					
Hatsukami	9.22	7.95	76	7.61	8.27	76	20.6%	1.61 [-0.97, 4.19]			+		
Lee SH	6.55	2.87	71	6.6	3.75	61	39.3%	-0.05 [-1.20, 1.10]			•		
Total (95% CI)			327			306	100.0%	1.11 [-0.41, 2.63]			•		
Heterogeneity: Tau ² =	= 1.18; 0	Chi² =	6.49, d	f = 2 (F	P = 0.0)4); I ² =	= 69%		100	- Ło			10
Test for overall effect	: Z = 1.4	43 (P =	0.15)						-100	-50 Favours [N	TI Favou	50 Irs [ENDS]	10



Mean reduction of cigarettes from baseline

	END	s	NR	г		Risk Ratio	Risk Ratio
Study or Subgroup		-		-	Weight	IV, Random, 95% CI	IV, Random, 95% CI
Bullen	107	241	96	215	43.5%	0.99 [0.81, 1.22]	+
Hatsukami	51	69	53	72	44.7%	1.00 [0.82, 1.22]	+
Lee SH	5	71	13	61	5.0%	0.33 [0.12, 0.87]	
Lee SM	11	20	4	9	6.7%	1.24 [0.54, 2.84]	_
Total (95% CI)		401		357	100.0%	0.96 [0.76, 1.20]	•
Total events	174		166				
Heterogeneity: Tau ² = Test for overall effect:				3 (P =	0.16); I ²	= 42%	0.01 0.1 1 10 100 Favours [NRT] Favours [ENDS]

Proportion of participants experiencing adverse events

Figure 3. Pooled results per outcome

Study or Subgroup Bullen Hajek Hatsukami Lee SH Total (95% CI) Total events	21	Total 289 438 76 75	17 44 13	295 446	21.6% 32.3%	1.83 [1.30, 2.58]	
Hajek Hatsukami Lee SH Total (95% CI) Total events	79 25	438 76	44 13	446	32.3%	1.83 [1.30, 2.58]	
Hatsukami Lee SH Total (95% CI) Total events	25	76	13				
Lee SH Total (95% CI) Total events				76	22 60/		
Total (95% CI) Total events	16	75			22.0%	1.92 [1.07, 3.47]	
Total events			21	75	23.4%	0.76 [0.43, 1.34]	
		878		892	100.0%	1.39 [0.92, 2.11]	•
	141		95				
Heterogeneity: Tau ² =	0.11; Chi ²	$^{2} = 7.$	74, df =	3 (P =	0.05); I ²	= 61%	0.01 0.1 1 10
Test for overall effect:	Z = 1.55 ((P = 0)).12)				Favours [NRT] Favours [ENDS]
e. 1. e. 1	ENDS		NRT			Risk Ratio	Risk Ratio
Study or Subgroup						IV, Random, 95% CI	IV, Random, 95% Cl
Bullen		289	17	295	91.3%	1.26 [0.68, 2.34]	-
Lee SM	5	20	1	10	8.7%	2.50 [0.34, 18.63]	
Total (95% CI)		309		305	100.0%	1.34 [0.74, 2.42]	•
Total events	26		18				
Heterogeneity: Tau ² =	0.00; Chi ²	= 0.4	41, df =	1 (P =	0.52); I ² =	= 0%	0.01 0.1 1 10
Helefogeneity. rau =	7 = 0.97 (1)	(P = 0)	.33)				Favours [NRT] Favours [ENDS]
Test for overall effect:	0.57 (
Test for overall effect:					-		
Test for overall effect:		g ces	sation,	comp	aring e-	cigarettes to nicot	ine patches only
		g ces	sation,	comp	aring e-	cigarettes to nicot	ine patches only

	END	S	NRT	Г		Risk Ratio		Risk	Ratio	
Study or Subgroup	Events	Total	Events	Total	Weight	IV, Random, 95% CI		IV, Rando	m, 95% CI	
Bullen	21	289	17	295	29.2%	1.26 [0.68, 2.34]		_	├ ■──	
Hajek	79	438	44	446	39.7%	1.83 [1.30, 2.58]				
Lee SH	16	75	21	75	31.1%	0.76 [0.43, 1.34]			+	
Total (95% CI)		802		816	100.0%	1.25 [0.73, 2.14]		•		
Total events	116		82							
Heterogeneity: Tau ² = Test for overall effect				2 (P =	0.03); I ²	= 71%	0.01	0.1 Favours [NRT]	1 10 Favours [ENI	100 DS]

Subgroup Analysis— Continuous/sustained abstinence, 6 months and greater only

Figure 4. Subgroup and Sensitivity Analyses

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Supplementary Material 1 Search strategies

MEDLINE, Embase, CENTRAL

 Note: Searches were conducted using an Ovid multi-database search and duplicate records were removed online giving preference to MEDLINE, then Embase, with no field preference. Lines 1-3 are optimized for MEDLINE and the main question constructs are broken out in separate lines for clarity. Lines 4-7 are optimized for Embase and lines 8-10 are optimized for CENTRAL. The next lines isolate the records to the database the search was designed for, combine those sets and then remove duplicate records and final isolate the records from each database again so each can be downloaded and imported into the citation manager using a database-specific import filter.

 Electronic Nicotine Delivery Systems/ or (e cig* or electr* cigar* or electronic nicotine).mp. or (vape or vaper or vapers or vaping or non-combustible nicotine-containing product).ti,ab,kf.
 exp "Tobacco Use Cessation Devices"/ or NRT.ti,ab,kf. or (nicotine adj2 (patch* or gum or nasal spray or mouth spray or mouth strips or lozenge* or tablet* or microtab* or sublingual or replac*)).mp. or (nicotine adj3 therapy).mp.

3. (1 and 2 and ((randomized controlled trial or controlled clinical trial).pt. or randomized.ab. or placebo.ab. or drug therapy.fs. or randomly.ab. or trial.ab. or groups.ab.)) not exp animals/ not humans.sh.

4. Electronic Cigarette/ or (e cig* or electr* cigar* or electronic nicotine).mp. or (vape or vaper or vapers or vaping or non-combustible nicotine-containing product).ti,ab,kw.

5. Nicotine Replacement Therapy/ or NRT.ti,ab,kw. or (nicotine adj2 (patch* or gum or nasal spray or mouth spray or mouth strips or lozenge* or tablet* or microtab* or sublingual or replac*)).mp. or (nicotine adj3 therapy).mp.

6. 4 and 5 and (Crossover-Procedure/ or Double-Blind Procedure/ or Randomized Controlled Trial/ or Single-Blind Procedure/ or (random* or factorial* or crossover* or cross over* or cross-over* or placebo* or (doubl* adj blind*) or (singl* adj blind*) or assign* or allocat* or volunteer*).ti,ab,kw.)

7. limit 6 to embase

8. (e cig* or electr* cigar* or electronic nicotine).mp. or (vape or vaper or vapers or vaping or non-combustible nicotine-containing product).ti,ab,kw.

9. NRT.ti,ab,kw. or (nicotine adj2 (patch* or gum or nasal spray or mouth spray or mouth strips or lozenge* or tablet* or microtab* or sublingual or replac*)).mp. or (nicotine adj3 therapy).mp.

- 10. 8 and 9
- 11. 3 use medall
- 12. 7 use emczd
- 13. 10 use cctr
- 14. 11 or 12 or 13
- 15. remove duplicates from 14
- 16. 15 use medall
- 17. 15 use emczd
- 18. 15 use cctr

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7	ClinicalTrials.gov
8	(electronic cigarette OR vape OR vaping OR electronic nicotine) AND (nicotine replacement OR
9	
10	NRT OR patch OR gum OR nasal spray OR mouth spray OR mouth strips OR lozenge OR tablet
11	OR microtab OR microtablet OR sublingual) Interventional Studies
12	91 records retrieved
13	91 records retrieved
14	
15	WHO ICTRP
16	electronic cigarette OR vape or vaping OR electronic nicotine
17	153 records retrieved with 20 remaining after records with a TrialID starting with NCT were
18	-
19	removed prior to screening
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21	Note: As the ICTED registry has limited search espekilities ³⁵ , only terms related to the
22	Note: As the ICTRP registry has limited search capabilities ³⁵ , only terms related to the
	intervention were used and protocols with a NCT number were removed from the retrieval, as
23	those protocols would also be included in ClinicalTrials.gov.
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Supplementary Material 2 Abstracted data

The abstracted data included the following:

1- study characteristics:

author names, year of publication, ties with tobacco industry, funding of study, country of study, study setting, study design, number of participating sites, recruitment procedures, enrolment dates, length of study period, random sequence generation, allocation sequence concealment, blinding, methods for preventing and controlling confounding, selection bias, information bias and missing bias, unit of analysis, covariates inclusion, funding, financial and conflict of interest disclosure including ties with industry, inclusion and exclusion criteria, sample size, number of participants that were analyzed, number of participants lost to follow up for each outcome and for the whole study, number of participants at study onset and randomized to each group, and type of analysis (intention to treat vs per protocol)

2- participant characteristics:

age, gender, comorbidities, ethnicities, socio-economic status, income, education, cigarettes smoked per day, Fagerström test for cigarette dependence

3- *intervention characteristics:*

type, model, brand and generation of ENDS, type and flavor of e-liquid, nicotine content, intervention protocol, length of time ENDS were provided free of charge, frequency of use, duration of intervention, integrity of intervention, description of co-interventions

4- comparator characteristics:

type of nicotine replacement therapy used, dose, frequency of use, nicotine content, control protocol, frequency of use, length of time supplies were provided free of charge, combination of products, frequency of use duration of control, integrity of control, description of co-interventions

5- outcomes:

smoking cessation, method of assessment for smoking cessation used (self-report vs biochemical), smoking abstinence definition, longest time point of smoking cessation, harms assessment, methods of harms assessment, definition of harms, withdrawal symptoms, method of assessment for withdrawal symptoms, reduction in cigarettes smoked, method of assessment of reduction in cigarettes smoked, number of quit attempts, method of quit attempt measurement, acceptance of ENDS/NRT, method of acceptance assessment, method of aggregation used for each outcome, timing of measurement for each outcome, summary data for each outcome, method of aggregation used for each outcome.

	Detailed description of the included studies
	able 1a. Characteristics of randomized controlled trials measuring smo
	onths or later Characteristics of randomized controlled trials measuring n at 6 months or later
Bullen, 2013	
Methods	Design: 3 parallel groups RCT
	Recruitment: Participants were recruited via community newspapers
	inviting people to call the study centre for eligibility pre-screening
	Setting: one single center in Auckland Australia
	Inclusion criteria: 18 years of age or older, smoked 10 or more cigare
	per day for the past year, and wanted to quit smoking.
	Exclusion criteria: Pregnant or breastfeeding women, people using
	smoking cessation drugs, those reporting heart attack, stroke, severe
	angina in the previous 2 weeks, and people with poorly controlled me
	disorders allergies, or other chemical dependence were excluded
Participants	Total N: 657 smokers were included in this study, but we only extract
	584 participants for our review (2 of the 3 groups) as the e-cigarette
	placebo group did not fit our eligibility criteria.
	Most participants were women (62%), of a mean age > 40. Approximately a state of the second s
	one third were of Maori descent, and a little over half had completed
	grade 12 or above education level. The average daily number of ciga smoked at study onset was around 18, and mean Fagerström test res
	to 10 scale) for cigarette dependence was > 5.
Interventions	Randomization: 4:4:1 ratio to nicotine e-cigarettes, nicotine patches
	placebo e-cigarette group
	Nicotine e-cigarette group
	Participants were couriered a first-generation e-cigarette, spare batte
	and charger, as well as cartridges containing 10 to 16mg of nicotine p
	mL (although labelled to contain 16 mg), plus simple instructions to u
	the e-cigarettes as desired from 1 week before until 12 weeks after the
	chosen quit day. Participants received on average around 20% of the
	nicotine obtained from cigarette smoking.
	Nicotine patch group
	Participants were sent exchange cards in the mail redeemable for nic
	patches 21 mg from community pharmacies, with instructions to use
	patches daily, from 1 week before until 12 weeks after their chosen q
	day. Vouchers were also supplied to participants to cover dispensing
	costs.
	Both groups

	Participants in all groups were also referred to telephone-based
	behavioural support
Outcomes	Continuous abstinence at 6 months after quit day, defined as self-reporte
	abstinence over the whole follow-up period allowing for 5 or less
	cigarettes in total, was self-reported, and verified with exhaled breath
	carbon monoxide of <10 ppm. Harms were both clinically assessed and
	self-reported, throughout the study period. Withdrawal symptoms were
	assessed at 1, 3, and 6 months. Reduction in daily cigarettes smoked was
	measured at 6 months, and acceptance of therapy was measured at 1 and
	6 months.
Notes	Some of this study's authors reported ties to e-cigarette manufacturers,
	and smoking cessation drug companies
Hajek, 2019	
Methods	Design: 2 parallel groups RCT
	Recruitment: Participants were recruited through stop smoking services,
	which included trial information in their advertising. Participants were
	also recruited through social media, and leaflets advertising the trial were
	delivered to local households.
	Setting: 3 sites in the United Kingdom
	Inclusion criteria: Adults, with no strong preference towards e-cigarette
	or NRT, who were not using either type of product at the time of study
	enrolment 💦
	Exclusion criteria: Pregnant women or breastfeeding women
Participants	Total N: 884 participants were included in this study
	Median age for both groups was 41, and women comprised 48% of
	participants. Most participants were White British, and the majority had
	post-secondary education. Median daily number of cigarettes smoked at
	study onset was 15, and mean Fagerström test result for cigarette
	dependence was 4.5 in the e-cigarette group and 4.6 in the NRT group.
Interventions	Randomization: nicotine-containing e-cigarettes of varying doses, and an
	choice of a list of NRT, in a 1:1 ratio
	E-cigarette group
	Participants were provided with a starter pack called One Kit, which
	included an atomizer, a battery, and one 30 mL bottle of Tobacco Royale
	flavor e-liquid. Participants were asked to purchase their future e-liquid
	online or from local vape shops and to buy a different e-cigarette device i
	the one supplied did not meet their needs. They were encouraged to
	experiment with e-liquids of different strengths and flavors. Those who
	were unable to obtain their own supply were provided with one further
	10-ml bottle, but this was not offered proactively. Participants received
	oral and written information on how to operate the e-cigarette.

$\begin{array}{c}1\\2\\3\\4\\5\\6\\7\\8\\9\\10\\11\\2\\31\\4\\15\\16\\17\\18\\19\\20\\21\\22\\32\\4\\25\\26\\7\\28\\9\\30\\31\\23\\34\\5\\36\\37\\38\\9\\0\\41\\42\\43\\44\\5\\6\end{array}$	
41 42 43 44	

NRT group
Participants were informed about the range of nicotine-replacement
products (patch, gum, lozenge, nasal spray, inhalator, mouth spray, mouth
strip, and microtabs) and selected their preferred product. Use of
combinations was encouraged, typically the patch and a faster-acting oral
product. Participants were also free to switch products.ps
Both groups
Participants in both groups were offered multisession behavioral support
as per UK stop smoking service practice, involving weekly one on one
session with local clinicians.
Participants were also asked to sign a commitment to not use the
unassigned treatment for 4 weeks
Continuous abstinence at 52 weeks after quit day, defined as self-reported
abstinence over the whole follow-up period allowing for 5 or less
cigarettes in total, was self-reported, and verified with exhaled breath
carbon monoxide of <8 ppm. Harms were self-reported throughout the
study period. Withdrawal symptoms were assessed at 1 and 4 weeks in
abstainers. Reduction in daily cigarettes smoked was also measured at 52
weeks, as well as acceptance of e-cigarettes and NRT
Some of this study's authors reported ties to smoking cessation drug
companies.
Design: 2 parallel groups PCT
Design: 2 parallel groups RCT
Recruitment: Participants were recruited from a motor company in the
Republic of Korea.
Setting: One site in Cheonan, Republic of Korea
Inclusion criteria: Participants were adults 18 years and above, male, who
smoked at least 10 cigarettes per day in the preceding year, and who were
motivated to stop smoking entirely or to reduce their cigarette
consumption
Exclusion criteria: Participants were excluded if they had a past medical
history of serious clinical diseases or had attempted to stop smoking in the
last 12 months by using other NRT.
Total N: 150 participants were included in the study
Mean age was 42 years and all participants were men. Almost 40% had
post-secondary education. Median daily number of cigarettes smoked at
study onset was 1 pack per day, and mean Fagerström test result for
study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4.
study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4.Randomization: nicotine-containing e-cigarettes, and nicotine gum in a
study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4.
study onset was 1 pack per day, and mean Fagerström test result for cigarette dependence was 4.Randomization: nicotine-containing e-cigarettes, and nicotine gum in a
_

	Participants received a 24-week supply of e-cigarettes eGo-C Ovale, Janty Korea Co., Janty-Asia Co., Seoul, Republic of Korea, nicotine 0.01 mg/mL.
	Nicotine gum group Participants received a 24-week supply of nicotine gum Nicoman, Daewoog Pharmaceutical, Seongnam, Republic of Korea, 2 mg/tablet
	Both groups Participants in both groups were offered 55-minute education sessions or smoking cessation aids
Outcomes	Continuous abstinence was defined as abstinence from smoking from 9 to 24 weeks, validated with end-expiratory carbon monoxide (<10 ppm) and a negative urine cotinine result. Harms were self-reported throughout the study period. Reduction in daily cigarettes smoked was also measured at 24 weeks.
Notes	None of the study authors were found to have ties to industry.
Lee SM, 2018	
Participants	 Recruitment: Participants were recruited from an anesthesia preoperative clinic for elective surgery. Setting: San Francisco Veterans' Affairs Medical Center, affiliated with the University of California in San Francisco United States of America Inclusion criteria: Participants were eligible if they presented to the clinic 3 or more days prior to elective surgery, smoked more than two cigarettee per day, and had smoked at least once in the last 7 days Exclusion criteria: Participants were excluded if they exclusively used other forms of tobacco (e.g. pipe tobacco) or marijuana only, were pregnant or breastfeeding, had an unstable condition, were using smokin cessation therapy at the time of study enrolment or were in another smoking cessation trial, or currently used e-cigarettes daily. Total N: 30 participants were included in this study
	Most participants were men (90%) in their 50's. Some had comorbidities including diabetes, hypertension, heart disease, and chronic obstructive pulmonary disease. Most were Caucasians. The average daily number of cigarettes smoked at study onset was 15.3 in the e-cigarette group, and 10.8 in the NRT group, and the mean Fagerström test result for cigarette dependence was 3.7 in the e-cigarette group and 2.5 in the NRT group.
Interventions	 Randomization: e-cigarettes and nicotine patches in a 2:1 ratio E-cigarette group Participants received a 6-week supply of NJOY e-cigarettes (Scottsdale, AZ USA), a disposable first-generation e-cigarette that is available in shops and online. They were issued a number of e-cigarettes corresponding to

	 the reported baseline cigarettes smoked per day, calculated assuming or NJOY e-cigarette was equivalent to 10 cigarettes. Participants were instructed to smoke bold (4.5%) e-cigarettes ad libitum for 3 weeks, then the Gold (2.4%) e-cigarettes ad libitum for 2 weeks, and then the Study (0%) e-cigarettes ad libitum for the final week. Nicotine patch group Participants randomized to the nicotine patches group were given a 6-week supply of Nicoderm CQ patches (5 weeks) and placebo patches (1 week) appropriate to baseline nicotine consumption. Those smoking an average of ten or more cigarettes per day were given a 21 mg/day patch for 3 weeks, a 14 mg/day patch for 1 week, a 7 mg/day patch for 1 week, and a 0 mg/day patch for 1 week. Participants who reported smoking an average of fewer than 10 cigarettes per day at baseline were given a 14
	 mg/day patch for 3 weeks, a 7 mg/day patch for 2 weeks, and a 0 mg/day patch for 1 week. Both groups Participants in both groups were given referral California Smokers' Helpline and were asked to refrain from the use of cigarettes during the study period.
Outcomes	Smoking cessation at 6 months was self-reported through 7-day point- prevalence abstinence and verified with exhaled breath carbon monoxide of <10 ppm. Harms and withdrawal symptoms were systematically collected at 8 weeks. Reduction in daily cigarettes smoked was also measured at 6 months, as well as acceptance of e-cigarettes and NRT.
Notes	None of the study authors were found to have ties to industry.

Supplementary Table 1b. Characteristics of randomized controlled trial measuring smoking cessation earlier than 6 months

Hatsukami, 2019	
Methods	Design: 4 parallel groups RCT Recruitment: Participants were culled from two sets of studies, one of which also included two groups randomized to snus (spitless smokeless tobacco); one was complete substitution with snus, and the other was ad libitum use. Due to recruitment challenges, the two snus groups were dropped midway through the study, resulting in four experimental groups: ad libitum use of e-cigarettes (participants may smoke as many cigarettes as they like), complete substitution with e-cigarettes (aiming for smoking

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Interventions
Participants

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12 13 14 15 16	
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22 23 24 25 26 27 28	
29 30 31	
32 33 34 35 36 37 38	
39 40 41 42	
43 44 45 46 47	
48 49 50 51 52	
53 54 55 56 57	
58 59 60	

	 Participants could choose between mint, cinnamon or fruit-flavored nicotine gum or nicotine lozenge, at a dose of 4 mg. If adverse effects were recorded, the dose was decreased to 2 mg. Both groups After randomization, participants were asked to complete daily diaries via interactive voice recording to chart the number of cigarettes smoked daily, as well as document assigned product use for the duration of the trial. Participants received a monetary bonus if they complied with the protocol; this included keeping an accurate record of product use, completing the daily diaries, and returning unused products. They also got a bonus payment if they had a carbon monoxide level ≤ 4 ppm at each visit. Participants also received a brief counseling session on how to avoid smoking.
Outcomes	Smoking cessation was determined by 7-day point prevalence at 8 weeks, mainly through biochemical verification but also by self-report Reduction in daily cigarettes smoked was also measured at 8 weeks, as well as acceptance of e-cigarettes and NRT. Harms were assessed systematically at 20 weeks, 12 weeks after the end of the study period. Withdrawal symptoms were assessed at weeks 1, 2, 4, 6, and 8.
Notes	One of the study authors is a member of the FDA Tobacco Products Scientific Advisory Committee and another one has served as an expert witness in tobacco company litigation.

Supplementary Table 1c. Characteristics of randomized controlled trial measuring other outcomes

outcomes	
Eisenhofer,	
2015	
Methods	Design: 2 parallel groups RCT
	Recruitment: Not specified
	Setting: Not specified
	Inclusion criteria: Veterans who met criteria for tobacco disorder as per
	the DSM
	Exclusion criteria: Not specified
Participants	Total N: 11 participants were included
	Mean age was 52, and 82% were males. The vast majority of participants
	were African American. The average daily number of cigarettes smoked at
	study onset was 26.5, and the mean Fagerström test result for cigarette
	dependence was 7.5.
Intervention	Randomization: e-cigarettes and nicotine patches
	E-cigarette group

	Participants received nicotine-containing e-cigarettes with 16 mg of nicotine per cartridge
	NRT group Participants received nicotine patch 16 mg daily
	Both groups All participants were instructed to smoke ad libitum during week 1, and to
Outcomes	smoke as little as possible during week 3.Reduction in cigarettes smoked per day was self-reported at 3 weeks and compared to week 1. Withdrawal symptoms were compared between week 1 and week 3.
Notes	This study was available as an abstract only therefore limited details are available.

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Supplementary Material 4 Details on Risk of Bias Assessment for each outcome of interest

Supplementary Table 2. Detailed description of concerns for each domain marked identified as "some concerns" or "high risk" on Risk of Bias Assessment

	Randomization	Deviations from intended	Missing of outcome	Measurement of the	Selection of the
	Process	intervention	data	outcome	reported result
Bullen 2013	Low risk	Adherence higher in the ENDS group compared to NRT group at all timepoints. At 6 months, 29% of ENDS group vs 8% of NRT group still using assigned treatment.	Low risk	Low risk	Low risk
Hajek 2019	Low risk	At 52 weeks among participants with 1-year abstinence, 80% were using e- cigarettes in the ENDS group vs 9% in the NRT group. Also, 6% of participants in the ENDS group reported using non-allocated NRT for at least five consecutive days in the past six months compared to 22% in the NRT group that reported using non-allocated product	Low risk	Low risk	Low risk
Hatsukami 2019	No information provided with regards to randomization process and allocation concealment. However, there were no	The NRT group had the highest dropout rates compared to the other groups in the study. At 8 weeks, 24% dropped out in the ENDS group compared to 30% in the NRT group.	Large number of dropouts; participants who did not stop smoking could be less motivated to continue with study follow up	Low risk	Low risk

	significant baseline differences between groups				
Lee, SH 2019	The use of constant block sizes of 2 makes it easy to determine order of randomization.	No participants discontinued the intervention. However, 4 and 14 participants in the ENDS and NRT group dropped out before treatment, respectively.	Although data was missing for 12% of randomized individuals, all dropouts occurred prior to the start of treatment. Missingness in this case less likely to be due to the value of the outcome as it happened prior to onset of therapy	Low risk	Low risk
Lee, SM 2018	Low risk	Low risk	Low risk	Low risk	Low risk
,					
	luction outcome		·		
Smoking red	Low risk	Refer to smoking cessation outcome	Sensitivity analyses conducted for the smoking cessation outcome were not performed for the smoking reduction outcome	Low risk	Low risk
Smoking red Bullen 2013 Eisenhofer		Refer to smoking cessation outcome Not enough information available in abstract	conducted for the smoking cessation outcome were not performed for the smoking reduction	Low risk	Low risk Not enough informatio available in abstract
	Low risk Not enough information available in	Not enough information available in	conducted for the smoking cessation outcome were not performed for the smoking reduction outcome Not enough information		Not enough informatic
Smoking red Bullen 2013 Eisenhofer 2015 Hajek 2019 Hatsukami	Low risk Not enough information available in abstract	Not enough information available in abstract	conducted for the smoking cessation outcome were not performed for the smoking reduction outcome Not enough information available in abstract	Low risk	Not enough informatic available in abstract
Smoking red Bullen 2013 Eisenhofer 2015	Low risk Not enough information available in abstract Low risk Refer to smoking	Not enough information available in abstract Refer to smoking cessation outcome	conducted for the smoking cessation outcome were not performed for the smoking reduction outcome Not enough information available in abstract	Low risk	Not enough informatic available in abstract

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Bullen 2013	Low risk	Differences in treatment adherence	No information on the	high likelihood that	Low risk
		could potentially lead to	proportion of	participants who were	
		discrepancies in harm reporting	participants on whom	unhappy with their	
			adverse events were	treatment allocation	
			collected; it is likely that	would report side	
			people who experienced	effects more often	
			more severe side effects	than their	
			did not continue with	counterparts.	
			study follow-up activities		
Hajek 2019	Low risk	Differences in treatment adherence	The authors reported	High likelihood that	Low risk
		could potentially lead to	harm data based on	participants who were	
		discrepancies in harm reporting	number of participants	unhappy with their	
			at randomization,	treatment allocation	
			however significant	would report side	
			dropout seen at 4-week	effects more often	
			follow up, raising	than their counterparts	
			concerns that adverse		
			event data not collected		
			on all participants		
Hatsukami	Refer to smoking	Differences in treatment adherence	No information on the	High likelihood that	Low risk
2019	cessation outcome	could potentially lead to	proportion of	participants who were	
		discrepancies in harm reporting	participants on whom	unhappy with their	
			adverse events were	treatment allocation	
			collected; it is likely that	would report side	
			people who experienced	effects more often	
			more severe side effects	than their counterparts	
			did not continue with		
			study follow-up activities		
Lee, SH 2019	Refer to smoking	Differences in treatment adherence	Low risk	High likelihood that	Low risk
	cessation outcome	could potentially lead to		participants who were	
		discrepancies in harm reporting		unhappy with their	
		however non-adherence happened		treatment allocation	
		prior to onset of treatment,		would report side	
		therefore less likely to have an		effects more often	
		impact		than their counterparts	
Lee, SM 2018	Low risk	Low risk	Low risk	High likelihood that	Low risk
				participants who were	
	1		1	unhappy with their	

				treatment allocation would report side effects more often than their counterparts	
Withdrawal	symptoms outco	me			
Eisenhofer 2015	Not enough information available in abstract	Not enough information available in abstract	Not enough information available in abstract	Not enough information available in abstract	Not enough informatio available in abstract
Hajek 2019	Low risk	Differences in treatment adherence could potentially lead to discrepancies in withdrawal symptoms reporting	Outcome not available for all randomized participants; likely that people who experienced more nicotine withdrawal symptoms did not continue with study follow-up activities	Given that the withdrawal measurements were self-reported, there is a high likelihood that participants who were unhappy with treatment allocation reported more withdrawal symptoms than their counterparts	Low risk
Hatsukami 2019	Refer to smoking cessation outcome	Differences in treatment adherence could potentially lead to discrepancies in withdrawal symptoms reporting	Outcome not available for all randomized participants; likely that people who experienced more nicotine withdrawal symptoms did not continue with study follow-up activities	Given that the withdrawal measurements were self-reported, there is a high likelihood that participants who were unhappy with treatment allocation reported more withdrawal symptoms than their counterparts	No information on how withdrawal symptom assessment was performed
Lee, SM 2018	Low risk	Low risk	Low risk	Given that the withdrawal measurements were self-reported, there is a high likelihood that participants who were	Low risk

				unhappy with treatment allocation reported more withdrawal symptoms than their counterparts	
Acceptance of	of therapy out	come			
Bullen 2013	Low risk	Differences in treatment adherence could potentially lead to discrepancies in acceptance of therapy outcome	Participants unhappy with their assigned therapy likely did not continue with study follow-up activities	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk
Hajek 2019	Low risk	Differences in treatment adherence could potentially lead to discrepancies in acceptance of therapy outcome	Participants unhappy with their assigned therapy likely did not continue with study follow-up activities	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk
Hatsukami 2019	Not enough information available in abstract	Differences in treatment adherence could potentially lead to discrepancies in acceptance of therapy outcome	Participants unhappy with their assigned therapy likely did not continue with study follow-up activities	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk
Lee, SM 2018	Low risk	Low risk	Low risk	Highly subjective outcome, inability to blind participants to assigned therapy	Low risk





PRISMA 2009 Checklist

Section/topic	#	Checklist item	Reported on page #			
TITLE		·				
Title	1	Identify the report as a systematic review, meta-analysis, or both.	0			
ABSTRACT						
Structured summary	2	Provide a structured summary including, as applicable: background; objectives; data sources; study eligibility criteria, participants, and interventions; study appraisal and synthesis methods; results; limitations; conclusions and implications of key findings; systematic review registration number.	1			
Rationale	3	Describe the rationale for the review in the context of what is already known.	4			
Objectives	4	Provide an explicit statement of questions being addressed with reference to participants, interventions, comparisons, outcomes, and study design (PICOS).	4			
20 21 METHODS						
Protocol and registration	5	Indicate if a review protocol exists, if and where it can be accessed (e.g., Web address), and, if available, provide registration information including registration number.	5			
Eligibility criteria	6	Specify study characteristics (e.g., PICOS, length of follow-up) and report characteristics (e.g., years considered, language, publication status) used as criteria for eligibility, giving rationale.	5			
Information sources	7	Describe all information sources (e.g., databases with dates of coverage, contact with study authors to identify additional studies) in the search and date last searched.	5,6			
Search	8	Present full electronic search strategy for at least one database, including any limits used, such that it could be repeated.	5,6, Supp material 1			
Study selection	9	State the process for selecting studies (i.e., screening, eligibility, included in systematic review, and, if applicable, included in the meta-analysis).	6			
Data collection process	10	Describe method of data extraction from reports (e.g., piloted forms, independently, in duplicate) and any processes for obtaining and confirming data from investigators.	6			
Data items	11	List and define all variables for which data were sought (e.g., PICOS, funding sources) and any assumptions and simplifications made.	6, Supp material 2			
Risk of bias in individual studies	12	Describe methods used for assessing risk of bias of individual studies (including specification of whether this was done at the study or outcome level), and how this information is to be used in any data synthesis.	6			
Summary measures	13	State the principal summary measures (ajgperistration difference in gmeans) es. xhtml	6			

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PRISMA 2009 Checklist

Synthesis of results	14	Describe the methods of handling data and combining results of studies, if done, including measures of consistency (e.g., I ²) for each meta-analysis.	7	
Page 1 of 2				
Section/topic	#	Checklist item	Reported on page #	
Risk of bias across studies	15	Specify any assessment of risk of bias that may affect the cumulative evidence (e.g., publication bias, selective reporting within studies).	6	
Additional analyses	16	Describe methods of additional analyses (e.g., sensitivity or subgroup analyses, meta-regression), if done, indicating which were pre-specified.	7	
RESULTS				
Study selection	17	Give numbers of studies screened, assessed for eligibility, and included in the review, with reasons for exclusions at each stage, ideally with a flow diagram.	7, Figure	
Study characteristics	18	For each study, present characteristics for which data were extracted (e.g., study size, PICOS, follow-up period) and provide the citations.	7,8, Supp material 3	
Risk of bias within studies	19	Present data on risk of bias of each study and, if available, any outcome level assessment (see item 12).	9, Figures 2a,b,c,d,e	
Results of individual studies	20	For all outcomes considered (benefits or harms), present, for each study: (a) simple summary data for each intervention group (b) effect estimates and confidence intervals, ideally with a forest plot.	9-13, Figures 3a,b,c,d,	
Synthesis of results	21	Present results of each meta-analysis done, including confidence intervals and measures of consistency.	9-12, Figures 3a,b,c,d	
Risk of bias across studies	22	Present results of any assessment of risk of bias across studies (see Item 15).	13	
Additional analysis	23	Give results of additional analyses, if done (e.g., sensitivity or subgroup analyses, meta-regression [see Item 16]).	13, Figures 4a,b,c	
DISCUSSION				
Summary of evidence	24	Summarize the main findings including the strength of evidence for each main outcome; consider their relevance to key groups (e.g., healthcare providers, users, and policy makers).	13-15	
Limitations	25	Discuss limitations at study and outcome level (e.g., risk of bias), and at review-level (e.g., incomplete retrieval of identified research, reporting bias), identified research, reporting bias), For peer review only - http://bmjopen.bmj.com/site/about/guidelines.xhtml	13-15	

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4	Conclusions	26	Provide a general interpretation of the results in the context of other evidence, and implications for future research.	13-15	
6	UNDING				
7 8 9	Funding	27	Describe sources of funding for the systematic review and other support (e.g., supply of data); role of funders for the systematic review.	0	
9↓ 10	From: Moher D, Liberati A, Tetzlaff doi:10.1371/journal.pmed1000097	J, Altm	an DG, The PRISMA Group (2009). Preferred Reporting Items for Systematic Reviews and Meta-Analyses: The PRISMA Statement. PLoS Med For more information, visit: www.prisma-statement.org. Page 2 of 2	6(7): e1000097.	
43 44 45 46 47			For peer review only - http://bmjopen.bmj.com/site/about/guidelines.xhtml		