Critical appraisal of interventional clinical trials assessing heated tobacco products: a systematic review and meta-regression.

Supplementary Materials

Supplementary Appendix 1. Coding of trial affiliation

Trials were coded as 'Industry-affiliated' if:

- the study sponsor named on the trial registration was a tobacco company or other organisation directly funded by a tobacco company; or
- funding statements in any of the trial literature indicated the trial was funded in part or in
 whole by a tobacco company or other organisation directly funded by a tobacco company; or
- author affiliations or conflict of interest statements indicated any author was an employee or funded by a tobacco company or other organisation directly funded by a tobacco company at the time of the trial.

Trials were coded as 'Independent' if:

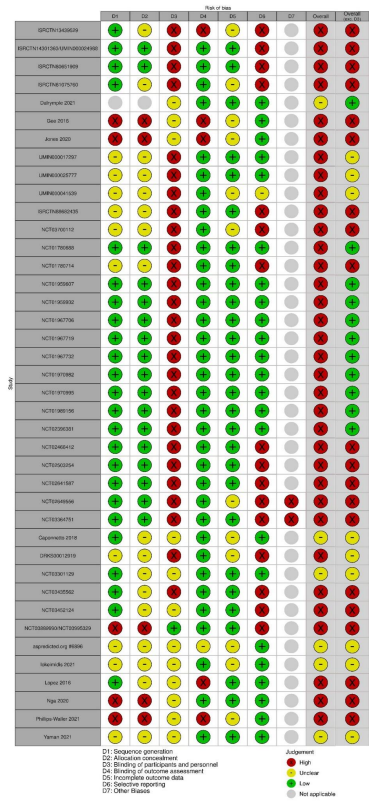
- the sponsor named on the trial registration had no known ties to the tobacco industry; and
- funding statements in any of the trial literature indicated the trial was not funded by a tobacco company or other organisation funded by a tobacco company; and
- author affiliations and conflict of interest statements indicated authors had no contemporary
 (i.e., while the study was being conducted, up to and including publication) ties to the tobacco industry.

Trials were coded as 'Unclear' if:

- There was insufficient information to determine affiliation; or
- Reviewers could not reach consensus.

In addition to conflict of interest and funding statements provided in the trial literature, we further investigated known ties and funding using the Tobacco Tactics website (www.tobaccotactics.org), relevant literature published by the Tobacco Control Research Group (University of Bath), and conflict of interest and funding statements in other contemporary work of the authors of included studies.

Supplementary Figure 1. Risk of bias summary: Review authors' judgments about risk of bias items for each included study



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Supplementary Table 1. Characteristics of included studies.

UMIN00001729	UMIN000017297			
Methods	Date o	f registration	: 27/04/2015	
	Submi	itted to peer-r	reviewed journal within 12 months: No	
	Publis	hed key outco	omes on trial registration within 12 months: No results posted	
	Design	: Crossover R	CT	
	Setting	g (Country): (Confinement (Japan)	
	Study	start date; st	udy end date: 11/05/2015; 27/05/2015	
	Interv	ention durati	on: 2 sessions of 10 puffs for 3 mins at approx 20 sec intervals	
Participants	Numb	er of particip	ants: 24 randomised, 0 withdrawn, 24 completed	
			s reported: N/A	
	male; I	Ethnicity/Natio	stics: N=24; Mean Age (SD): 39 years (SD not reported); Sex: 100% onality: 100% Japanese.	
	Key in	clusion criter	ia: Health status: "good health"; ≥11 CPD; smoked for ≥1 year	
Interventions	Interv	entions: HTP	(Prototype novel tobacco vapor product), CC (unknown brand)	
	Co-int	erventions: n	one	
	Mode	of exposure:	direct restricted	
Outcomes		ary: Time to reach nicotine Cmax, Maximal nicotine concentration, Area under the entration curve from start of product use to time of last quantifiable concentration		
	chemis	stry, haematolo	Events/Serious Adverse Events, Physical examination, Clinical ogy and urine analysis safety panel, Vital signs, Terminal half-life of l exposure to nicotine.	
Analyses	Sampl	e size calcula	tion reported: Yes	
·	Primary analysis population: Per-protocol population defined as "completed subjects who completed the study and who did not deviate from the protocol were included in the statistical analysis"			
		f analysis: Inc	lividuals	
Study funding		-	national (Industry-affiliated)	
Notes	Not inc	cluded in meta	i-regression analysis	
Risk of bias			8	
Bias		Authors'	Support for judgement	
Dias		judgement	Support for judgement	
Random sequence generation	e	Unclear	Beyond stating the study was 'randomised', no further information provided.	
Allocation conce		Unclear	No information provided.	
Blinding of participants and personnel		High	"Blinding: Open-no one is blinded". Included non-active comparator (cigarettes).	
Blinding of outcome assessment		Low	"Blinding: Open-no one is blinded". All primary outcomes were objectively measured.	
Incomplete outco	ome	Low	All subjects randomised completed the study and were included in the analyses.	
Selective reporting	ng	Low	3 safety profile parameters were not reported, but adverse events data were reported. All other outcomes listed in the methods and on the trial registration are reported on in at least one literature source.	
UMIN00002577	7			

Methods Date of registration: 20/01/2017

Submitted to peer-reviewed journal within 12 months: Yes

Published key outcomes on trial registration within 12 months: No results posted

Design: Parallel RCT

Setting (Country): Confinement (Japan)

Study start date; study end date: 21/01/2017; 22/02/2017

Intervention duration: 5 days

Participants		Number of participants: 60 randomised (HTP 20, CC 20, Cess 20), 0 withdrawn, 60 completed (HTP 20, CC 20, Cess 20)		
	Withd	rawal reasons	s reported: N/A	
			stics: N=60; Mean Age (SD): HTP 32.7 (12.3) years, CC 30.9 (12.5) 6); Sex: 70% male; Ethnicity/Nationality: 100% Japanese.	
	Key in	clusion criter	ia: Health status: "good health"; ≥11 CPD; smoked for ≥1 year	
Interventions	Interv	terventions: HTP (novel tobacco vapor product), CC (own brand), smoking cessation		
	Co-int	Co-interventions: None		
	Mode	of exposure: 1	Direct ad libitum	
Outcomes	butano monoh hydrox aminoi aminoi	Primary: Exhaled Carbon monoxide, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, total N-nitrosonornicotine, Nicotine equivalents, monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, total 1-hydroxypyrene, S-phenylmercapturic acid, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene, 2-hydroxyethylmercapturic acid, 3-hydroxybenzo[a]pyrene, 1-aminonaphthalene, 4-hydroxybutyl-2-mercapturic acid		
	Dependent safety Topogram	endary: Daily product consumption, Fagerström Test for Nicotine/Cigarette endence, Physical examination, Clinical chemistry, haematology and urine analysis ty panel, Vital signs, Minnesota Nicotine Withdrawal Scale, Human Puffing/Smoking ography (inc. puff count), Product Liking Questionnaire, Adverse Events/Serious erse Events, Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of oking Urges		
Analyses	Prima at leas	Sample size calculation reported: Yes Primary analysis population: Full analysis set defined as "randomized subjects who had at least one BoE assessment after post-randomization" Unit of analysis: Individuals		
Study funding		-	national (Industry-affiliated)	
Notes			ression analysis. Data obtained from published literature.	
Risk of bias			rate in the second seco	
Bias		Authors'	Support for judgement	
generation	Random sequence generation		Beyond stating the study was 'randomised', no further information provided.	
	Allocation concealment		No information provided.	
and personnel			"Blinding: Open-no one is blinded". Included non-active comparator (cigarettes).	
assessment	Blinding of outcome assessment		"Blinding: Open-no one is blinded". All primary outcomes were objectively measured.	
Incomplete outco	ome	Low	All subjects randomised completed the study and were included in the analyses.	
Selective reporting	ng	Low	3 safety profile parameters were not reported, but adverse events data were reported. All other outcomes listed in the methods and on the trial registration are reported on in at least one literature source.	

Caponnetto,	2018
Caponinetto,	4010

Methods	Date of registration: Not registered
	Submitted to peer-reviewed journal within 12 months: Unclear
	Published key outcomes on trial registration within 12 months: Unclear
	Design: Crossover RCT
	Setting (Country): Confined (Unknown)
	Study start date; study end date: Not reported
	Intervention duration: 3 sessions of 2x 10 puffs with 30 sec intervals and 5 min interround break
Participants	Number of participants: 12 randomised, 0 withdrawn, 12 completed Withdrawal reasons reported: N/A

Baseline characteristics: N=12; Mean Age (SD): 28.6 years (SD not reported); Sex: 50%

	male; Ethnicity/Nationality: not reported				
	Key inclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥5 years				
Interventions	Interventions: HTP (IQOS), HTP (Glo), CC (Own brand)				
inter ventions	Co-interventions: None				
	Mode of exposure: Direct restricted				
0-4	Primary: Exhaled Carbon monoxide				
Outcomes		•	arbon monoxide		
	Secondary: N/A				
Analyses	_		tion reported: No		
	1		opulation: Not specified		
	Unit of	f analysis: Ind	lividuals		
Study funding	Univer	sity of Catania	a (Industry-affiliated)		
Notes	Include	ed in meta-reg	ression analysis. Data obtained from study authors.		
Risk of bias					
Bias		Authors'	Support for judgement		
		judgement			
Random sequence	e	Low	"The randomization sequence was computer-generated"		
generation Allocation conce	alment	Unclear	No information provided.		
Blinding of partic			No information on blinding. Included non-active comparator		
and personnel	cipunts	Unclear	(cigarettes).		
Blinding of outco	ome	Low	No information on blinding, but only outcome was objectively		
assessment		Low	measured.		
Incomplete outco	ome	Unclear	The authors state 12 subjects "took part" in the study but it is unclear		
data Selective reporting	200	Low	whether more than 12 were initially randomised. Only outcome measured (eCO) is reported on in the results.		
		Low	Only outcome measured (eCO) is reported on in the results.		
aspredicted.org	I				
Methods	I	f registration			
	I	_	reviewed journal within 12 months: Unclear		
	I	-	omes on trial registration within 12 months: Unclear		
	-	: Crossover R			
	_	-	Confined (Belgium)		
			ady end date: Not reported		
			on: 3 sessions of single use of one cigarette or tobacco stick		
Participants	l .		ants: randomised not reported, 0 withdrawn not reported, 34		
		completed			
	l	Withdrawal reasons reported: N/A Baseline characteristics: N=30; Mean Age (SD); 22 (3.09) years; Sex: 67% male;			
	l		Nationality: 14 Belgium, 16 Other		
	1	-	ia: Health status: cannot have "one or more severe medical		
			D; smoked for ≥3 years		
Interventions			(IQOS), CC (Own brand), EC (Eleaf iStick)		
21102 (011010115		erventions: N			
			Direct ad libitum		
Outcomes					
Outcomes		Primary: Exhaled Carbon monoxide, Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, Fagerström Test for Nicotine/Cigarette			
			ota Nicotine Withdrawal Scale, A visual analogue scale (VAS)		
	assessing cigarette craving, Product preference				
	1				
	1	lary: N/A			
Analyses	Second	lary: N/A	tion reported: No		
Analyses	Second Sample	lary: N/A e size calculat	tion reported: No opulation: Not specified or unclear		
Analyses	Second Sample Primar	lary: N/A e size calculat	opulation: Not specified or unclear		

Study funding	KU Le	uven and Thor	mas More University of Applied Sciences (Independent)	
Notes	Although number of participants randomised not reported, the authors stated 46 signed up for intake session. Also 34 completed all sessions, but 4 were excluded from the analyses for not meeting inclusion criteria. Included in meta-regression analysis. Data obtained from published literature.			
Risk of bias				
Bias		Authors' judgement	Support for judgement	
Random sequence generation	e	Unclear	Beyond stating the study was 'randomised', no further information provided.	
Allocation conce		Unclear	No information provided.	
Blinding of partic	•	Unclear	Presence of blinding not described. Included non-active comparator (cigarettes.	
Blinding of outco	ome	Unclear	Presence of blinding not described. Some primary outcomes were subjectively measured.	
Incomplete outco	ome	Unclear	The authors explained "46 signed up for the intake session, of whom 34 completed all sessions", but number of participants randomised was not reported.	
Selective reporting	ng	Low	All outcomes reported on in at least one literature source.	
NCT03435562				
Methods	Date o	f registration	: 19/02/2018	
	Submi	tted to peer-r	reviewed journal within 12 months: No publication	
	Publis	hed key outco	omes on trial registration within 12 months: Yes	
	Design	: Crossover R	CT	
	Setting	g (Country): (Confined (United States of America)	
	Study	start date; stı	udy end date: 03/03/2018; 16/09/2019	
	Interv bout	ention duratio	on: 3 sessions of a 10-puff product use bout and a 90 mins ad lib use	
Participants			ants: 22 randomised, 4 withdrawn, 18 completed	
	Withdrawal reasons reported: No Baseline characteristics: N=18; Mean Age (SD): 36.8 (9.3) years; Sex: 72% male; Ethnicity/Nationality: 7 Black or African America, 8 White, 2 more than one race, 1 unknown or not reported			
	Key in		ia: Health status: "healthy"; unspecified CPD; unspecified smoking	
Interventions	Interv	entions: HTP	(IQOS), CC (Own brand), EC (JUUL)	
		erventions: N		
	Mode	of exposure: 1	Direct restricted and direct ad libitum	
Outcomes		ry: Nicotine		
	Secondary: Exhaled Carbon monoxide, Questionnaire of Smoking Urges, Minnesota Nicotine Withdrawal Scale, Heart rate, The Direct Effects of Nicotine Questionnaire, Blood pressure			
Analyses	Sampl	e size calculat	tion reported: Yes	
•	_		opulation: Not specified or unclear	
		f analysis: Ind		
Study funding	Virginia Commonwealth University and National Institute on Drug Abuse, Center for the Study of Tobacco Products (Independent)			
Notes			ression analysis. Data obtained from published literature.	
Risk of bias	1			
Bias		Authors'	Support for judgement	
Dias		judgement	Support for Judgement	
Random sequence generation	e	Low	"Order of the products used in each session will be assigned using Latin-square order procedure"	
-				

Allocation conce	alment	Unclear	No information provided.	
Dlinding of participants			"Masking: None (Open Label)". Included non-active comparator	
and personnel		High	(cigarettes).	
Blinding of outco	ome	Low	"Masking: None (Open Label)". Primary outcome objectively measured.	
Incomplete outco	ome	Low	Overall attrition = 18.18%. All participants who completed the study were included in the analysis.	
Selective reporting	ng	High	Results data for heart rate and blood pressure have not been reported.	
NCT03889990/N	NCT039	95329		
Methods	Date o	f registration	: 26/03/2019 (NCT03889990); 24/06/2019 (NCT03995329)	
	Submi	tted to peer-r	reviewed journal within 12 months: Yes	
	Publis	hed key outco	omes on trial registration within 12 months: No results posted	
	Design	: 2 non-rando	mised single group assignment trials	
	Setting	g (Country): (Confined (Greece)	
		start date; sti 2019 (NCT039	udy end date: 01/01/2018; 01/01/2019 (NCT03889990), 19/06/2019; 995329)	
	Interv	vention duration: 1 session of up to 14 puffs over 5-6 mins		
Participants	Numb	er of particip	ants: 65 enrolled, 0 withdrawn, 50 completed	
	Withd	hdrawal reasons reported: No		
			stics: N=50; Mean Age (SD): Smokers 40.3 (13.2) years, Non-	
		ers 37.4 (10.4) years; Sex: 100% male; Ethnicity/Nationality: not reported inclusion criteria: Health status: "healthy"; ≥5 pack years		
T				
Interventions		Interventions: HTP (IQOS) in smokers and non-smokers		
		aterventions: None e of exposure: Direct restricted		
		_		
Outcomes	Primary: Exhaled Carbon monoxide, Forced expiratory volume in one second, Forced vital capacity, Forced expiratory flow at 25–75% of forced vital capacity, Total lung capacity, Residual volume, Forced expiratory volume in one second/forced vital capacity, Heart rate, Functional residual capacity, Diffusion Capacity, Peak Expiratory Flow, [Mean] Arterial Blood Pressure, Total respiratory resistances, Respiratory impedance, Oxygen Saturation, Maximal Mid-Expiratory Flow, Expiratory reserve volume Secondary: N/A			
Analyses	_		tion reported: Yes	
	Prima	ry analysis po	opulation: Not specified or unclear	
	Unit of	f analysis: Inc	lividuals	
Study funding	Aristot	le University	Of Thessaloniki (Independent)	
Notes	The authors reported enrolling 25 subjects in each trial, but on the registration of one trial (NCT03889990) it was reported that 40 participants had in fact enrolled. It is not clear when or why 15 subjects were removed from the study. Not included in meta-regression analysis.			
Risk of bias				

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation	High	Non-randomised trial.
Allocation concealment	High	Non-randomised trial.
Blinding of participants and personnel	Low	Both arms received the same intervention, and the arms were from two separately conducted single-group assignment trials.
Blinding of outcome assessment	Low	All primary outcomes were objectively measured.
Incomplete outcome data	Low	NCT03889990 attrition=37.5%; NCT03995329 attrition =0%, but both arms received the same intervention.
Selective reporting	High	Blood pressure and heart rate were listed as primary outcomes on the non-smoker trial registration (NCT03995329) but results data for these have not been reported.

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NCT03301129			
Methods	Submi Publish Design Setting Study	hed key outco :: Crossover R g (Country): (start date; stu	reviewed journal within 12 months: Yes omes on trial registration within 12 months: No results posted
Participants	Number of participants: 20 randomised, 0 withdrawn, 20 completed Withdrawal reasons reported: N/A Baseline characteristics: N=20; Mean Age (SD): 35 (13) years; Sex: 30% male; Ethnicity/Nationality: not reported Key inclusion criteria: Health status: "healthy"; unspecified CPD; unspecified smoking duration		
Interventions	Co-int	erventions: N	(IQOS2.2), CC (Marlboro Gold), EC (Blu Pro) Jone Direct ad libitum
Outcomes	Primary: Soluble Nox2-derived peptide, Flow-mediated dilation Secondary: Cotinine, Vitamin E, Soluble P-selectin, Soluble CD40 ligand, nitric oxide bioavailability, H2O2 production, H2O2 breakdown activity, Systolic blood pressure, Diastolic blood pressure, 8-iso-prostaglandin F2alpha, Product Satisfaction Questionnaire		
Analyses	Sample size calculation reported: Yes Primary analysis population: Not specified or unclear Unit of analysis: Individuals		
Study funding	University of Roma La Sapienza (Independent)		
Notes	Not inc	cluded in meta	regression analysis.
Risk of bias			
Bias		Authors' judgement	Support for judgement
Random sequence generation		Low	"The randomization list was computer generated"
Allocation concealment Blinding of participants and personnel		Unclear	No information provided. Despite describing the trial as "Double" blinded on the trial registration, only "Investigator" and "Outcome Assessor" are noted as being masked, not participants.
Blinding of outco	ome	Low	"Masking: Double (Investigator, Outcomes Assessor)". Primary outcomes were objectively measured
Incomplete outco	ome	Low	The 30 subjects excluded were excluded pre-randomisation. No subjects who were randomised withdrew or were excluded from the final analysis population.
Selective reportin	ng	Low	All outcomes reported on in at least one literature source.
Methods	Submi Publish Design Setting Study	hed key outco a: Parallel RCT g (Country): A start date; sta	reviewed journal within 12 months: No publication omes on trial registration within 12 months: No T Ambulatory (Japan) udy end date: 07/11/2017; 12/06/2019
Participants	Number 170 cos	mpleted (86 H	ants: 172 randomised (87 HTP, 85 CC), 2 withdrawn (1 HTP, 1 CC),

	Baseline characteristics: N=172; Mean Age (SD): HTP 48.1 years, CC 46.5 years, Dual Use 54.4 years, Other use 54 years (SDs not reported); Sex: 81% male; Ethnicity/Nationality: 100% Japanese Key inclusion criteria: Health status: must have generalized chronic periodontitis; ≥10 CPD; smoked for ≥5 years
Interventions	Interventions: HTP (IQOS), CC (Own brand)
	Co-interventions: Mechanical periodontal therapy
	Mode of exposure: Direct ad libitum
Outcomes	Primary: Mean Periodontal Pocket Depth (PD) reduction in all sites with initial PD ≥ 4 mm after mechanical periodontal therapy Secondary: Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, Nicotine equivalents, Daily product consumption, Adverse Events/Serious Adverse Events, Mean PD change in sites with initial PD≥4 mm after mechanical periodontal therapy, mean PD change in sites with initial PD<4mm, 4-5 mm, 5-6 mm, 6-7 mm, and ≥7 mm, Mean clinical attachment level (CAL) change in sites with initial PD≥4mm after mechanical periodontal therapy, mean CAL change in sites with initial PD<4 mm, 4-5 mm, 5-6 mm, 6-7 mm, and ≥7mm, change in tooth mobility (grade), change in the number of sites with PD<4 mm, 4-5mm, 5-6 mm, 6-7 mm, and ≥7 mm, change in plague control record, change in mean full-mouth PD, change in mean full-mouth CAL, change in gingival inflammation (GI) score, change in bleeding on probing scores Pro-inflammatory and immuno-regulatory mediators (sCD40L, CRP, EGF,
	Eotaxin/CCL11, Flt3 ligand, GM-CSF, GRO, IFNα2, IL-1α, IL-1β, IL-1Ra, IL-2, IL-3, IL-4, IL-5, IL-6, IL-7, IL-8/CXCL8, IL-9, IL-10, IL-12 (p40), IL-12 (p70), IL-13, IL-15, IL-17A/CTLA8, IP10/CXCL10, MCP-1/CCL2, MCP-3/CCL7, MDC/CCL22, MIP-1α/CCL3, MIP-1β/CCL4, MMP-1, MMP-8, MMP-9, MMP-10, MMP-12, MMP-13, osteoprotegerin, PDGF-AA, PDGF-AB/BB, RANKL, RANTES/CCL5, TGFα, TIMP-1, TNFα, TNFβ/LT-α), Microbiological status, Full transcriptomics profile
Analyses	Sample size calculation reported: Yes
·	Primary analysis population: Full analysis set (as exposed) defined as "all randomized subjects with at least one product use experience and at least one valid non-safety assessment. Subjects were analyzed based on their actual self-reported product use. Some participants were excluded from analysis for protocol deviations (including, but not limited to, missing measurements)" Unit of analysis: Individuals
Study funding	Philip Morris International (Industry-affiliated)
Notes	Not included in meta-regression analysis.
	1 Not included in incla-regression analysis.
Risk of bias	

Risk of bias	Risk of bias			
Bias	Authors' judgement	Support for judgement		
Random sequence generation	Low	"Randomization will be done through the Interactive Web and Voice Response System (IXRS)"		
Allocation concealment	Low	"Randomization will be done through the Interactive Web and Voice Response System (IXRS)"		
Blinding of participants and personnel	High	"Masking: Single (Investigator)". Included non-active comparator (cigarettes).		
Blinding of outcome assessment	Low	"Masking: Single (Investigator)". Primary outcome objectively assessed.		
Incomplete outcome data	Low	Attrition: IQOS=1.15% CC=1.18%, overall=1.16%. Exclusion: IQOS=19.54% CC=1.18%, overall=1.74%.		
Selective reporting	High	The following outcomes listed in the protocols have not been reported on: measurement of pro-inflammatory and immunoregulatory mediators; microbiological status; full transcriptomics profile.		
Other	High	Only reported data grouped by participant product use not randomisation.		

Methods	Date of registration: 29/12/2015				
	Submitted to peer-reviewed journal within 12 months: No				
	Published key outcomes on trial registration within 12 months: No				
	Design: Parallel RCT				
	Setting (Country): Confined & Ambulatory (Poland)				
	Study start date; study end date: January 2016; July 2017				
	Intervention duration: 90 Days (5 days confinement + 85 days ambulatory)				
Participants	Number of participants: 120 randomised (80 HTP, 40 CC), 5 withdrawn (4 HTP, 1 CC), 115 completed (76 HTP, 39 CC)				
	Withdrawal reasons reported: Yes				
	Baseline characteristics: N=120; Mean Age (SD): HTP 38.9 (8.9) years, CC 39.0 (8.0) years; Sex: 53% male; Ethnicity/Nationality: 100% Caucasian Key inclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥10 years				
Interventions					
interventions	Interventions: HTP (carbon heated tobacco product 1.2), CC (Own brand) Co-interventions: None				
~ .	Mode of exposure: Direct ad libitum				
Outcomes	Primary: S-phenylmercapturic acid, monohydroxybutenylmercapturic acid, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 3-hydroxypropylmercapturic acid, Carboxyhemoglobin				
	Secondary: 2-cyanoethylmercapturic acid, Modified Cigarette/Product Evaluation				
	Questionnaire, Questionnaire of Smoking Urges, total N-nitrosonornicotine, Nicotine				
	equivalents, Exhaled Carbon monoxide, total 1-hydroxypyrene, o-toluidine, 4-				
	aminobiphenyl, 2-aminonaphthalene, Daily product consumption, Fagerström Test for				
	Nicotine/Cigarette Dependence, 3-hydroxy-1-methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, Respiratory symptoms (inc. cough assessment), Nicotine,				
	Physical examination, Clinical chemistry, haematology and urine analysis safety panel,				
	Vital signs, Electrocardiogram, 3-hydroxybenzo[a]pyrene, Spirometry, Concomitant				
	medications, Cotinine, 1-aminonaphthalene, 8-epi-prostaglandin F2alpha, 11-				
	dehydrothromboxane B2, Cytochrome P450 2A6 activity, Ames mutagenicity test				
	(YG1024+S9), White blood cell count, Systolic blood pressure, Soluble intercellular				
	adhesion molecule-1, High-density lipoprotein cholesterol, Forced expiratory volume in				
	one second, Diastolic blood pressure, Weight, Waist circumference, Low-density				
	lipoprotein cholesterol, Homocysteine, High-sensitivity C-reactive protein, Fibrinogen,				
	Platelet count, Hemoglobin glycosylated (Hemoglobin A1C), Forced vital capacity, Forced expiratory flow at 25–75% of forced vital capacity, Triglycerides, Total cholesterol,				
	Apolipoprotein B, Apolipoprotein A1, Blood glucose, Forced expiratory volume in one				
	second/forced vital capacity, Myeloperoxidase, Intention to use [HTP] Questionnaire, Total				
	anti-oxidant capacity, 8-Hydroxy-2'-deoxyguanosine, Prochaska "Stage of Change"				
	Questionnaire, 4-Hydroxy-2-nonenal, Adverse Events/Serious Adverse Events				
Analyses	Sample size calculation reported: Yes				
	Primary analysis population: Per-protocol population defined as "randomized subjects				
	who fulfilled product adherence criteria and had no major protocol deviations impacting				
	evaluability, such as violation of eligibility criteria or insufficient duration of urine				
	collection. Separate PP populations were defined for the analysis at Day 5 and Day 90.				
	Non-adherence to CHTP was defined as an average cigarette use of > 0.5 cigarettes/day from Day 1 to the end of the respective period (Day 5 or Day 90) or use of > 2 cigarettes				
	on a single day within a week prior to the assessments."				
	Unit of analysis: Individuals				
Study funding	Philip Morris International (Industry-affiliated)				
Notes	Not included in meta-regression analysis.				
Risk of bias					
Bias	Authors' Support for judgement				

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Random sequence generation			"subjects will be randomized using an interactive web and voice response system (IxRS)"			
Allocation concealment		Low	"subjects will be randomized using an interactive web and voice response system (IxRS)"			
Blinding of parti and personnel	Blinding of participants and personnel		"Masking: None (Open Label)". Included non-active comparator (cigarettes).			
Blinding of outcomes assessment		Low	"Masking: None (Open Label)". All primary outcomes objectively measured.			
Incomplete outco data	ome	Low	Attrition: IQOS=5% CC=2.5%, overall=4.17%. Exclusion: IQOS=3.75% CC=12.5%, overall=6.6%.			
Selective reporti	Selective reporting		"Samples for 4-HNE analysis have been collected but will not be analyzed due to the failure to develop a selective and quantitative assay." QSU, Intent to Use of CHTP 1.2, Prochaska "Stage of Change" Questionnaire, MCEQ, and pre- and post-bronchodilator FVC, FEV1/FVC, FEF 25-75 were not reported in any literature sources.			
NCT02396381						
Methods	Date o	f registration	: 24/03/2015			
		_	reviewed journal within 12 months: No			
	1	•	omes on trial registration within 12 months: No			
	_	: Parallel RC				
	1	ting (Country): Ambulatory (United States of America)				
	1	y start date; study end date: 12/03/2015; 01/08/2017 vention duration: 26 weeks				
Participants			ants: 984 randomised (488 HTP, 496 CC), 127 withdrawn (74 HTP,			
Farticipalits		C), 857 completed (414 HTP, 443 CC)				
	Withdrawal reasons reported: Yes					
	Baseli years, Ethnic Indian		aseline characteristics: N=857; Mean Age (SD): HTP 44.2 (9.64) years, CC 45.2 (9.55) ears, Dual Use 43.8 (9.77) years, Other use 44.2 (8.14) years; Sex: 58.8% male; thnicity/Nationality: 79.2% White, 17.6% Black or African American, 0.7% American idian or Alaska Native, 0.9% Asian, 0.4% Native Hawaiian or Other Pacific Islander, 2% unknown or not reported			
			ia: Health status: "healthy"; ≥10 CPD; smoked for ≥1 year			
Interventions			terventions: HTP (IQOS2.2), CC (Own brand)			
	Co-int	o-interventions: None				
	Mode	of exposure: 1	Direct ad libitum			
Outcomes	(methyl Soluble expirate Seconda Questio		Primary: 8-epi-prostaglandin F2alpha, 11-dehydrothromboxane B2, Total 4-methylnitrosamino)-1-(3-pyridyl)-1-butanol, Carboxyhemoglobin, White blood cell count, Soluble intercellular adhesion molecule-1, High-density lipoprotein cholesterol, Forced expiratory volume in one second			
			Secondary: 2-cyanoethylmercapturic acid, Modified Cigarette/Product Evaluation Questionnaire, total N-nitrosonornicotine, Nicotine equivalents,			
hydro Deper cough urine Spiro blood Diaste chole Hemo		monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, total 1-hydroxypyrene, Daily product consumption, Fagerström Test for Nicotine/Cigarette Dependence, 3-hydroxy-1-methylpropylmercapturic acid, Respiratory symptoms (inc. cough assessment), Nicotine, Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, 3-hydroxybenzo[a]pyrene, Spirometry, Concomitant medications, Cotinine, Cytochrome P450 2A6 activity, Systolic blood pressure, Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Diastolic blood pressure, Weight, Waist circumference, Low-density lipoprotein cholesterol, Homocysteine, High-sensitivity C-reactive protein, Fibrinogen, Platelet count, Hemoglobin glycosylated (Hemoglobin A1C), Forced vital capacity, Forced expiratory flow at 25–75% of forced vital capacity, Apolipoprotein B, Apolipoprotein A1, Total lung				
	capaci		relaperayidase Vital capacity, Inspiratory capacity, Functional residual capacity,			

Myeloperoxidase, Vital capacity, Inspiratory capacity, Functional residual capacity, Intention to use [HTP] Questionnaire, bronchodilator reversibility in FEV1, Albumin

Analyses | Sample size calculation reported: Yes

Primary analysis population: Full analysis set (as exposed) defined as "Subjects in FAS-AR who had at least 1 record of reported product use diary post-randomization. The exposure assignment was actual product exposure, as defined by the product use pattern categories estimated during the 6 month period: •THS-use: ≥ 1 THS or CC, and $\geq 70\%$ THS use over the analysis period, and $\geq 70\%$ THS use on >50% of days in the analysis period •Dual-use: ≥ 1 THS or CC and, 1% \leq THS<70% over the analysis period, or THS-use and CC-use categories do not apply to 50% of these days •CC-use: ≥ 1 THS or CC use, and <1% THS use over the entire analysis period and <1% THS use on $\geq 50\%$ of days in the analysis period. •Other-use: Subjects with missing product use, or using e-cigarettes or other tobacco products, quitters, or subjects who switched across different use patterns between consecutive analysis periods"

Unit of analysis: Individuals

Study funding

Philip Morris International (Industry-affiliated)

Notes

Included in meta-regression analysis. Data obtained from published literature.

Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation	Low	"Randomization was done through the interactive voice and web response system (IXRS)"	
Allocation concealment	Low	"Randomization was done through the interactive voice and web response system (IXRS)"	
Blinding of participants and personnel	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).	
Blinding of outcome assessment	Low	"Masking: None (Open Label)". All primary outcomes objectively measured	
Incomplete outcome data	Low	Attrition: IQOS=15.16% CC=10.69%, overall=2.91%. Although not the main analysis population, full analysis set (as randomised) results data were also presented in the published literature.	
Selective reporting	Low	All outcomes reported on in at least one literature source.	

NCT02466412

110102100112	
Methods	Date of registration: 09/06/2015
	Submitted to peer-reviewed journal within 12 months: No publication
	Published key outcomes on trial registration within 12 months: No
	Design: Crossover RCT
	Setting (Country): Confined (Japan)
	Study start date; study end date: 08/05/2015; November 2015
	Intervention duration: 2 sessions of single use of one cigarette or tobacco stick
Participants	Number of participants: 48 randomised (24 HTP-CC, 24 CC-HTP), 0 withdrawn, 48 completed (24 HTP-CC, 24 CC-HTP)
	Withdrawal reasons reported: N/A
	Baseline characteristics: N=47; Mean Age (SD): HTP-CC 44.7 (10.03) years, CC-HTP 40.7 (11.48) years; Sex: 47% male; Ethnicity/Nationality: 100% Japanese
	Key inclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years
Interventions	Interventions: HTP (carbon heated tobacco product 1.1 M), CC (Own brand M)
	Co-interventions: None
	Mode of exposure: Direct ad libitum
Outcomes	Primary: Maximal nicotine concentration, Area under the concentration curve from start of product use to time of last quantifiable concentration
	Secondary: Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, Carboxyhemoglobin, Respiratory symptoms (inc. cough assessment), Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Time to reach nicotine Cmax
	Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Spirometry

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	Concomitant medications, Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity, Partial AUC, AUC from start of product use up to 12 hours, Terminal half-life				
Analyses	Sample size calculation reported: Yes				
	Primary analysis population: Pharmacokinetic population defined as "all the randomized subjects who give informed consent, completed at least one of the single use Day 1 or Day 3, and for whom at least one PK parameter can been derived. Only subjects without major protocol deviations that impact evaluability of the data (to be defined in the SAP) will be included in the PK analysis sets" Unit of analysis: Individuals				
C4 J £ J:					
Study funding	_		tional (Industry-affiliated)		
Notes	concer		ed from the analyses (sequence HTP-CC) due to all plasma nicotine rements being below the quantification limit. Not included in meta-		
Risk of bias					
Bias		Authors' judgement	Support for judgement		
Random sequence generation	ce	Low	"Randomization to product exposure sequence will be done through IxRS"		
Allocation conce		Low	"Randomization to product exposure sequence will be done through IxRS"		
Blinding of parti and personnel	-	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).		
Blinding of outco		Low	"Masking: None (Open Label)". All primary outcomes were objectively measured.		
Incomplete outco	ome	Low	Attrition was 0%. Exclusion: mCHTP-mCC=4.16% mCC-mCHTP=0%, overall=2.1%		
Selective reporti	ng	High	Only results data for the two primary outcomes have thus far been published.		
NCT02503254					
Methods	Date of registration: 20/07/2015 Submitted to peer-reviewed journal within 12 months: No Published key outcomes on trial registration within 12 months: Yes Design: Parallel RCT Setting (Country): Confined (Poland) Study start date; study end date: 04/07/2015; March 2016				
	1	ention durati	-		
Participants		Number of participants: 80 randomised (41 HTP, 39 CC), 0 withdrawn, 80 completed (41 HTP, 39 CC)			
Baseli (10.97		Withdrawal reasons reported: N/A			
		Baseline characteristics: N=80; Mean Age (SD): HTP 34.1 (10.45) years, CC 32.7 (10.97) years; Sex: 49% male; Ethnicity/Nationality: 100% Caucasian			
	-		ia: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years		
Interventions			(carbon heated tobacco product 1.0), CC (Own brand)		
		erventions: N			
_		_	Direct ad libitum		
Outcomes	Carbox	Primary: monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, Carboxyhemoglobin, S-phenylmercapturic acid			
	Secondary: Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of				

Smoking Urges, total N-nitrosonornicotine, Nicotine equivalents, Exhaled Carbon monoxide, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, total 1-hydroxypyrene, Adverse Events/Serious Adverse Events, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene, Daily product consumption, 3-hydroxy-1-methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, Respiratory symptoms (inc. cough assessment), Nicotine, Physical examination, Clinical chemistry, haematology

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	and urine analysis safety panel, Vital signs, 3-hydroxybenzo[a]pyrene, Spirometry, Concomitant medications, Cotinine, 1-aminonaphthalene, Minnesota Nicotine Withdrawal Scale, Cytochrome P450 2A6 activity, Human Puffing/Smoking Topography (inc. puff count), Ames mutagenicity test (YG1024+S9), Human Puffing/Smoking Topography Questionnaire				
Analyses	Sample size calculation reported: Yes Primary analysis population: Full analysis set defined as "all the randomized subjects who had at least one post-randomization product use experience, if randomized to CHTP 1.0 or CC, and have at least one valid non safety assessment." Unit of analysis: Individuals				
Study funding	Philip N	Morris Interna	tional (Industry-affiliated)		
Notes	Not incl	luded in meta	-regression analysis.		
Risk of bias					
Bias		Authors' judgement	Support for judgement		
Random sequence generation	e	Low	"subjects were randomized by an interactive web and voice response system"		
Allocation conce		Low	"subjects were randomized by an interactive web and voice response system"		
Blinding of partiand personnel	cipants	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).		
Blinding of outco	ome	Low	"Masking: None (Open Label)". All primary outcomes objectively measured.		
Incomplete outco	ome	Low	Attrition and exclusion both 0%.		
Selective reporting	ng	High	Several outcomes listed in the study protocol were not reported on in the main results article. Only one was reported on in a poster instead.		
NCT02649556			•		
Methods	Date of registration: 07/01/2016 Submitted to peer-reviewed journal within 12 months: No publication Published key outcomes on trial registration within 12 months: No Design: Parallel RCT Setting (Country): Ambulatory (United States of America) Study start date; study end date: 30/09/2015; 20/12/2017 Intervention duration: 26 weeks				
Participants	Number of participants: 857 started (230 HTP, 424 CC, 152 Dual use, 51 Other use), 248 withdrawn (63 HTP, 112 CC, 50 Dual use, 23 Other use), 609 completed (167 HTP, 312 CC, 102 Dual use, 28 Other use) Withdrawal reasons reported: No Baseline characteristics: N=857; Mean Age (SD): HTP 43.8 (9.68) years, CC 45.2 (9.54) years, Dual use 44.2 (9.76) years, Other use 44.5 (8.21) years; Sex: 58.8% male; Ethnicity/Nationality: 79.2% White, 17.6% Black or African American, 0.7% American Indian or Alaska Native, 0.9% Asian, 0.4% Native Hawaiian or Other Pacific Islander, 1.2% unknown or not reported Key inclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥1 year				
Interventions	Co-inte	erventions: N	(IQOS2.2), CC (Own brand) fone Direct ad libitum		
Outcomes	1-(3-pyi intercell volume Second	Primary: Carboxyhemoglobin, 8-epi-prostaglandin F2alpha, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 11-dehydrothromboxane B2, White blood cell count, Soluble intercellular adhesion molecule-1, High-density lipoprotein cholesterol, Forced expiratory volume in one second Secondary: Modified Cigarette/Product Evaluation Questionnaire, total N-			
		nitrosonornicotine, Nicotine equivalents, Daily product consumption, Fagerström Test for Nicotine/Cigarette Dependence, Respiratory symptoms (inc. cough assessment), Nicotine,			

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Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Concomitant medications, Cotinine, Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Weight, Waist circumference, Lowdensity lipoprotein cholesterol, Homocysteine, High-sensitivity C-reactive protein, Fibrinogen, Platelet count, Hemoglobin glycosylated (Hemoglobin A1C), Forced vital capacity, Forced expiratory flow at 25–75% of forced vital capacity, Apolipoprotein B, Apolipoprotein A1, Total lung capacity, Forced expiratory volume in one second/forced vital capacity, Myeloperoxidase, Vital capacity, Inspiratory capacity, Functional residual capacity, Intention to use [HTP] Questionnaire, bronchodilator reversibility in FEV1, Albumin, Blood pressure

Analyses

Sample size calculation reported: Yes

Primary analysis population: Full analysis set (as exposed) defined as "The FAS-EX consists of all subjects in FAS-AR who have at least one record of reported product use diary post randomization. The exposure assignment for the FAS-EX will be actual product exposure, as defined by the product use pattern categories estimated during the 12 month period JV4, V16"But note "Some participants were excluded from analysis for protocol deviations (including, but not limited to, missing measurements)."

Unit of analysis: Individuals

Study funding

Philip Morris International (Industry-affiliated)

Notes

This is an extension to NCT02396381. 672 (309 in the THS arm and 363 in the CC arm) subjects enrolled in the extension study; the 857 subjects in the Full Analysis Set - As Exposed (FAS-EX) included subjects for combined analyses from the original six-month study who did not enter the extension study. The analysis was performed according to subjects' exposure over the 12-month period. Not included in meta-regression analysis.

Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation	Low	"Randomization was done during the original study at V4 through the interactive voice and web response system (IXRS)."	
Allocation concealment	Low	"Randomization was done during the original study at V4 through the interactive voice and web response system (IXRS)."	
Blinding of participants and personnel	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).	
Blinding of outcome assessment	Low	"Masking: None (Open Label)". All primary outcomes objectively measured.	
Incomplete outcome data	Unclear	672 subjects enrolled into the extension study (309 THS and 363 CC). However, it is unclear how many completed the study as the data is combined with the previous 6-month trial.	
Selective reporting	High	Only results data for the primary outcomes have been published.	
Other	High	Only reported data grouped by participant product use not randomisation.	

NCT01967706

Methods Date of registration: 23/10	/20	1
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Submitted to peer-reviewed journal within 12 months: No publication **Published key outcomes on trial registration within 12 months:** No

Design: Crossover RCT

Setting (Country): Confined (Japan)

Study start date; study end date: 01/08/2013; May 2014

Intervention duration: 2 sessions of single use of one cigarette, tobacco stick or piece of

gum for 35 ± 5 mins

Participants

Number of participants: 62 randomised (44 HTP/CC, 18 HTP/NRT), 1 withdrawn (1 HTP/CC), 61 randomised (43 HTP/CC, 18 HTP/NRT)

Withdrawal reasons reported: Yes

Baseline characteristics: N=61; Mean Age (SD): HTP/CC 33.4 (10.03) years, HTP/NRT

30.7 (7.8) years; Sex: 52% male; Ethnicity/Nationality: 100% Japanese

Key inclusion criteria: Health status: "healthy"; \geq 10 CPD; smoked for \geq 3 years

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Interventions	Interventions: HTP (IQOS2.2 M), CC (Own brand M), NRT (Nicorette Gum) Co-interventions: None				
Outcomes	Mode of exposure: Direct ad libitum Primary: Maximal nicotine concentration, Area under the concentration curve from start of product use to time of last quantifiable concentration Secondary: Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, Carboxyhemoglobin, Fagerström Test for Nicotine/Cigarette Dependence, Respiratory symptoms (inc. cough assessment), Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Time to reach nicotine Cmax, Spirometry, Concomitant medications, Terminal half-life of nicotine, Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity, Partial AUC, Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events Sample size calculation reported: Yes Primary analysis population: Pharmacokinetic population defined as "all the randomized subjects who give informed consent, completed at least one of the single use Day 1 or Day 3, and for whom at least one PK parameter can been derived. Only subjects without major protocol deviations (to be defined in the SAP) will be included"				
		f analysis: Inc			
Study funding	_		ntional (Industry-affiliated)		
Notes	Include	ed in meta-reg	ression analysis. Data obtained from published literature.		
Risk of bias					
Bias		Authors' judgement	Support for judgement		
Random sequence generation	e	Low	"Randomization to product exposure sequence was done through an Interactive Telephone and Web Response System"		
Allocation conce		Low	"Randomization to product exposure sequence was done through an Interactive Telephone and Web Response System"		
Blinding of partic and personnel		High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).		
Blinding of outco		Low	"Masking: None (Open Label)". All primary outcomes objectively measured.		
data			Attrition: IQOS-CC=2.27% IQOS-NRT=0%, overall=1.61%. No subjects who completed the study were excluded from the analysis.		
Selective reporting	ng	Low	All outcomes reported on in at least one literature source.		
NCT01780688					
Methods		f registration			
		-	reviewed journal within 12 months: No		
		•	omes on trial registration within 12 months: No results posted		
		: Crossover R			
			(Country): Confined (United Kingdom)		
	•		study start date; study end date: May 2012; December 2012		
	Intervention duration: 2 sessions of single use of one cigarette or tobacco stick and 1 da of <i>ad lib</i> use				
Participants			ants: 28 randomised (14 HTP-CC, 14 CC-HTP), 0 withdrawn, 28 CC, 14 CC-HTP)		
	Withd	rawal reasons	s reported: N/A		
	Baseline characteristics: N=28; Mean Age (SD): HTP-CC 30.0 (4.9) years, CC-H' (4.0) years; Sex: 50% male; Ethnicity/Nationality: 100% Caucasian		stics: N=28; Mean Age (SD): HTP-CC 30.0 (4.9) years, CC-HTP 29.1 % male; Ethnicity/Nationality: 100% Caucasian		
	Key inclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years				
Interventions	Interventions: HTP (IQOS2.1), CC (Own brand)				

Co-interventions: None

Mode of exposure: Direct restricted and ad libitum

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Primary: Maximal nicotine concentration, Area under the concentration curve from start

Outcomes

	of product use to time of last quantifiable concentration					
	_	Secondary: Questionnaire of Smoking Urges, Fagerström Test for Nicotine/Cigarette				
		Dependence, Respiratory symptoms (inc. cough assessment), Time to reach nicotine Cmax,				
		Terminal half-life of nicotine, Time to nicotine Cpeak, Maximum observed nicotine				
			ving ad lib use), Lowest observed plasma concentration during the			
			val in which Cpeak was observed, Adverse Events/Serious Adverse			
	Events, Modified Cigarette/Product Evaluation Questionnaire					
Analyses	Sample size calculation reported: Yes					
			opulation: Per-protocol population defined as "all randomized			
		ects who did not deviate from the protocol, who completed at least one of the single use I libitum days, and had at least one estimable pharmacokinetic parameter derived				
		the single or ad libitum days"				
	Unit of	of analysis: Individuals				
Study funding	Philip I	Morris Interna	tional (Industry-affiliated)			
Notes	Not inc	cluded in meta	-regression analysis.			
Risk of bias						
Bias		Authors'	Support for judgement			
		judgement	The state of the s			
Random sequence generation	e	Low	"Randomization was performed using an Interactive Web Response System"			
Allocation conce	alment	Low	"Randomization was performed using an Interactive Web Response System"			
Blinding of partic	cipants	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).			
Blinding of outco	ome	Low	"Masking: None (Open Label)". All primary outcomes objectively measured.			
Incomplete outco	ome		All participants randomised completed the trial and no participants			
data	,,,,,	Low	were excluded from the analysis.			
Selective reporting	ng	Low	All outcomes reported on in at least one literature source.			
NCT01780714						
Methods	Date of	f registration	: 31/01/2013			
	Submi	tted to peer-r	reviewed journal within 12 months: No			
		-	omes on trial registration within 12 months: No results posted			
	_	esign: Parallel RCT				
	_	etting (Country): Confined (Poland)				
		udy start date; study end date: June 2012; December 2012 tervention duration: 5 days				
			•			
Participants		Number of participants: 40 randomised (20 HTP, 20 CC), 0 withdrawn, 40 completed (20 HTP, 20 CC)				
		Withdrawal reasons reported: N/A				
	Key in		·			
	Interventions: HTP (IQOS2.1), CC (Own brand)					
Interventions		4. 33	Co-interventions: None			
Interventions	Co-inte					
	Co-into	of exposure:	Direct ad libitum			
Interventions Outcomes	Co-into Mode o Prima	of exposure: l ry: monohydr	Direct ad libitum oxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid,			
	Co-into Mode of Primar Carbox	of exposure: lary: monohydr xyhemoglobin,	Direct <i>ad libitum</i> oxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, S-phenylmercapturic acid			
	Co-into Mode of Primar Carbox Second	of exposure: lary: monohydrayhemoglobin, lary: Question	Direct <i>ad libitum</i> oxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, S-phenylmercapturic acid nnaire of Smoking Urges, total N-nitrosonornicotine, Nicotine			
	Co-into Mode of Primar Carbox Second equival	of exposure: I ry: monohydr ryhemoglobin, lary: Question lents, total 1-h	Direct <i>ad libitum</i> oxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, S-phenylmercapturic acid nnaire of Smoking Urges, total N-nitrosonornicotine, Nicotine ydroxypyrene, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene,			
	Co-into Mode of Primar Carbox Second equival Daily p Respira	of exposure: 1 ry: monohydr ryhemoglobin, lary: Question lents, total 1-h product consur atory sympton	Direct <i>ad libitum</i> oxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, S-phenylmercapturic acid nnaire of Smoking Urges, total N-nitrosonornicotine, Nicotine ydroxypyrene, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene, nption, Fagerström Test for Nicotine/Cigarette Dependence, ns (inc. cough assessment), Nicotine, Cotinine, 11-			

	activity, Human Puffing/Smoking Topography (inc. puff count), Adverse Events/Serious Adverse Events, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, Modified Cigarette/Product Evaluation Questionnaire		
Analyses	Sample size calculation reported: Yes		
	Primary analysis perecord of at least one assessment"		opulation: Full analysis set defined as "randomized subjects who had e post-randomization product use and at least one valid biomarker
C4 - 1 - C 1'		f analysis: Inc	
Study funding			ational (Industry-affiliated)
Notes	Not inc	cluded in meta	regression analysis.
Risk of bias			
Bias		Authors' judgement	Support for judgement
Random sequence generation		Unclear	Beyond stating the study was 'randomised', no further information provided.
Allocation conce		Unclear	No information provided.
Blinding of partical and personnel	cıpants	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).
Blinding of outco	ome	Low	"Masking: None (Open Label)". All primary outcomes objectively measured.
Incomplete outco	ome	Low	All participants randomised completed the trial and no participants were excluded from the analysis.
Selective reporting	ng	High	Data for 4 outcomes listed in the protocol (Cytochrome P450 2A6 activity, Questionnaire of Smoking Urges, Minnesota Nicotine Withdrawal Scale, Respiratory symptoms) were not reported.
ISRCTN886824	35		windra war board, respiratory symptoms, were not reported.
Methods	1	f registration	• 06/10/2015
	Date of registration: 06/10/2015 Submitted to peer-reviewed journal within 12 months: No publication Published key outcomes on trial registration within 12 months: No results posted Design: Crossover RCT Setting (Country): Confined (United Kingdom) Study start date; study end date: 06/01/2015; 10/10/2015		
	Interv	ention durati	on: 2 sessions of 10 puffs at 20 sec intervals
Participants	Number of participants: 25 randomised, 1 withdrawn, 24 completed Withdrawal reasons reported: Yes Baseline characteristics: N=25; Mean Age (SD): 33.1 (7.34) years; Sex: 52% male; Ethnicity/Nationality: not reported Key inclusion criteria: Health status: "good general health"; ≥10 CPD; smoked for ≥1 year		
Interventions	Interventions: HTP (HNB2.1), CC (Unknown) Co-interventions: None Mode of exposure: Direct restricted		
Outcomes	Second Second	tration curve flary: Termina	ach nicotine Cmax, Maximal nicotine concentration, Area under the from start of product use to time of last quantifiable concentration al half-life of nicotine, Area under the plasma concentration-time curve
		, Mouth level	use extrapolated from time of last quantifiable concentration to exposure to nicotine, Inhalation to non-inhalation ratios during HTP
Analyses	Prima		tion reported: No opulation: Not specified or unclear lividuals
Study funding		<u>-</u>	national (Industry-affiliated)
Notes	_		
000	Not included in meta-regression analysis.		

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Risk of bias					
Bias		Authors' judgement	Support for judgement		
Random sequence generation		Unclear	Beyond stating the study was 'randomised', no further information provided.		
Allocation conce		Unclear	No information provided.		
Blinding of partic and personnel		High	Study described as "open label". Included non-active comparator (cigarettes).		
Blinding of outco	ome	Low	Study described as "open label". All primary outcomes objectively measured.		
Incomplete outcome data		Low	Attrition: NHTP-CC=0%, CC-NHTP=8%. All 24 subjects who completed the study were included in the analyses.		
Selective reporting	ng	High	2 outcomes listed on the trial registration (mouth level exposure to nicotine and inhalation to non-inhalation ratios) were not reported.		
Nga, 2020					
Submi		tted to peer-r ned key outco	: Not registered reviewed journal within 12 months: Unclear remes on trial registration within 12 months: Unclear		
	_		nised quasi-experimental (Parallel)		
	_	_	Confined (Malaysia)		
	1 -		ady end date: Not reported on: 1 session of 2 10-puff rounds at 30 sec intervals and 5 min inter-		
	round b		on: 1 session of 2 10-puri founds at 50 sec intervals and 5 min inter-		
Participants	comple	Number of participants: 45 enrolled (15 HTP, 15 CC, 15 EC), 0 withdrawn, 45 completed (15 HTP, 15 CC, 15 EC)			
		Vithdrawal reasons reported: N/A			
male;		aseline characteristics: N=45; Mean Age (SD): 43.6 years (SDs not reported); Sex: 87% ale; Ethnicity/Nationality: 51% Chinese, 22% Malay, 20% Indian, 7% Other ey inclusion criteria: Health status: not specified; ≥10 CPD; smoked for ≥5 years			
Interventions	I		(IQOS), CC (Own brand), EC (Aspire AVP)		
	Co-interventions: 1 Mode of exposure:				
Outcomes	Primary: Exhaled Carbon monoxide		arbon monoxide		
		lary: None			
Analyses	Sample size calculat Primary analysis po Unit of analysis: Ind		pulation: Not specified or unclear		
Study funding		-	l University (Independent)		
Notes			-regression analysis.		
Risk of bias	1 2 2 2 110				
Bias		Authors'	Support for judgement		
Random sequence generation	e	High	Non-randomised trial.		
Allocation conce	alment	High	Non-randomised trial.		
Blinding of partic and personnel	cipants	Unclear	No information provided on blinding. Included a non-active comparator (cigarettes).		
Blinding of outco	ome	Low	No information provided on blinding. Primary outcome objectively measured.		
Incomplete outcome			All participants enrolled completed the trial and no participants were		
data	ome	Low	excluded from the analysis.		
-		Low Low			

Participants

Methods	Date o	f registration	: Not registered		
	Submitted to peer-reviewed journal within 12 months: Unclear				
	Published key outcomes on trial registration within 12 months: Unclear				
	Design	: Crossover R	CT		
	Setting	g (Country): (Confined (United States of America)		
	Study	start date; stu	udy end date: Not reported		
	Interv	ention durati	on: 3 sessions of 2 10-puff bouts at 30 sec intervals and 60 min inter-		
	bout b	reak			
Participants	Numb	er of particip	ants: 24 randomised, 9 withdrawn, 15 completed		
	Withd	rawal reasons	s reported: Yes		
	Ethnic Asian,	ity/Nationality 7% unknown	stics: N=15; Mean Age (SD): 33.6 (11.8) years; Sex: 80% male; v: 47% White or Caucasian, 40% Black or African American, 7%		
	Key in	clusion criter	ia: Health status: "healthy"; ≥10 CPD; unspecified smoking duration		
Interventions	Interv	entions: HTP	(PAX), CC (Own brand), EC (eGo)		
	Co-int	erventions: N	Ione		
	Mode	of exposure: 1	Direct restricted		
Outcomes	Primary: Exhaled Carbon monoxide, Questionnaire of Smoking Urges, Nicotine, Minnesota Nicotine Withdrawal Scale, The Direct Effects of Nicotine Questionnaire, The Direct Effects of Product scale				
			öm Test for Nicotine/Cigarette Dependence, Heart rate		
Analyses	_	Sample size calculation reported: No			
	Primary analysis population: Not specified or unclear				
	Unit of analysis: Individuals				
Study funding	National Institute on Drug Abuse of the National Institutes of Health and the Center for Tobacco Products of the U.S. Food and Drug Administration (Independent)				
Notes	Includ	Included in meta-regression analysis. Data obtained from published literature.			
Risk of bias					
Bias		Authors' judgement	Support for judgement		
Random sequence generation	ee	Low	"Participants completed each of the three, Latin-square ordered, ~2.5-h sessions"		
Allocation conce	alment	Unclear	No information provided.		
Blinding of parti	cipants	Unclear	No information provided on blinding. Included a non-active		
and personnel		Oncicui	comparator (cigarettes).		
Blinding of outco	ome	High	No information provided on blinding. Some primary outcomes		
assessment Incomplete outco	ome	_	subjectively measured. Overall attrition = 37.5%. No subjects who completed the study were		
data	лис	Low	excluded from the analysis.		
Selective reporting	ng	Low	All outcomes reported on in at least one literature source.		
ISRCTN810757					
Methods		f registration	: 31/01/2018		
	Date of registration: 31/01/2018 Submitted to peer-reviewed journal within 12 months: Yes				
		_	omes on trial registration within 12 months: No results posted		
		: Parallel RC			
	_				
	Setting (Country): Ambulatory (United Kingdom) Study start date; study end date: 15/02/2018; 31/03/2020				
	Intervention duration: 12-months (day 90 interim analysis)				

Number of participants: 411 enrolled (Glo 105, CC 42, Cess 190, NS 40, THD 34)

Baseline characteristics: N=280 (baseline characteristics for THD arm not reported); Mean Age (SD): Glo 39 (8.8) years, CC 38 (9.3) years, Cess 38 (9.0) years, NS 40 (9.9)

Withdrawal reasons reported: Unclear

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years; Sex: 55% male; Ethnicity/Nationality: 90.7% White, 3.6% Asian, 2.5% Black or African American, 3.2% Other **Key inclusion criteria:** Health status: "good health"; 10-30 CPD; smoked for ≥5 years

Interventions

Interventions: HTP (Glo1.1), CC (Own brand), smoking cessation (aided if necessary), NS, HTP (THD2.4T20)

Co-interventions: None

Mode of exposure: Direct ad libitum

Outcomes

Primary: Augmentation index, 8-epi-prostaglandin F2alpha, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol

Secondary: 2-cyanoethylmercapturic acid, total N-nitrosonornicotine, Nicotine equivalents, monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, total 1-hydroxypyrene, S-phenylmercapturic acid, o-toluidine, 4-aminobiphenyl, 2aminonaphthalene, Daily product consumption, Fagerström Test for Nicotine/Cigarette Dependence, 3-hydroxy-1-methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, Respiratory symptoms (inc. cough assessment), Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Spirometry, 11-dehydrothromboxane B2, White blood cell count, Soluble intercellular adhesion molecule-1, High-density lipoprotein cholesterol, Forced expiratory volume in one second, Weight, Waist circumference, Low-density lipoprotein cholesterol, Homocysteine, Highsensitivity C-reactive protein, Fibrinogen, Forced vital capacity, Forced expiratory flow at 25-75% of forced vital capacity, Triglycerides, Total cholesterol, N-(2-cyanoethyl)valine haemoglobin adducts, Pulse wave velocity, Peak Expiratory Flow, Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Product Satisfaction Questionnaire, 4-Hydroxy-2-nonenal, Blood pressure, Tissue plasminogen activator, Plasminogen activator inhibitor-1, Nitric oxide, Monocyte chemotactic protein 1/C-C motif chemokine ligand 2, Glucose, E-selectin, Endothelin-1, 3-nitrotyrosine, Finger plethysmography, 6-minute walking test, Smoking cessation quality of life questionnaire

Analyses

Sample size calculation reported: Yes

Primary analysis population: Per-protocol population defined as "all subjects who had a valid assessment of a biomarker variable and completed the study (to day 90) according to the protocol. This population excludes subjects in Groups B and D who had major protocol deviations or a significant level of self-reported smoking" and CEVal-compliant population defined as "excludes subjects in Groups B and D who were considered noncompliant with smoking restrictions, based on CEVal levels above predetermined thresholds"

Unit of analysis: Individuals

Study funding

British American Tobacco (Industry-affiliated)

Notes

The published data was from an interim analysis at day 90. Data for the full 12-months has not yet been published. The number of participants randomised/withdrawn/completed at Day 90 was only reported for one arm (THD2.4T20) in which all 34 randomised participants were excluded from the study without explanation. Included in meta-regression analysis. Data obtained from study authors.

Risk of bias		
Bias	Authors' judgement	Support for judgement
Random sequence generation	Low	"randomised using blocks of computer-generated random number sequences"
Allocation concealment	Unclear	No information provided.
Blinding of participants and personnel	High	"This study will not be blinded". Included non-active comparator (cigarettes).
Blinding of outcome assessment	Low	"This study will not be blinded". All primary outcomes objectively measured.
Incomplete outcome data	Unclear	Number of subjects randomised, attrition and exclusions were not reported, neither were reasons for exclusion.
Selective reporting	High	The 90-day interim publication is the only reporting of results from this 12-month trial. In this publication, only a small selection of outcomes listed in the trial registration and protocol are reported, including only 1 primary outcome.

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ISRCTN13439529				
Methods	Date o	f registration	: 07/08/2018	
	Submi	itted to peer-r	reviewed journal within 12 months: No publication	
	Published key outcomes on trial registration within 12 months: No results posted			
	Design: Crossover RCT			
	Setting (Country): Confined (Italy)			
	Study	start date; stu	udy end date: 01/01/2018; 30/09/2018	
	Interv	ention durati	on: 4 sessions of single use of one cigarette, tobacco stick or cartridge	
Participants			ants: 32 randomised, withdrawn/completed not reported s reported: N/A	
		ine characteristics: N= 32; Mean Age (SD): 35.8 (9.66) years; Sex: 72% male;		
		ity/Nationality		
			ria: Health status: normal biochemistry, haematology, urinalysis, ECG PD; smoked for ≥ 1 year	
Interventions	Interv	entions: HTP	(Glo1.0), HTP (Glo1.1), CC (Own brand), NRT (Nicorette inhaler)	
		erventions: N		
	Mode	of exposure: 1	Direct ad libitum	
Outcomes	Primary: Time to reach nicotine Cmax, Maximal nicotine concentration, Area under the concentration curve from start of product use to time of last quantifiable concentration, Intention to use [HTP] Questionnaire, Product Liking Questionnaire, Urge To Smoke questionnaire, Urge For Product questionnaire Secondary: Product Evaluation Scale, Human Puffing/Smoking Topography (inc. puff count), Adverse events			
Analyses	Sample size calculation reported: Yes			
rinary ses	1 -		opulation: Not specified or unclear	
		f analysis: Inc	-	
Study funding		-	bacco (Industry-affiliated)	
Notes	Not inc	cluded in meta	-regression analysis	
Risk of bias	ı			
Bias		Authors' judgement	Support for judgement	
Random sequence	ce	Low	"The order of use will be assigned by a pre-defined computer-	
generation	almant	Unclear	generated randomisation schedule" No information provided.	
Allocation concealment Blinding of participants			•	
and personnel		High	"open-label". Included non-active comparator.	
Blinding of outcome assessment		High	"open-label". Some primary outcomes subjectively measured.	
Incomplete outco	Incomplete outcome		While the number of participants randomised is reported, the number withdrawn/completed and included in the analysis was not reported.	
Selective reporting		High	The two secondary outcomes (Puff count during 5 minute investigational product use session and Product evaluation using the Product Evaluation Scale (PES)) were not reported.	

ISRCTN14301360/UMIN000024988

Methods

Date of registration: 14/12/2016 (ISRCTN), 24/11/2016 (UMIN) **Submitted to peer-reviewed journal within 12 months:** Yes

Published key outcomes on trial registration within 12 months: No results posted

Design: Parallel RCT

Setting (Country): Confined (Japan)

Study start date; study end date: 01/08/2016; 30/06/2017

Intervention duration: 5 days

Participants	Number of participants: 182 (30 Glo R, 30 Glo M, 30 CC R, 30 CC M, 30 Cess, 30 IQOS R, 2 unknown), 2 withdrawn (2 unknown), 180 completed (30 Glo R, 30 Glo M, 30 CC R, 30 CC M, 30 Cess, 30 IQOS R) Withdrawal reasons reported: Yes
	Baseline characteristics: N= 180; Mean Age (SD): Glo R 34 (10.1) years, Glo M 31 (7.7) years, CC R 32 (8.2) years, CC M 33 (8.6) years, Cess 35 (10.0) years, IQOS R 33 (9.5) years; Sex: 50% male; Ethnicity/Nationality: 100% Japanese Key inclusion criteria: Health status: "good health"; 10-30CPD; smoked for ≥3 years
Interventions	Interventions: HTP (Glo 1.0 R), HTP (Glo 1.0 M), HTP (IQOS R), CC (Lucky Strike R), CC (Lucky Strike M), tobacco and nicotine cessation Co-interventions: None
	Mode of exposure: Direct ad libitum
Outcomes	Primary: Exhaled Carbon monoxide, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, total N-nitrosonornicotine, Nicotine equivalents, monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, total 1-hydroxypyrene, S-phenylmercapturic acid, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene, 3-hydroxy-1-methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, N-acetyl-S-(2-hydroxy-2-carbamoylethyl)cysteine, N-acetyl-S-(2-carbamoylethyl)cysteine Secondary: Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Electrocardiogram, Time to reach nicotine Cmax, Maximal nicotine concentration, Area under the concentration curve from start of product use to time of last quantifiable concentration, Spirometry, 8-epi-prostaglandin F2alpha, Human Puffing/Smoking Topography (inc. puff count), White blood cell count, Nicotine molar metabolic ratio, Product Satisfaction Questionnaire, Medical history, Adverse Events/Serious Adverse Events, Daily product consumption, Vital signs
Analyses	Sample size calculation reported: Yes
	Primary analysis population: Per protocol population defined as "All subjects who had valid assessment of a biomarker variable and completed study according to the protocol will be used for biomarker analyses" and pharmacokinetic population defined as "All subjects who had sufficient data to calculate at least 1 pharmacokinetic parameter and completed study according to the protocol will be used for PK data analyses". Unit of analysis: Individuals
Study funding	British American Tobacco (Industry-affiliated)
Notes	2 participants were randomised but withdrew before the exposure period. The groups these 2 belonged to were not reported. Included in meta-regression analysis. Data obtained from published literature and study authors.
Risk of bias	
Rias	Authors' Support for judgement

Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation	Low	"The randomisation will be performed by Covance"	
Allocation concealment	Low	"The randomisation will be performed by Covance and the clinics will enrol the participants and assign them to interventions"	
Blinding of participants and personnel	High	"open-label". Included non-active comparator (cigarettes).	
Blinding of outcome assessment	Low	"open-label". All primary outcomes objectively measured.	
Incomplete outcome data	Low	Overall attrition = 1.1%. No subjects who completed the study were excluded from the primary analyses.	
Selective reporting	High	There were several outcomes listed in the protocol, namely biomarkers of effect and pharmacokinetic measures, that were not reported on.	
DDT7C00044040			

DRKS00012919

Methods Date of registration: 29/08/2017

	1	_	reviewed journal within 12 months: Unclear		
	1	-	omes on trial registration within 12 months: Unclear		
	-	: Crossover R			
	1	-	Confined (Germany)		
	1		udy end date: 01/06/2016; not reported		
		ention duration of the second	on: 4 sessions of single use of one cigarette or tobacco stick at 1 puff puffs		
Participants	Numbe	er of particip	ants: 20 randomised, 0 withdrawn, 20 completed		
	Withd	rawal reason	s reported: N/A		
			stics: N= 20; Mean Age (SD): 21.9 (2.6) years; Sex: 50% male; root reported		
	Key in not rep		ria: Health status: no disorders or diseases; CPD and smoking duration		
Interventions	Interve		(IQOS2.2), CC (Marlboro Gold), EC (eGo nicotine), EC (eGo no		
	Co-int	erventions: N	Ione		
	Mode	of exposure:	Direct ad libitum		
Outcomes	Prima	ry: Nicotine, S	Systolic blood pressure		
	Second	Secondary: Heart rate, Pulse wave velocity, Augmentation index, [Mean] Arterial Blood Pressure			
Analyses	Sampl	e size calcula	tion reported: No		
	Prima	ry analysis po	opulation: Not specified or unclear		
	Unit of	f analysis: Inc	lividuals		
Study funding	Univer	sitätsklinikum	Schleswig-Holstein Campus Lübeck (Independent)		
Notes			i-regression analysis.		
Risk of bias	'				
Bias		Authors' judgement	Support for judgement		
Random sequence generation	ce	Unclear	Beyond stating the study was 'randomised', no further information provided.		
Allocation conce		Unclear	No information provided.		
Blinding of partial and personnel	cipants	High	Only the e-cigarette arms were blinded. Included non-active comparator (cigarettes).		
Blinding of outco	ome		Only the e-cigarette arms were blinded. All primary outcomes		
assessment		Low	objectively measured.		
Incomplete outco data	ome	Unclear	In the protocol the target for enrolment was 55. It is unclear whether 55 completed the study and only 20 were included in the analyses.		
Selective reporting	ng	High	In the trial registration, the authors state outcomes relating to "endothelial dysfunction and inflammatory markers" were measured. No specific measures were given and no relevant data were reported.		
ISRCTN806519	000		No specific measures were given and no relevant data were reported		
	T .	C	. 00/02/2017		
Methods	1	f registration			
	Submitted to peer-reviewed journal within 12 months: No				
		=	omes on trial registration within 12 months: No results posted		
	_	Design: Parallel RCT			
	_		Confined (United Kingdom)		
		•	udy end date: 01/08/2016; 03/10/2017		
		ention durati	•		
Participants	withdra	awn (2 Glo, 2	ants: 148 randomised (30 Glo, 30 CC, 30 EC, 29 Cess, 29 HTP), 7 EC, 2 Cess, 1 HTP), 143 (28 Glo, 30 CC, 28 EC, 29 Cess, 28 HTP)		
	vvitnd	rawai reason	s reported: Yes		

Baseline characteristics: N= 148; Mean Age (SD): Glo 37.4 (11.48) years, CC 35.6 (8.93) years, EC 36.7 (9.1) years, Cess 37.2 (9.09) years, HTP (32.8 (8.78) years; Sex: 59% male; Ethnicity/Nationality: 100% White **Key inclusion criteria:** Health status: "good health"; 10-30CPD; smoked for \geq 3 years

Interventions

Interventions: HTP (Glo1.0), CC (Lucky Strike Regular), EC (prototype IS1.0[TT]), tobacco and nicotine cessation, HTP (unknown)

Co-interventions: None

Mode of exposure: Direct ad libitum

Outcomes

Primary: Exhaled Carbon monoxide, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, total N-nitrosonornicotine, Nicotine equivalents, monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, total 1-hydroxypyrene, S-phenylmercapturic acid, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene, N-acetyl-S-(2-hydroxy-2-carbamoylethyl)cysteine, N-acetyl-S-(2-carba-moylethyl)cysteine, 3-hydroxy-1-methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, 8-epi-prostaglandin F2alpha, White blood cell count, Nicotine molar metabolic ratio

Secondary: Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Time to reach nicotine Cmax, Maximal nicotine concentration, Area under the concentration curve from start of product use to time of last quantifiable concentration, Spirometry, Product Satisfaction Questionnaire, Adverse Events/Serious Adverse Events, Daily product consumption

Analyses

Sample size calculation reported: Yes

Primary analysis population: Not specified or unclear

Unit of analysis: Individuals

Study funding

Notes

British American Tobacco (Industry-affiliated)

According to the published study literature, 29 participants were randomised to the cessation and 29 completed this study, yet 2 were said to have withdrawn. It is not clear if these 2 were replaced or if this was a mistake. Data from the unknown HTP arm was excluded from the analysis because the authors "wished to focus on the exposure continuum". Included in meta-regression analysis. Data obtained from published literature and study authors.

Risk of bias			
Bias	Authors' judgement	Support for judgement	
Random sequence generation	Low	"The randomization will be computer-generated using SAS Version 9.3"	
Allocation concealment	Low	"A randomisation scheme was provided for the clinical site to recruit 30 participants for each arm, giving a total of 150 participants"	
Blinding of participants and personnel	High	"open-label". Included non-active comparator (cigarettes).	
Blinding of outcome assessment	Low	"open-label". All primary outcomes objectively measured.	
Incomplete outcome data	Low	Attrition: Glo=6.67% CC=0% EC=6.67% Cess=0% HTP=3.45%, overall=3.38%. Exclusion: Glo=6.67% CC=0% EC=6.67% Cess=0% HTP=N/A, overall=3.34%.	
Selective reporting	High	No data reported for an entire study arm (C: "switching to a non-BAT commercial product"). No quantitative data reported for two biomarker of effect outcomes (WBC count & 8-epi-PGF2α Type III). No data reported for pharmacokinetic outcomes measured	

UMIN000041539

Methods

Date of registration: 25/08/2020

Submitted to peer-reviewed journal within 12 months: No publication

Published key outcomes on trial registration within 12 months: No results posted

Design: Parallel RCT

Setting (Country): Confined (Japan)

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	Ctudu	stant datas str	udy and data. Santanahan 2020, Oatahan 2020		
	Study start date; study end date: September 2020; October 2020 Intervention duration: 5 days				
HTP, 1		Tumber of participants: 90 randomised (15 Ploom Tech+, 15 Ploom S2.0, 15 unknown TP, 15 unknown HTP, 15 CC, 15 Cess), withdrawn/completed not reported			
			s reported: N/A		
			stics: not reported ia: Health status: "good health"; unspecified CPD; smoked for ≥1		
	year	iciusion criter	ia. Heathi status. good heathi , unspecified ci D, sinoked foi \(\frac{1}{2}\)1		
Interventions	(unkno	own), CC (Ow	(Ploom Tech+), HTP (Ploom S2.0), HTP (unknown), HTP n brand), smoking cessation		
		Co-interventions: None Mode of exposure: Unclear			
Outcomes	Primary: Exhaled Carbon monoxide, Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, total N-nitrosonornicotine, monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, total 1-hydroxypyrene, S-phenylmercapturic acid, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene, 3-hydroxy-1-methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, 3-hydroxybenzo[a]pyrene, 1-aminonaphthalene Secondary: None				
Analyses		•	tion reported: No		
-	Prima	ry analysis po	opulation: Not specified or unclear		
	Unit of	f analysis: Inc	lividuals		
Study funding	Japan Tobacco International (Industry-affiliated)				
Notes	Data requested from study authors, but no data received. Therefore, not included in meta-regression analysis.				
Risk of bias		ı			
Bias		Authors' judgement	Support for judgement		
Random sequence generation	ee	Unclear	Beyond stating the study was 'randomised', no further information provided.		
Allocation conce		Unclear	No information provided.		
Blinding of partical and personnel	cipants	High	"Open -no one is blinded". Included non-active comparator (cigarettes).		
Blinding of outco	ome	Low	"Open -no one is blinded". All primary outcomes objectively measured.		
Incomplete outcome data		Unclear	While the number of participants randomised was reported, the number completed and included in the analysis was not.		
Selective reporting		Unclear	As the trial registration does not explicitly list all outcomes measured in this trial and there is no publicly available protocol, it is difficult to determine whether the 15 biomarkers of exposure were the only measures of the study. Moreover, data is thus far only presented in a graph.		
NCT03700112					
Methods	Date o	f registration	: 09/10/2018		
		_	reviewed journal within 12 months: No publication		
	Published key outcomes on trial registration within 12 months: No results pos Design: Crossover RCT				
		-	Confined (New Zealand)		
	Study start date; study end date: 04/12/2018; 09/04/2019 Intervention duration: 8 sessions of 10 puffs at 30 second intervals and 8 sessions of <i>ad lib</i> use for 4.5 minutes				
Participants	Number of participants: 25 randomised, 0 withdrawn, 25 completed Withdrawal reasons reported: N/A				
			•		

	Baseline characteristics: N= 25; Mean Age (SD): 30.44 (10.18) years; Sex: 72% male; Ethnicity/Nationality: not reported		
	Key inclusion criteria: Health status: "healthy"; ≥8 CPD; smoked for ≥1 year		
Interventions	Interventions: EC (JUUL), EC (myblu Original 2.4%), EC (MarkTen Bold Classic 4.0%),		
Title veneralis	EC (VUSE Solo Original 4.8%), EC (PHIX Original Tobacco 5.0%), EC (NJOY Daily EXTRA Rich Tobacco 6.0%), HTP (IQOS), CC (Marlboro Red)		
	Co-interventions: None		
		•	Direct restricted and ad libitum
Outcomes	Primary: Time to reach nicotine Cmax, Maximal nicotine concentration, Baseline adjusted Cmax, Baseline adjusted AUC1hour, Area under the concentration curve from start of product use to 60 minutes		
	Questi		Carbon monoxide, Modified Cigarette/Product Evaluation an Puffing/Smoking Topography (inc. puff count), Rate of plasma of absorption)
Analyses			tion reported: No
rinary ses	_		opulation: Not specified or unclear
		f analysis: Inc	-
Study funding		•	ustry-affiliated)
Notes			-regression analysis.
Risk of bias	1.00 1110	iii iiicla	. reg. evoluti utuu julu.
Bias		Authors'	Support for judgement
Dias		judgement	Support for juagement
Random sequence generation	e	Unclear	Beyond stating the study was 'randomised', no further information provided.
Allocation conce		Unclear	No information provided.
Blinding of partic	cipants	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).
	nding of outcome		"Masking: None (Open Label)". All primary outcomes objectively measured.
Incomplete outcome data		Unclear	Attrition was 0%. Exclusion=0-8% as the analysis population stated under the tables on poster was "N=24-25" or "N=23-25". However, the exact N for each outcome analysis is not specified and reasons for excluding some subjects from the analyses are not provided.
Selective reporting	ng	High	Total number of puffs during exposure session and exhaled CO - both measures listed on the trial registration - were not reported.
NCT01970995			
Methods	Date o	f registration	: 28/10/2013
			reviewed journal within 12 months: No
		-	omes on trial registration within 12 months: No
		: Parallel RC	_
	Setting	g (Country): (Confined and Ambulatory (Japan)
		-	udy end date: 01/08/2013; November 2014
	Interv	ention durati	on: 90 Days (5 days confinement + 85 days ambulatory)
Participants			ants: 160 randomised (78 HTP, 42 CC, 40 Cess), 5 withdrawn (2 155 (76 HTP, 41 CC, 38 Cess)
			s reported: Yes
	Baseline characteristics: N= 160; Mean Age (SD): HTP 37.1 (10.58) years, CC 37.4 (11.23) years, Cess 37 (9.96) years; Sex: 57.5% male; Ethnicity/Nationality: 100% Japanese		
			ia: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years
Interventions	Interventions: HTP (IQOS2.2 M), CC (Own brand M), smoking cessation (aided if necessary)		
	Co-interventions: None		

Mode of exposure: Direct restricted and ad libitum

Outcomes

Primary: Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, Carboxyhemoglobin, S-phenylmercapturic acid

Secondary: Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, 2cyanoethylmercapturic acid, Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, total N-nitrosonornicotine, Nicotine equivalents, total 1hydroxypyrene, o-toluidine, 4-aminobiphenyl, 2-aminonaphthalene, Daily product consumption, Fagerström Test for Nicotine/Cigarette Dependence, 3-hydroxy-1methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, Respiratory symptoms (inc. cough assessment), Nicotine, Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, 3-hydroxybenzo[a]pyrene, Spirometry, Concomitant medications, Cotinine, 1-aminonaphthalene, 8-epi-prostaglandin F2alpha, 11-dehydrothromboxane B2, Minnesota Nicotine Withdrawal Scale, Cytochrome P450 2A6 activity, Human Puffing/Smoking Topography (inc. puff count), Ames mutagenicity test (YG1024+S9), White blood cell count, Systolic blood pressure, Soluble intercellular adhesion molecule-1. High-density lipoprotein cholesterol, Diastolic blood pressure, Time to nicotine Cpeak, Maximum observed nicotine concentration (following ad lib use), S-benzylmercepturic acid, Weight, Waist circumference, Low-density lipoprotein cholesterol, Homocysteine, High-sensitivity C-reactive protein, Fibrinogen, Platelet count, Hemoglobin glycosylated (Hemoglobin A1C), Potential combustion occurances in tobacco plugs, Weighted average nicotine concentration over 24 hours, Human Puffing/Smoking Topography Questionnaire, Triglycerides

Total cholesterol, Blood glucose

Analyses

Sample size calculation reported: Yes

Primary analysis population: Per-protocol population defined as "all randomized subjects who - have had compliance to their randomized arm if randomized to THS 2.2 Menthol or SA arms. Non-compliance will be defined over a period (confinement period, J Day6-Day 30 Visit], JDay 30 Visit-Day 60 Visit], JDay 60 Visit-Day 90 Visit] and will be defined as having smoked than 3 CC during a single day in that period or having smoked on average over that period more that, not including 0.5 cigarettes per day. - have not been misrandomized. - and have no major protocol deviation"

Unit of analysis: Individuals

Study funding Notes

Philip Morris International (Industry-affiliated)

Included in meta-regression analysis. Data obtained from published literature.

Risk of bias

RISK OF DIAS			
Bias	Authors' judgement	Support for judgement	
Random sequence generation	Low	"randomization was performed through the Interactive Web and Voice Response System"	
Allocation concealment	Low	"randomization was performed through the Interactive Web and Voice Response System"	
Blinding of participants and personnel	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).	
Blinding of outcome assessment	Low	"Masking: None (Open Label)". All primary outcomes objectively measured.	
Incomplete outcome data	Low	Attrition: IQOS=2.56% CC=2.38% Cess=5%, overall=3.12%. Exclusion: IQOS=10.26% CC=2.4% Cess=7.5%, overall=7.5%.	
Selective reporting	Low	All outcomes reported in at least one literature source.	

NCT01989156

Methods

Date of registration: 20/11/2013

Submitted to peer-reviewed journal within 12 months: No Published key outcomes on trial registration within 12 months: No

Design: Parallel RCT

Setting (Country): Confined and Ambulatory (United States of America)

Study start date; study end date: 17/12/2013; May 2015

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			on: 91 Days (5 days confinement + 86 days ambulatory)		
Participants		nber of participants: 160 (80 HTP, 41 CC, 39 Cess), 21 withdrawn (7 HTP, 6 CC, 8 s), 139 completed (73 HTP, 35 CC, 31 Cess)			
		-	s reported: Yes		
			stics: N= 160; Mean Age (SD): HTP 39.2 (11.72) years, CC 33.7		
			8.8 (11.42) years; Sex: 60% male; Ethnicity/Nationality: 62% White,		
			n American, 6% other, 1% missing		
	Key in	nclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years			
Interventions	Interv	entions: HTP	(IQOS2.2 M), CC (Own brand M), smoking cessation (aided if		
	necessary)				
		erventions: N			
	Mode	of exposure: 1	Direct ad libitum		
Outcomes			nethylnitrosamino)-1-(3-pyridyl)-1-butanol,		
			Imercapturic acid, 3-hydroxypropylmercapturic acid,		
			S-phenylmercapturic acid Carbon monoxide, Adverse Events/Serious Adverse Events, 2-		
			ric acid, Modified Cigarette/Product Evaluation Questionnaire,		
		•	oking Urges, total N-nitrosonornicotine, Nicotine equivalents, total 1-		
			luidine, 4-aminobiphenyl, 2-aminonaphthalene, Daily product		
			tröm Test for Nicotine/Cigarette Dependence, 3-hydroxy-1-		
			turic acid, 2-hydroxyethylmercapturic acid, Respiratory symptoms		
		(inc. cough assessment), Nicotine, Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, 3-hydroxybenzo[a]pyrene, Concomitant medications, Cotinine, 1-aminonaphthalene, 8-epi-prostaglandin F2alpha, 11-dehydrothromboxane B2, Minnesota Nicotine Withdrawal Scale, Cytochrome P450 2A6 activity, Human Puffing/Smoking Topography (inc. puff count), Ames mutagenicity test (YG1024+S9), White blood cell count, Systolic blood pressure, Soluble intercellular adhesion molecule-1, High-density lipoprotein cholesterol, Forced expiratory volume in one second, Diastolic blood pressure, Time to nicotine Cpeak, Maximum observed nicotine			
	one sec				
			ving ad lib use), S-benzylmercepturic acid, Weight, Waist		
			density lipoprotein cholesterol, Homocysteine, High-sensitivity C-		
		e protein, Fibrinogen, Platelet count, Hemoglobin glycosylated (Hemoglobin A1C), al combustion occurances in tobacco plugs, Weighted average nicotine			
concer vital c Total o		entration over 24 hours, Human Puffing/Smoking Topography Questionnaire, Forced			
		apacity, Forced expiratory flow at 25–75% of forced vital capacity, Triglycerides,			
		cholesterol, Apolipoprotein B, Apolipoprotein A1, Total lung capacity, Blood			
		ose, Residual volume, Vital capacity, Inspiratory capacity, Diffusion Capacity, Carbon oxide transfer coefficient, Oxysterols (6α-hydroxy-5α-cholestanol, 7α-			
		roxycholesterol, 5α,6αepoxycholestanol, 7-ketocholesterol, 7β-hydroxycholesterol,			
		3,6β-epoxycholestanol, 24(R)-hydroxycholesterol, 25-hydroxycholesterol, 22(R)-			
		ydroxycholesterol, 4ßhydroxycholesterol, and 27-hydroxycholesterol), Prochaska "Stage f Change" Questionnaire			
Analyses	_		tion reported: Yes		
			pulation: Per-protocol population defined as "all randomized had compliance to their randomized arm; Have not been		
			Have no major protocol deviation"		
		it of analysis: Individuals			
Study funding	Philip	Morris Interna	itional (Industry-affiliated)		
Notes	Include	ed in meta-reg	ression analysis. Data obtained from published literature.		
Risk of bias					
Bias		Authors'	Support for judgement		
D 1		judgement			
Random sequence	ee	Low	"randomization was done through the Interactive Web and Voice		
generation		-	Response System (IWRS)" "randomization was done through the Interactive Web and Voice		
Allocation concealment		Low	Response System (IWRS)"		

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Blinding of partical and personnel	cipants	High	"This is an open-label study". Included non-active comparator (cigarettes).	
Blinding of outcome assessment		Low	"This is an open-label study". All primary outcomes objectively measured.	
			Attrition: IQOS=9%, CC=15%, SA=21%. Although the primary	
Incomplete outco	ome	Low	analysis used per-protocol populations, results data for the full	
		T	analysis set were also provided in the clinical study report.	
Selective reporting	ng	Low	All outcomes reported in at least one literature source.	
NCT01970982	I			
Methods		registration		
		_	eviewed journal within 12 months: No	
		-	omes on trial registration within 12 months: No	
		Parallel RCT	Confined (Japan)	
	_	-	idy end date: 23/07/2013; July 2014	
	1	ention duration		
Participants			ants: 160 randomised (80 HTP, 40 CC, 40 Cess), 2 withdrawn (2	
rarticipants	Cess), 1	58 completed	1 (80 HTP, 40 CC, 38 Cess)	
			s reported: Yes	
			stics: N= 160; Mean Age (SD): HTP 37.6 (11.7) years, CC 37.2 (11.7) 6) years; Sex: 50% male; Ethnicity/Nationality: 100% Japanese	
	-		ia: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years	
Interventions	-		· · · · · · · · · · · · · · · · · · ·	
interventions		erventions: N	(IQOS2.2), CC (Own brand), tobacco and nicotine cessation	
	1	Iode of exposure: Direct ad libitum		
Outcomes	Primary: monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid,			
Outcomes			S-phenylmercapturic acid	
	1	Secondary: Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Total 4-		
		methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, Modified		
			aluation Questionnaire, Questionnaire of Smoking Urges, total N-	
			ficotine equivalents, total 1-hydroxypyrene, o-toluidine, 4- ninonaphthalene, Daily product consumption, Fagerström Test for	
			ependence, 3-hydroxy-1-methylpropylmercapturic acid, 2-	
	hydroxy	yethylmercapt	turic acid, Respiratory symptoms (inc. cough assessment), Nicotine,	
			n, Clinical chemistry, haematology and urine analysis safety panel,	
			urdiogram, 3-hydroxybenzo[a]pyrene, Spirometry, Concomitant e, 1-aminonaphthalene, 8-epi-prostaglandin F2alpha, 11-	
			e B2, Minnesota Nicotine Withdrawal Scale, Cytochrome P450 2A6	
	activity.	, Human Puff	ing/Smoking Topography (inc. puff count), Ames mutagenicity test	
		(YG1024+S9), Time to nicotine Cpeak, Maximum observed nicotine concentration		
), S-benzylmercepturic acid, Potential combustion occurrences in nted average nicotine concentration over 24 hours, Human	
			pography Questionnaire	
Analyses	_		tion reported: Yes	
J 	_		pulation: Full analysis set defined as "all the randomized subjects	
	who had	d at least one	post-randomization product use experience, if randomized to THS 2.2	
			east one valid nonsafety assessment"	
		analysis: Ind		
Study funding	_		tional (Industry-affiliated)	
Notes		-	study authors, but no data received. Therefore, not included in meta-	
D2.1 01.*	regressi	on analysis.		
Risk of bias		A 114h c ?	Summent for independent	
Bias		Authors'	Support for judgement	
Dias	judgement Support for judgement			

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Random sequence generation	e	Low	"randomization was performed through an Interactive Web and Voice Response System"
Allocation concealment		Low	"randomization was performed through an Interactive Web and Voice Response System"
Blinding of participants and personnel		High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).
Blinding of outco	ome	Low	"Masking: None (Open Label)". All primary outcomes objectively measured.
Incomplete outco		Low	Attrition: IQOS=0% CC=0% Cess=5%, overall=1.25%. All subjects who completed the study were included in the analysis.
Selective reporting	ng	Low	All outcomes reported in at least one literature source.
NCT01959932 Methods	Doto	f vocistuation	. 10/10/2012
Wethous	Date of registration: 10/10/2013 Submitted to peer-reviewed journal within 12 months: No Published key outcomes on trial registration within 12 months: No Design: Parallel RCT Setting (Country): Confined (Poland) Study start date; study end date: 29/06/2013; June 2014		
	1	ention durati	
Participants	HTP),	158 completed	ants: 160 randomised (80 HTP, 41 CC, 39 Cess), 2 withdrawn (1 d (79 HTP, 41 CC, 39 Cess)
years, Cess 33.6 (11.		ne characteris Cess 33.6 (11.	stics: N= 160; Mean Age (SD): HTP 35.4 (9.4) years CC 32.6 (10.06) 51) years; Sex: 50% male; Ethnicity/Nationality: 100% White ia: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years
Interventions	Interventions: HTP (IQOS2.2), CC (Own brand), tobacco and nicotine cessation Co-interventions: None Mode of exposure: Direct ad libitum		
Outcomes	Carbox Second Total 4 Modifit total N aminot methyl (inc. cc and uri Spirom F2alph P450 2 mutage concern occura	Primary: monohydroxybutenylmercapturic acid, 3-hydroxypropylmercapturic acid, Carboxyhemoglobin, S-phenylmercapturic acid Secondary: Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol, 2-cyanoethylmercapturic acid, Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, total N-nitrosonornicotine, Nicotine equivalents, total 1-hydroxypyrene, o-toluidine, 4- aminobiphenyl, 2-aminonaphthalene, Daily product consumption, 3-hydroxy-1- methylpropylmercapturic acid, 2-hydroxyethylmercapturic acid, Respiratory symptoms (inc. cough assessment), Nicotine, Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, 3-hydroxybenzo[a]pyrene, Spirometry, Concomitant medications, Cotinine, 1-aminonaphthalene, 8-epi-prostaglandin F2alpha, 11-dehydrothromboxane B2, Minnesota Nicotine withdrawal Scale, Cytochrome P450 2A6 activity, Human Puffing/Smoking Topography (inc. puff count), Ames mutagenicity test (YG1024+S9), Time to nicotine Cpeak, Maximum observed nicotine concentration (following ad lib use), S-benzylmercepturic acid, Potential combustion occurances in tobacco plugs, Weighted average nicotine concentration over 24 hours, Human Puffing/Smoking Topography Questionnaire	
Analyses	Sample size calculation reported: Yes Primary analysis population: Full analysis set defined as "all randomized participants who used the allocated product at least once after randomization and with at least one valid value for a biomarker of exposure" Unit of analysis: Individuals		opulation: Full analysis set defined as "all randomized participants ed product at least once after randomization and with at least one marker of exposure"
Study funding	Philip Morris International (Industry-affiliated)		
Notes	Data requested from study authors, but no data received. Therefore, not included in meta-		

regression analysis.

Risk of bias

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Bias		Authors' judgement	Support for judgement	
Random sequence generation		Low	"randomization was done through an Interactive Web and Voice Response System"	
Allocation concealment		Low	"randomization was done through an Interactive Web and Voice Response System"	
Blinding of partic	cipants	High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).	
	Blinding of outcome		"Masking: None (Open Label)". All primary outcomes objectively measured.	
Incomplete outco	Incomplete outcome		Attrition: IQOS=1.25% CC=0% Cess=0%, overall=0.62%. All subjects who completed the study were included in the analysis.	
Selective reportir	ng	Low	All outcomes reported in at least one literature source.	
NCT01959607				
Methods	Date of registration: 10/10/2013 Submitted to peer-reviewed journal within 12 months: No Published key outcomes on trial registration within 12 months: No Design: Crossover RCT Setting (Country): Confined (Japan) Study start date; study end date: 31/07/2013; April 2014 Intervention duration: 2 sessions of 14 puffs (6 minutes)			
Participants	Number of participants: 62 randomised (44 HTP/CC, 18 HTP/NRT), 2 withdrawn (2 HTP/CC), 60 completed (42 HTP/CC, 18 HTP/NRT) Withdrawal reasons reported: Yes Baseline characteristics: N= 60; Mean Age (SD): HTP/CC 33.2 (8.61) years, HTP/NRT 35.8 (10.44) years; Sex: 55% male; Ethnicity/Nationality: 100% Japanese Key inclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years			
Interventions	Interventions: HTP (IQOS2.2), CC (Own brand), NRT (Nicorette gum) Co-interventions: None Mode of exposure: Direct restricted			
Outcomes Prim		mary: Maximal nicotine concentration, Area under the concentration curve from start roduct use to time of last quantifiable concentration		
	Secondary: Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, Carboxyhemoglobin, Respiratory symptoms (inc. cough assessment), Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Time to reach nicotine Cmax, Spirometry, Concomitant medications, Terminal half-life of nicotine, Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity, Partial AUC			
Analyses	Sample size calculation reported: Yes			
	Primary analysis population: Pharmacokinetic population defined as "all randomised subjects who gave informed consent, completed at least 1 of the single-use days (Day 1 or 3), and for whom at least 1 PK parameter was derived. Subjects with major protocol deviations that impacted the evaluability of the results were excluded from the PK analysis sets." Unit of analysis: Individuals			
Study funding	Philip Morris International (Industry-affiliated)			
Notes				
Risk of bias				
Bias		Authors' judgement	Support for judgement	
Random sequenc generation	Random sequence Low		"Randomization to each product exposure sequence was done through an Interactive Telephone and Web Response System."	

Allocation conce	alment	Low	"Randomization to each product exposure sequence was done			
Blinding of participants		High	through an Interactive Telephone and Web Response System." "This was an open-label study". Included non-active comparator			
and personnel		nigii	(cigarettes).			
Blinding of outcome assessment		Low	"This was an open-label study". All primary outcomes objectively measured.			
Incomplete outcome data		Low	Attrition: IQOS-CC=5%, IQOS-NRT=0%. No participants who completed the trial were excluded from the analyses.			
Selective reporting	ng	Low	All outcomes reported in at least one literature source.			
NCT01967732						
Methods	Date o	Date of registration: 23/10/2013				
	Submitted to peer-reviewed journal within 12 months: No publication					
		_	omes on trial registration within 12 months: No			
			n: Crossover RCT			
_		g (Country): Confined (United Kingdom)				
	Study	start date; stı	udy end date: 01/11/2013; July 2014			
	Interv	ention durati	on: 2 sessions of single use of one cigarette, tobacco stick or 1 nasal			
	spray i	n each nostril				
		ber of participants: 62 randomised (44 HTP/CC, 18 HTP/NRT), 2 withdrawn (2 CC), 60 completed (42 HTP/CC, 18 HTP/NRT)				
	Withd	ithdrawal reasons reported: Yes				
	1	aseline characteristics: N= 60; Mean Age (SD): HTP/CC 32.1 (8.98) years, HTP/NRT 0.6 (5.8) years; Sex: 58% male; Ethnicity/Nationality: 100% Japanese				
	Key in	clusion criter	ia: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years			
Interventions	Interv	entions: HTP	(IQOS2.2), CC (Own brand), NRT (Nicotrol nasal spray)			
Co-interventions: None			Tone			
	Mode	of exposure: 1	Direct ad libitum			
Outcomes	Primary: Maximal nicotine concentration, Area under the concentration curve from start of product use to time of last quantifiable concentration					
	Secondary: Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, Carboxyhemoglobin, Fagerström Test for Nicotine/Cigarette Dependence, Respiratory symptoms (inc. cough assessment), Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Time to reach nicotine Cmax, Spirometry, Concomitant medications, Terminal half-life of nicotine, Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity, Partial AUC					
Analyses	Sampl	e size calculat	tion reported: Yes			
	Primary analysis population: Pharmacokinetic population defined as "all the randomized subjects who give informed consent, completed at least one of the single use Day 1 or Day 3, and for whom at least one PK parameter can been derived. Only subjects without major protocol deviations" Unit of analysis: Individuals					
Study funding	Philip	Morris Interna	ational (Industry-affiliated)			
Notes	_		ression analysis. Data obtained from published literature.			
Risk of bias						
Bias		Authors'	Support for judgement			
		judgement				
Random sequence generation		Low	"Randomization to product exposure sequence was performed through an Interactive Telephone and Web Response System"			
Allocation concealment		Low	"Randomization to product exposure sequence was performed through an Interactive Telephone and Web Response System"			
Blinding of partical and personnel	Blinding of participants and personnel		"Masking: None (Open Label)". Included non-active comparator (cigarettes).			

Blinding of outco	ome	Low	"Masking: None (Open Label)". All primary outcomes objectively measured.
Incomplete outcome data		Low	Attrition: IQOS/CC=4.55% IQOS/NRT=5.56%, overall=4.84%. Exclusion: IQOS/CC=6.81% IQOS/NRT=5.5%, overall=6.45%.
Selective reporting		Low	All outcomes reported in at least one literature source.
NCT01967719			
Methods	Date of registration: 23/10/2013 Submitted to peer-reviewed journal within 12 months: No publication Published key outcomes on trial registration within 12 months: No Design: Crossover RCT Setting (Country): Confined (United States of America) Study start date; study end date: 02/10/2013; May 2014 Intervention duration: 2 sessions of single use of one cigarette, tobacco stick or 1 nasal spray in each nostril		
Participants	Number of participants: 62 randomised (44 HTP/CC, 18 HTP/NRT), 3 withdrawn (2 HTP/CC, 1 HTP/NRT), 60 completed (42 HTP/CC, 17 HTP/NRT) Withdrawal reasons reported: Yes Baseline characteristics: N= 62; Mean Age (SD): HTP/CC 37.2 (10.2) years, HTP/NRT 33.1 (7.3) years; Sex: 53% male; Ethnicity/Nationality: not reported Key inclusion criteria: Health status: "healthy"; ≥10 CPD; smoked for ≥3 years		
Interventions	Interventions: HTP (IQOS2.2 M), CC (Own brand M), NRT (Nicotrol nasal spray) Co-interventions: None Mode of exposure: Direct ad libitum		
Outcomes	Primary: Maximal nicotine concentration, Area under the concentration curve from start of product use to time of last quantifiable concentration Secondary: Exhaled Carbon monoxide, Adverse Events/Serious Adverse Events, Modified Cigarette/Product Evaluation Questionnaire, Questionnaire of Smoking Urges, Carboxyhemoglobin, Respiratory symptoms (inc. cough assessment), Physical examination, Clinical chemistry, haematology and urine analysis safety panel, Vital signs, Electrocardiogram, Time to reach nicotine Cmax, Spirometry, Concomitant medications, Cotinine, Terminal half-life of nicotine, Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity, Partial AUC		
Analyses	Sample size calculation reported: Yes Primary analysis population: Pharmacokinetic population defined as "all the randomized subjects who give informed consent, completed at least one of the single use Day 1 or Day 3, and for whom at least one PK parameter can been derived. Only subjects without major protocol deviations will be included in the PK analysis sets." Unit of analysis: Individuals		
Study funding	Philip Morris International (Industry-affiliated)		
Notes	_		ression analysis. Data obtained from published literature.
Risk of bias	I		•
Bias		Authors' judgement	Support for judgement
Random sequence generation	Random sequence generation		"Randomization to each product exposure sequence was done through an Interactive Telephone and Web Response System"
Allocation concealment		Low	"Randomization to each product exposure sequence was done through an Interactive Telephone and Web Response System"
Blinding of participants and personnel		High	"Masking: None (Open Label)". Included non-active comparator (cigarettes).
Blinding of outcome assessment		Low	"Masking: None (Open Label)". All primary outcomes objectively measured.
Incomplete outcome data Selective reporting		Low	Attrition: IQOS/CC=4.55% IQOS/NRT=0%, overall=3.23%. No subjects who completed the study were excluded from the analysis.

All outcomes reported in at least one literature source.

Selective reporting

Low

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Gee et al., 2018	(Actual	Use Study)				
Methods	Date o	f registration	: not registered			
		_	reviewed journal within 12 months: Unclear			
		shed key outcomes on trial registration within 12 months: Unclear				
		: Actual use s	_			
	_		Confined and Ambulatory (Japan)			
	1	-	udy end date: not reported			
	Interv	ention duration	on: Group $1 = 13$ days, Groups 2 and $3 = 9$ days, Group $4 = 1$ day			
Participants		per of participants: 208 (52 Group 1, 52 Group 2, 52 Group 3, 52 Group 4) drawal reasons reported: N/A				
			stics: N=208; Age, n participants: 21-29=58, 30-44=109, 45-65=40; nicity/Nationality: 100% Japanese			
	Key inclusion criteria: Health status: not specified; smokers ≥5 CPD, smoked for ≥6 months; THS users ≥5 product use sessions per day, using for ≥3 months					
Interventions	Interv	entions: Grou	p 1 (smokers): CC (T189 R), HTP (Glo1.0 R), HTP (IQOS R)			
			CC (322 M), HTP (Glo1.0 M)			
	Group	3 (THS users)	: HTP (Glo1.0 R), HTP (IQOS R)			
	Group	oup 4 (smokers): HTP (Glo1.0 R)				
	Co-int	erventions: N	Ione			
	Mode of exposure: Direct ad libitum					
Outcomes	Prima	Primary: Puffing topography, Mouth level exposure to nicotine free dry				
	particu	particulate matter, nicotine and menthol, Daily product consumption, Mouth insertion				
	depth					
	Secondary: None					
Analyses	Sampl	Sample size calculation reported: No				
	Prima	Primary analysis population: Not specified or unclear				
	Unit of analysis: Individuals					
Study funding	British American Tobacco (Industry-affiliated)					
Notes	Not inc	cluded in meta	-regression analysis.			
Risk of bias						
Bias		Authors'	Support for judgement			
Dius		judgement	Support for juagement			
Random sequence	ee	High	Non-randomised trial.			
generation						
Allocation conce Blinding of parti		High	Non-randomised trial.			
and personnel	•	Unclear	No information is provided in the text regarding blinding. Nonactive (CC) comparator.			
Blinding of outco		TT: 1	No information is provided in the text regarding blinding. Some			
assessment		High	primary outcomes were subjectively measured.			
Incomplete outco	ome	Unclear	Number of participants enrolled, completed and withdrawn was not			
data			reported.			
Selective reporting		Low	All outcomes listed in methods were reported on in the main results.			
Jones et al., 202						
Methods	Date of registration: not registered Submitted to peer-reviewed journal within 12 months: Unclear Published key outcomes on trial registration within 12 months: Unclear Design: Actual use study.					
	1	-	Confined and Ambulatory (Italy)			
	Study start date; study end date: not reported					
		ervention duration: Group 1 = 15 days, Group 2 = 10 days, Group 3 = 5 days				
Participants	Numb	er of participa	ants: 152 (50 Group 1, 50 Group 2, 52 Group 3)			

	Withdrawal reasons reported: N/A					
	Baseline characteristics: N=152; Age, n participants: 25-29=21, 30-44=67, 45-65=64;					
	l	50% male; Ethnicity/Nationality: 100% Italian				
		aclusion criteria: Health status: not specified; smokers ≥8 CPD, smoked for ≥7 vapers ≥1 product use per day, using for ≥6 months				
Interventions	Interv	erventions: Group 1 (smokers): EC (IS1.0[T]), HTP (IQOS2.4), CC (C651)				
	Group	2 (vapers): EO	C (Is1.0[T])			
	Group	o 3 (smokers): HTP (Glo1.0), CC (C651)				
	Co-int	terventions: None				
	Mode	of exposure: Direct ad libitum				
Outcomes	Prima	ary: Puffing topography, Mouth level exposure to nicotine free dry				
	particu	late matter and	d nicotine, Daily product consumption, Sensory questionnaire			
	Second	Secondary: None				
Analyses	Sampl	e size calculat	tion reported: No			
	Prima	ry analysis po	opulation: Not specified or unclear			
	Unit of	f analysis: Inc	lividuals			
Study funding	British	American To	bacco (Industry-affiliated)			
Notes	Not inc	cluded in meta	-regression analysis.			
Risk of bias						
Bias		Authors' judgement	Support for judgement			
Random sequence	e		Non-randomised trial.			
generation	_	High				
Allocation conce		High	Non-randomised trial.			
Blinding of partic	cipants	Unclear	No information is provided in the text regarding blinding. One active (EC) and one non-active (CC) comparator.			
Blinding of outco	ome	TT: -1-	No information is provided in the text regarding blinding. Some			
assessment		High	primary outcomes were subjectively measured.			
Incomplete outco	ome	Unclear	Number of participants enrolled, completed and withdrawn was not			
data Selective reporting	າຕ	Low	reported. All outcomes listed in methods were reported on in the main results.			
Dalrymple, 2022		LOW	7411 Outcomes fisted in methods were reported on in the main results.			
Methods		f registration	: not registered			
Methous	I	_				
		itted to peer-reviewed journal within 12 months: unclear shed key outcomes on trial registration within 12 months: unclear n: repeated measures				
	l					
	_	ng (Country): Confined (Germany)				
Study		udy start date; study end date: not reported				
		Intervention duration: 3 sessions of 32 puffs of Glo, ePen 3 or N491 cigarette				
Participants	Numb	er of particip	ants: 10 enrolled, 0 withdrawn, 10 completed			
•	Withdrawal reasons reported: N/A					
	Baseline characteristics: N=10; Age, n participants: 52.8; Sex: 30% male;					
	Ethnicity/Nationality: not reported					
	Key in	clusion criter	ia: Health status: "healthy"; non-smokers			
Interventions	Interventions: HTP (Glo), CC (N491), EC (ePen 3)					
	Co-interventions: None					
	Mode of exposure: Direct restricted					
Outcomes	Primary: Malondialdehyde; Catalase; Squalene; Squalene monohydroperoxide; Squalene					
	monohydroperoxide/Squalene ratio; L* (lightness); a* (green-red); b* (blue-yellow); Total					
	difference in colour from control (ΔΕ)					
	Secondary: Adverse Events/Serious Adverse Events					
Analyses	Sampl	e size calculat	tion reported: No			

	Primary analysis population: Not specified or unclear Unit of analysis: areas of skin					
Study funding	British American Tobacco (Industry-affiliated)					
Notes	Not included in meta-regression analysis.					
Risk of bias						
Bias						
Random sequence generation		N/A	Cochrane RoB tools designed to assess trials in which the unit of randomisation is people, rather than multiple sites on one individual,			
Allocation conce		N/A	selection bias cannot be fairly assessed using this tool on this study.			
Blinding of partic	cipants	Unclear	There is insufficient information provided in the text regarding blinding. One active (EC) and one non-active (CC) comparator.			
Blinding of outco		Low	No information is provided in the text regarding blinding, but all primary outcomes objectively measured.			
Incomplete outco	ome	Low	All participants completed the study and none withdrew.			
Selective reporting	ng	Low	All outcomes were reported on.			
NCT03452124			•			
Methods	Date o	f registration:	: 02/03/2018			
	Submi	tted to peer-r	eviewed journal within 12 months: Unclear			
		-	omes on trial registration within 12 months: Unclear			
	_	Design: Randomised controlled crossover followed by case control study				
	_	etting (Country): Confined and ambulatory (Greece)				
	_		art date; study end date: 30/03/2018; not reported			
			on: acute: 3x 7 minute sessions of sham cigarette, IQOS or cigarette			
D 411		c: 1 month				
Participants			ants: acute: 50 randomised, 0 withdrawn, 50 completed			
		ronic: 25 enrolled, 0 withdrawn, 25 completed ithdrawal reasons reported: N/A				
		seline characteristics: N=75; Age, n participants: 48 (acute) 26 (chronic); Sex: 48%				
			ile; Ethnicity/Nationality: not reported			
	Key in	clusion criter	clusion criteria: Health status: "healthy"; smokers ≥5 CPD			
Interventions	Interv	entions: Acute	e: HTP (IQOS), CC (Marlboro Red), sham cigarette			
	Chroni	hronic: HTPs (IQOS), CC (unknown brand)				
	Co-int	erventions: N	one			
	Mode	of exposure: I	Direct ad libitum			
Outcomes	subling reserve	qual arterial mi	e velocity; Exhaled Carbon Monoxide; Perfused boundary region of icrovessels; Global longitudinal strain of left ventricle; Coronary flow			
	Secondary: 11-dehydrothromboxane B2; Systolic blood pressure; Central Systolic blood pressure; Heart rate; Diastolic blood pressure; Protein carbonyls; Malondialdehyde; Myocardial work; Total arterial compliance; Augmentation index; Vital signs; Electrocardiogram; High-sensitivity C-reactive protein; Transforming growth factor-b; lipoprotein associated phospholipase A2; Tumor necrosis factor-a; Interleukin 6; Interleukin 10; Procollagen propeptide type III; Matrix metalloproteinase 2; Matrix metalloproteinase 9; Macrophage-colony stimulating factor; Flow-mediated dilation					
Analyses	_		tion reported: Yes			
			pulation: Not specified or unclear			
		f analysis: Ind				
Study funding		_	strian University of Athens (Independent)			
Notes	Not inc	cluded in meta	-regression analysis.			
Risk of bias	Risk of bias					

Bias		Authors' judgement	Support for judgement
Random sequence generation		Low	"Randomization was performed by an attending research nurse using a table of random numbers as reproduced from the online randomization software http://www.graphpad.com/quickcalcs/index.cfm"
Allocation concealment		Unclear	There is insufficient information provided to determine whether intervention allocation was concealed
Blinding of partic	cipants	Unclear	Trial registration states "Masking: Quadruple (Participant, Care Provider, Investigator, Outcomes Assessor)", but in the publication the only blinding described is in regard to outcome assessors.
Blinding of outco	ome	Low	"examinations were executed by a single, blindedto-treatment and to values of measured biomarkers, operator". Outcomes were physiological measures.
Incomplete outco	ome	Low	All participants completed the study and none withdrew.
Selective reporting	ng	High	Not all outcomes measured were reported on.
Iokeimidis, 2021	1		
Methods	Submi Publis Design Setting Study	tted to peer-r hed key outco a: Randomised g (Country): (start date; stu	eviewed journal within 12 months: unclear omes on trial registration within 12 months: unclear controlled crossover Confined (Greece) ady end date: note reported; not reported
	Interv	ention duratio	on: 3 sessions of 5 minutes use of IQOS, cigarette or cham cigarette
Turtespunts	Withdrawal reasons Baseline characteris Ethnicity/Nationality		stics: N=22; Age, n participants: 33, n=22; Sex: 45% male;
Interventions	Interventions: HTP (IQOS), Co-interventions: none Mode of exposure: direct ad		
Outcomes	Augme velocit	entation index;	Brachial systolic blood pressure; Aortic systolic blood pressure; Carotid–femoral pulse wave velocity; Brachial-ankle pulse wave
Analyses	Sampl	e size calculat	tion reported: Yes
randy ses	_		opulation: not specified or unclear
		f analysis: ind	1
Study funding		·	pol, Hippokration Hospital (ndependent)
Notes			-regression analysis.
	NOU INC	ruucu III IIIela	-regression analysis.
Risk of bias			S 4 6 3 4
Bias		Authors' judgement	Support for judgement
Random sequence generation	ee	Unclear	Whether or how participants were randomised is unclear.
Allocation conce	alment	Unclear	How interventions were allocated is not described.
Blinding of partic		Unclear	No information is provided in the text regarding blinding. Nonactive (CC) comparator.
Blinding of outco		Low	Outcomes were objectively measured.
Incomplete outco		Unclear	The authors state they "studied 22 current smokers" but it is unclear whether more than 22 were initially randomised or enrolled.
Selective reporting	ng	Low	Results data for all outcomes were reported.

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Yaman, 2021	Yaman, 2021					
Methods	Methods Date of registration: not registered					
	Submi	itted to peer-r	reviewed journal within 12 months: unclear			
	Published key outcomes on trial registration within 12 months: unclear					
			controlled crossover			
	١ `	.	confined (Cyprus)			
	l		udy end date: Not reported; not reported			
		Intervention duration: 3 sessions of 5 minutes use of IQOS or cigarettes				
Participants	1	Number of participants: 27 randomised, 0 withdrawn, 27 completed Withdrawal reasons reported: N/A				
		ne characteris ity/Nationality	stics: N=27; Age, n participants: 39.2, n=27; Sex: 59% male; root reported			
	Key in	clusion criter	ia: Health status: "healthy"; smoking history criteria not reported			
Interventions	Interv	entions: HTP	(IQOS), CC (own brand)			
	Co-int	erventions: n	one			
	Mode	of exposure:	Direct restricted			
Outcomes	Primary: A wave velocity; Diastolic blood pressure; E wave velocity; E/A ratio; Em/Am ratio; Heart rate; Left atrium diameter; Left ventricle ejection fraction; Left ventricle global circumferential strain; Left ventricle global longitudinal strain; Left ventricular end-diastolic diameter; Peak early diastolic velocity of the left ventricle; Peak late diastolic velocity of the left ventricle; Right atrium diameter; Right ventricle diameter; Right ventricle free wall strain; Right ventricle global longitudinal strain; Right ventricle peak early diastolic velocity; Right ventricle peak late diastolic velocity; Right ventricle systolic myocardial velocity; Right ventricle Em/Am ratio; Systolic blood pressure; Systolic myocardial velocity of the left ventricle; Tricuspid annular plane systolic excursion Secondary: none					
Analyses			e size calculation reported: No			
·	_	rimary analysis population: Not specified or unclear				
	Unit of	of analysis: individuals				
Study funding	Near E	ast University	and Mersin City Training and Research Hospital (Independent)			
Notes	Not inc	cluded in meta	ı-regression analysis.			
Risk of bias	'		·			
Bias		Authors' judgement	Support for judgement			
Random sequence generation	e	Unclear	Despite explaining the subjects were randomised, the sequence generation was not described in any of the study literature.			
Allocation conce		Unclear	Staff asked participants to use products, ie. They were aware. It is not clear if the order of interventions was randomised.			
Blinding of particular and personnel		Unclear	No information is provided in the text regarding blinding. Nonactive (CC) comparator.			
Blinding of outco		Low	Outcomes were physiological measures.			
data			Reasons for withdrawal are clearly described.			
Selective reporting		Low	All outcomes were reported on.			
	Phillips-Waller, 2021					
Methods		_	: not registered			
	l	_	reviewed journal within 12 months: unclear			
		•	omes on trial registration within 12 months: unclear			
Design: Non-randomised controlled crossover						
	1		confined (UK)			
	Study	start date; sti	udy end date: not reported; not reported			

		ention durati , Innokin iTas	on: 5 sessions of single use of IQOS, cigarette, JUUL, KangerTech te MVP 2	
Participants	Numb	er of particip	ants: 22 enrolled, 0 withdrawn, 22 completed	
	Withd	rawal reasons	s reported: N/A	
		ine characteristics: N=22; Age, n participants: 31, n=22; Sex: 82% male; city/Nationality: not reported		
	Key in	clusion criter	ia: Health status: "No serious illnesses"; smokers & vapers <1 CPD	
Interventions		entions: HTPS MVP 2)	S (IQOS), CC (own brand), EC (JUUL, KangerTech EVOD, Innokin	
	Co-int	erventions: n	one	
	Mode	of exposure:	direct ad libitum	
Outcomes Primary: Human Put concentration; Time t of product use to 30 m			offing/Smoking Topography (inc. puff count); Maximal nicotine to reach nicotine Cmax; Area under the concentration curve from start minutes; Nicotine; Nicotine boost effect; Urge To Smoke tandard questionnaire on user experience	
	Second	dary: none		
Analyses	Sampl	e size calculat	tion reported: no	
	Prima	ry analysis po	opulation: not specified or unclear	
	Unit of	f analysis: ind	lividuals	
Study funding	Tobaco	co Advisory G	roup project grant, Cancer Research UK (Independent)	
Notes	Not inc	cluded in meta	-regression analysis.	
Risk of bias	'			
Bias		Authors' judgement	Support for judgement	
Random sequence generation		High	Non-randomised trial	
Allocation concealment		High	Non-randomised trial	
Blinding of parti and personnel	•	Unclear	No information is provided in the text regarding blinding. One active (EC) and one non-active (CC) comparator.	
Blinding of outcome assessment		High	No information is provided in the text regarding blinding. Some primary outcomes were subjectively measured.	
Incomplete outcome		Unclear	The authors state they "studied 22 current smokers" but it is unclear	

Abbreviations: HTP=heated tobacco product; CC=combustible cigarette; EC=electronic cigarette; $Cess = cessation; \ NS = non-smoker; \ NRT = nicotine \ replacement \ the rapy; \ R = regular, \ M = menthol;$ CPD=cigarettes per day

Unclear

Low

data

Selective reporting

Supplementary Table 2. Outcomes measured and reported in heated tobacco product interventional trials.

All outcomes were reported on.

whether more than 22 were initially enrolled.

Outcome	Number of trials (measured)	Number of trials (reported)
Biomarkers of exposure		
Exhaled Carbon monoxide	26	21
2-cyanoethylmercapturic acid	14	14
Nicotine	14	11
Nicotine equivalents (molar sum of free nicotine, nicotine-glucuronide, free cotinine, cotinine-glucuronide, free trans-3'-hydroxycotinine, trans-3'-		
hydroxycotinine-glucuronide)	14	13
total N-nitrosonornicotine	14	13
3-hydroxypropylmercapturic acid	13	13

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monohydroxybutenylmercapturic acid	13	13
S-phenylmercapturic acid	13	12
total 1-hydroxypyrene	13	13
2-aminonaphthalene	12	12
4-aminobiphenyl	12	12
o-toluidine	12	12
2-hydroxyethylmercapturic acid	11	11
3-hydroxy-1-methylpropylmercapturic acid	11	11
Cotinine	10	8
3-hydroxybenzo[a]pyrene	9	9
1-aminonaphthalene	8	8
Cytochrome P450 2A6 activity	8	7
Ames mutagenicity test (YG1024+S9)	6	6
S-benzylmercepturic acid	4	4
N-acetyl-S-(2-carba-moylethyl)cysteine	2	2
N-acetyl-S-(2-hydroxy-2-carbamoylethyl)cysteine	2	2
4-hydroxybutyl-2-mercapturic acid	1	1
Cotinine	1	1
N-(2-cyanoethyl)valine haemoglobin adducts	1	1
Carboxyhemoglobin*	14	13
Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol*	15	15
Biomarkers of potential harm		
11-dehydrothromboxane B2	10	10
8-epi-prostaglandin F2alpha	9	9
Systolic blood pressure	8	8
White blood cell count	8	7
Diastolic blood pressure	7	7
Heart rate	7	5
High-sensitivity C-reactive protein	7	5
Fibrinogen Forced expiratory flow at 25–75% of forced vital capacity (aka Maximal mid-	6	4
expiratory flow)	6	3
Forced expiratory volume in one second	6	6
Forced vital capacity	6	3
High-density lipoprotein cholesterol	6	6
Homocysteine	6	4
Low-density lipoprotein cholesterol	6	4
Soluble intercellular adhesion molecule-1	6	6
Waist circumference	6	4
Weight	6	4
Hemoglobin glycosylated (Hemoglobin A1C)	5	4
Platelet count	5	4
Apolipoprotein A1	4	3
Apolipoprotein B	4	3
Augmentation index	4	2

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Forced expiratory volume in one second/forced vital capacity	4	2
Total cholesterol	4	3
Total lung capacity	4	3
Triglycerides	4	3
Blood glucose	3	3
Blood pressure	3	0
Functional residual capacity	3	2
Inspiratory capacity	3	2
Myeloperoxidase	3	2
Pulse wave velocity	3	2
Residual volume	3	3
Vital capacity	3	2
[Mean] Arterial Blood Pressure	2	1
4-Hydroxy-2-nonenal	2	0
Albumin	2	1
bronchodilator reversibility in FEV1	2	1
Carbon monoxide transfer coefficient	2	2
Diffusion Capacity	2	2
Flow-mediated dilation	2	2
Malondialdehyde	2	2
Peak Expiratory Flow	2	1
3-nitrotyrosine	1	0
8-Hydroxy-2'-deoxyguanosine	1	1
8-iso-prostaglandin F2alpha	1	1
A wave velocity	1	1
Aortic systolic blood pressure	1	1
Brachial systolic blood pressure	1	1
Brachial-ankle pulse wave velocity	1	1
Carotid–femoral pulse wave velocity	1	1
Catalase	1	1
Central Systolic blood pressure	1	1
change in bleedng on probing scores	1	1
change in gingival inflammation (GI) score	1	1
Change in mean full-mouth CAL	1	1
change in mean full-mouth PD	1	1
change in plaque control record	1	1
change in the number of sites with PD<4 mm, 4-5mm, 5-6 mm, 6-7 mm, and ≥7		
mm	1	1
change in tooth mobility (grade)	1	1
Coronary flow reserve	1	1
E wave velocity	1	1
E/A ratio	1	1
Em/Am ratio	1	1
Endothelin-1	1	0
<u>E-selectin</u>	1	0

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Expiratory reserve volume	1	1
Forced expiratory flow at X%	1	1
Global longitudinal strain of left ventricle	1	1
Glucose	1	0
H2O2 breakdown activity	1	1
H2O2 production	1	1
Interleukin 10	1	0
Interleukin 6	1	0
Left atrium diameter	1	1
Left ventricle ejection fraction	1	1
Left ventricle global circumferential strain	1	1
Left ventricle global longitudinal strain	1	1
Left ventricular end-diastolic diameter	1	1
lipoprotein associated phospholipase A2	1	0
Macrophage-colony stimulating factor	1	0
Matrix metalloproteinase 2	1	0
·	1	0
Matrix metalloproteinase 9 mean CAL change in sites with initial PD<4 mm, 4-5 mm, 5-6 mm, 6-7 mm, and	1	U
≥7mm	1	1
Mean clinical attachment level (CAL) change in sites with initial PD≥4mm after		
mechanical periodontal therapy	1	1
mean PD change in sites with initial PD<4mm, 4-5 mm, 5-6 mm, 6-7 mm, and ≥7	1	4
mm Mean PD change in sites with initial PD≥4 mm after mechanical periodontal	1	1
therapy	1	1
Mean Periodontal Pocket Depth (PD) reduction in all sites with initial PD ≥ 4 mm		
after mechanical periodontal therapy	1	1
Microbiological status	1	0
Monocyte chemotactic protein 1/C-C motif chemokine ligand 2	1	0
Myocardial work	1	1
Nitric oxide	1	1
nitric oxide bioavailability	1	1
Oxygen Saturation	1	1
Peak early diastolic velocity of the left ventricle	1	1
Peak late diastolic velocity of the left ventricle	1	1
Perfused boundary region of sublingual arterial microvessels	1	0
Plasminogen activator inhibitor-1	1	0
Procollagen propeptide type III	1	0
Pro-inflammatory and immuno-regulatory mediators (sCD40L, CRP, EGF,		
Eotaxin/CCL11, Flt3 ligand, GM-CSF, GRO, IFNα2, IL-1α, IL-1β, IL-1Ra, IL-2, IL-3, IL-		
4, IL-5, IL-6, IL-7, IL-8/CXCL8, IL-9, IL-10, IL-12 (p40), IL-12 (p70), IL-13, IL-15, IL-		
17A/CTLA8, IP10/CXCL10, MCP-1/CCL2, MCP-3/CCL7, MDC/CCL22, MIP-1α/CCL3, MIP-1β/CCL4, MMP-1, MMP-8, MMP-9, MMP-10, MMP-12, MMP-13,		
osteoprotegerin, PDGF-AA, PDGF-AB/BB, RANKL, RANTES/CCL5, TGFα, TIMP-1,		
TNF α , TNF β / LT- α)	1	0
Protein carbonyls	1	1
Respiratory impedance	1	1
Right atrium diameter	1	1
g diameter		

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Right ventricle diameter	1	1
Right ventricle Em/Am ratio	1	1
Right ventricle free wall strain	1	1
Right ventricle global longitudinal strain	1	1
Right ventricle peak early diastolic velocity	1	1
Right ventricle peak late diastolic velocity	1	1
Right ventricle systolic myocardial velocity	1	1
Soluble CD40 ligand	1	1
Soluble Nox2-derived peptide	1	1
Soluble P-selectin	1	1
Squalene	1	1
Squalene monohydroperoxide	1	1
Squalene monohydroperoxide/Squalene ratio	1	1
Systolic myocardial velocity of the left ventricle	1	1
Tissue plasminogen activator	1	0
Total anti-oxidant capacity	1	1
Total arterial compliance	1	1
Total respiratory resistances		1
Transforming growth factor-b	1	0
Tricuspid annular plane systolic excursion	1	1
Tumor necrosis factor-a	1	0
Vitamin E	1	1
Pharmacokinetic outcomes		
Time to reach nicotine Cmax	13	10
Maximal nicotine concentration	13 12	10 10
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last	12	10
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration	12	10
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine	12	10
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use	12 11 8	9 7
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity	12 11 8	10 9 7 5
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use)	12 11 8 6 5	10 9 7
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity	12 11 8	10 9 7 5 5
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC	12 11 8 6 5 5	10 9 7 5 5 4
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak	12 11 8 6 5 5 5	10 9 7 5 5 4 5
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes	12 11 8 6 5 5 5 4	10 9 7 5 5 4 5
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes	12 11 8 6 5 5 5 4 2	10 9 7 5 5 4 5 4
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours	12 11 8 6 5 5 5 4 2 1 1	10 9 7 5 5 4 5 4 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour	12 11 8 6 5 5 5 4 2 1 1 1	10 9 7 5 5 4 5 4 1 0 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax	12 11 8 6 5 5 5 4 2 1 1	10 9 7 5 5 4 5 4 1 0 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in	12 11 8 6 5 5 4 2 11 1 1 1	10 9 7 5 5 4 5 4 1 0 1 0 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in which Cpeak was observed	12 11 8 6 5 5 5 4 2 1 1 1 1 1	10 9 7 5 5 4 5 4 1 0 1 0 1 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in which Cpeak was observed Nicotine boost effect	12 11 8 6 5 5 5 4 2 1 1 1 1 1 1 1	10 9 7 5 5 4 5 4 1 0 1 1 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in which Cpeak was observed Nicotine boost effect Rate of plasma nicotine rise (speed of absorption)	12 11 8 6 5 5 5 4 2 1 1 1 1 1	10 9 7 5 5 4 5 4 1 0 1 0 1 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in which Cpeak was observed Nicotine boost effect Rate of plasma nicotine rise (speed of absorption) Questionnaires/Subjective effects	12 11 8 6 5 5 5 4 2 1 1 1 1 1 1 1	10 9 7 5 5 4 5 4 1 0 1 1 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in which Cpeak was observed Nicotine boost effect Rate of plasma nicotine rise (speed of absorption) Questionnaires/Subjective effects Modified Cigarette/Product Evaluation Questionnaire	12 11 8 6 5 5 5 4 2 1 1 1 1 1 1 1 1	10 9 7 5 5 4 5 4 1 0 1 0 1 1 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in which Cpeak was observed Nicotine boost effect Rate of plasma nicotine rise (speed of absorption) Questionnaires/Subjective effects	12 11 8 6 5 5 5 4 2 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	10 9 7 5 5 4 5 4 1 0 1 0 1 1 1 1
Maximal nicotine concentration Area under the concentration curve from start of product use to time of last quantifiable concentration Terminal half-life of nicotine Area under the plasma concentration-time curve from start of product use extrapolated from time of last quantifiable concentration to infinity Maximum observed nicotine concentration (following ad lib use) Partial AUC Time to nicotine Cpeak Weighted average nicotine concentration over 24 hours Nicotine molar metabolic ratio Area under the concentration curve from start of product use to 60 minutes Area under the concentration curve from start of product use to 60 minutes AUC from start of product use up to 12 hours Baseline adjusted AUC1hour Baseline adjusted Cmax Lowest observed plasma concentration during the same sampling interval in which Cpeak was observed Nicotine boost effect Rate of plasma nicotine rise (speed of absorption) Questionnaires/Subjective effects Modified Cigarette/Product Evaluation Questionnaire Questionnaire of Smoking Urges	12 11 8 6 5 5 4 2 1 1 1 1 1 1 1 1 1 1 1 1	10 9 7 5 5 4 5 4 1 0 1 1 1 1 1 1 1 1 1 1 1

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Intention to use [HTP] Questionnaire	4	2
Product Satisfaction Questionnaire	4	1
Prochaska "Stage of Change" Questionnaire	2	1
Product Liking Questionnaire	2	2
The Direct Effects of Nicotine Questionnaire	2	2
Urge To Smoke questionnaire	2	2
A visual analogue scale (VAS) assessing cigarette craving	1	1
Inhalation to non-inhalation ratios during HTP use	1	0
Product Evaluation Scale	1	0
Product preferene	1	1
Questionnaire (Other)	1	1
Sensory questionnaire	1	1
Smoking cessation quality of life questionnaire	1	0
The Direct Effects of Product scale	1	1
Urge For Product questionnaire	1	1
Safety Profile		
Adverse Events/Serious Adverse Events	23	23
Vital signs	19	11
Clinical chemistry, hematology and urine analysis safety panel	18	10
Physical examination	18	10
Electrocardiogram	16	10
Respiratory symptoms (inc. cough assessment)	16	11
Spirometry	14	9
Concomitant medications	13	9
Medical history	1	0
Other outcomes		
Daily product consumption	16	14
Human Puffing/Smoking Topography (inc. puff count)	13	10
Mouth level exposure to nicotine	4	3
Potential combustion occurances in tobacco plugs	4	4
Mouth level exposure to NFDPM	2	2
6-minute walking test	1	0
a* (green-red)	1	1
b* (blue-yellow)	1	1
Finger plethysmography	1	0
Full transcriptomics profile	1	0
L* (lightness)	1	1
Mouth insertion depth	1	1
Mouth level exposure to menthol	1	1
Oxysterols (6α-hydroxy-5α-cholestanol, 7α-hydroxycholesterol,		
5α,6αepoxycholestanol, 7-ketocholesterol, 7β-hydroxycholesterol, 5β,6β-		
epoxycholestanol, 24(R)-hydroxycholesterol, 25-hydroxycholesterol, 22(R)-		
hydroxycholesterol, 4ßhydroxycholesterol, and 27-hydroxycholesterol)	1	1
Total difference in colour from control (ΔE)	1	1
*Also measured as biomarkers of potential harm in one study		

Supplementary Table 3. Direction of effect in primary outcomes compared between heated tobacco and cigarette arms.

Trial ID	Primary Outcome(s)	Time point	Data	EoE between group difference*
767	Time to reach nicotine Cmax (min)	N/A	PNTV (median, range): 3.83, 2.83-7.83) CC (median, range): 3.83, 2.83-4.83	↔ (Positive)
000172	Maximal nicotine concentration (ng/mL)	N/A	PNTV (mean, 95% CI): 5.39, 4.34;6.69 CC (mean, 95% CI): 11.8, 9.49;14.6	(Negative)
UMIN000017297	Area under the concentration curve from start of product use to time of last quantifiable concentration (ng.h/mL)	N/A	PNTV (mean, 95% CI): 4.12, 3.43;4.95 CC (mean, 95% CI): 6.03, 5.02;7.25	↓ (Negative)
	3-hydroxypropylmercapturic acid (ug/24hr)	Day 5	NTV=484 ± 256 CC=1579 ± 696	↓ (Positive)
	2-cyanoethylmercapturic acid (ug/24hr)	Day 5	NTV=12.4 ± 6.6 CC=118.1 ± 64.7	↓ (Positive)
	4-aminobiphenyl (ng/24hr)	Day 5	NTV=1.8 ± 1.0 CC=12.3 ± 5.7	↓ (Positive)
	1-aminonaphthalene (ng/24hr)	Day 5	$NTV=5.7 \pm 3.2 CC=93.6 \pm 45.8$	↓ (Positive)
11.	2-aminonaphthalene (ng/24hr)	Day 5	$NTV=2.5 \pm 0.8 CC=26.3 \pm 12.2$	↓ (Positive)
UMIN000025777	S-phenylmercapturic acid (ng/24hr)	Day 5	NTV=276 ± 102 CC=2741 ± 1939	(Positive)
00NII	3-hydroxybenzo[a]pyrene (pg/24hr)	Day 5	NTV=48.7 ± 29.5 CC=156.3 ± 82.2	(Positive)
5	monohydroxybutenylmercapturi c acid (ng/24hr)	Day 5	NTV=219 ± 85 CC=1921 ± 1588	↓ (Positive)
	Exhaled Carbon Monoxide (ppm)	Day 5	NTV= 3.7 ± 1.8 CC= 25.6 ± 10.6	(Positive)
	4-hydroxybutyl-2-mercapturic acid (ug/24hr)	Day 5	$NTV=75.7 \pm 22.0 \text{ CC}=346.3 \pm 160.9$	↓ (Positive)
	2-hydroxyethylmercapturic acid (ng/24hr)	Day 5	NTV=844 ± 364 CC=3023 ± 2252	(Positive)
	Total 4-(methylnitrosamino)-1-(3-pyridyl)-1-butanol (ng/24hr)	Day 5	NTV=41.5 ± 30.3 CC=116.6 ± 75.1	(Positive)

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	Total N-nitrosonornicotine (pg/24hr)	Day 5	NTV=955 ± 604 CC=4986 ± 6644	↓ (Positive)
	o-Toluidine (ng/24hr)	Day 5	NTV=50.8 ± 20.2 CC=154.0 ± 41.0	(Positive)
	Total 1-hydroxypyrene (ng/24hr)	Day 5	NTV=208.7 ± 89.0 CC=332.4 ± 135.1	(Positive)
	Nicotine equivalents (mg/24hr)	Day 5	NTV=5.0 ± 3.0 CC=10.5 ± 4.8	↓ (Negative)
Caponnetto , 2018	Exhaled Carbon monoxide (ppm)	45 mins	Specific quantitative data was not provided, however based on the graph provided eCO was substantially lower in the HTP arms compared to the CC arm at all time points past baseline, with no overlapping error bars. Moreover, "repeated-measures ANOVA post-hoc comparisons showed significant differences between-product effect (iQOS/GLO vs own brand cigarette; P < 0.0001"	↓ (Positive)
	Exhaled Carbon monoxide (ppm)	55 mins	IQOS(mean, SE)=3.07, 0.32 CC(mean, SE)=6.47, 0.41	(Positive)
9689	Modified Cigarette/Product Evaluation Questionnaire	5 mins	All subscales of the mCEQ ("Smoking satisfaction", "Psychological reward", "Aversion", "Enjoyment of respiratory tract sensations", and "Craving reduction") were rated lower for the IQOSTM than for the tobacco cigarette.	↓ (Negative)
aspredicted.org #6896	Questionnaire of Smoking Urges	55 mins	"At T1 and T5, smoking resulted in lower craving scores compared to vaping (all ps < 0.01) and compared to using the IQOSTM (all ps < 0.01)"	↑ (Negative)
dicted	Fagerström Test for Nicotine/Cigarette Dependence		No relevant comparison (only reported at baseline)	N/A
aspre	Minnesota Nicotine Withdrawal Scale	55 mins	"At T5, no differences in withdrawal symptoms were present between smoking and using the IQOS [] ps>0.11"	↔ (Positive)
	A visual analogue scale (VAS) assessing cigarette craving	55 mins	IQOS(mean, SE)=58.20, 3.89 CC(mean, SE)=45.33, 4.05	↑ (Negative)
	Product preference		No relevant comparison (no HTP v CC comparison for outcome)	N/A
NCT03435 562	Nicotine (ng/mL)	5 mins post restricted use and 1-hour post <i>ad lib</i> use	Post-puff bout (mean, SD): IQOS=10.65 (6.20), CC=18.31 (11.39) Post ad lib (mean, SD): IQOS=5.97 (7.70), CC=12.23 (9.26)	↓ (Negative)

NCT03889990/ NCT03995329	No relevant comparison (no HTP						
30112	Soluble Nox2-derived peptide (pg/mL)			↓ (Positive)			
NCT0330112 9	Flow-mediated dilation (%)	Immediately after product use	IQOS (mean, SD)= 3.79 ± 2.68 CC (mean, SD)= 2.40 ± 1.89	↑ (Positive)			
NCT03364 751	Mean Periodontal Pocket Depth (PD) reduction in all sites with initial PD ≥ 4 mm after mechanical periodontal therapy (mm)	Month 6	IQOS (mean 95%CI)=-1.046, -1.194;-0.898 CC (mean, 95%CI)=-1.114, -1.258;-0.970. Mean difference=0.068 (-0.06; 0.196), p=0.297	↔ (Negative)			
	S-phenylmercapturic acid (pg/mg creat)	Day 90	CHTP (mean, 95%CI)=467, 365;597 CC (mean, 95%CI)=2652, 1853;3795	↓ (Positive)			
87	monohydroxybutenylmercapturi c acid (pg/mg creat)	Day 90	CHTP(mean, 95%CI)=420, 365;483 CC (mean, 95%CI)=2552, 1802;3612	(Positive)			
NCT02641587	Total 4-(methylnitrosamino)-1- (3-pyridyl)-1-butanol (pg/mg creat)	Day 90	CHTP(mean, 95%CI)=39.7, 29.3;53.7 CC (mean, 95%CI)=196.7, 117;245.0	↓ (Positive)			
NC	3-hydroxypropylmercapturic acid (ng/mg creat)	Day 90	CHTP(mean, 95%CI)=378.2, 334.6;427.6 CC (mean, 95%CI)=966.0, 786.4;1187	↓ (Positive)			
	Carboxyhemoglobin (%)	Day 90	CHTP(mean, 95%CI)=1.94, 1.78;2.13 CC (mean, 95%CI)=4.33, 3.69; 5.07	(Positive)			
	8-epi-prostaglandin F2alpha (pg/mg creat)	Month 6	FAS-AR (mean 95%CI): IQOS=330 (316;345) CC=349 (335;364) FAS-EX (mean 95%CI): IQOS=326 (309;345) CC=350 (336;365) [p=0.018]	$\leftrightarrow \leftrightarrow$ (Negative)			
6381	11-dehydrothromboxane B2 (pg/mg creat)	Month 6	FAS-AR (mean 95%CI): IQOS=511 (475;549) CC=527 (492;565) FAS-EX (mean 95%CI): IQOS=502 (458;550) CC=527 (491;564) [p=0.193]	$\leftrightarrow \leftrightarrow$ (Negative)			
NCT02396381	Total 4-(methylnitrosamino)-1- (3-pyridyl)-1-butanol	Month 6	FAS-AR (mean 95%CI): IQOS=198 (178;220) CC=282 (254;312) FAS-EX (mean 95%CI): IQOS=159 (140;181) CC=281 (254;311) [p<0.001*]	↓↓ (Favourable)			
NC	Carboxyhemoglobin (%)	Month 6	FAS-AR (mean 95%CI): IQOS=3.46 (3.18;3.77) CC=4.40 (4.06;4.78) FAS-EX (mean 95%CI): IQOS=2.95 (2.69;3.24) CC=4.35 (4.03;4.70) [p<0.001*]	↓↓ (Positive)			
	White blood cell count (GI/L)	Month 6	FAS-AR (mean 95%CI): IQOS=7.26 (7.05;7.48) CC=7.53 (7.33;7.74)	↔ ↓			

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			FAS-EX (mean 95%CI): IQOS=7.06 (6.81;7.31) CC=7.48 (7.28;7.68) [p=0.001*]	(Unclear)
	Soluble intercellular adhesion molecule-1 (ng/mL)	Month 6	FAS-AR (mean 95%CI): IQOS=260 (253;266) CC=264 (257;271) FAS-EX (mean 95%CI): IQOS=257 (249;265) CC=265 (258;271) [p0.030]	$\leftrightarrow \leftrightarrow$ (Negative)
	High-density lipoprotein cholesterol (mg/dL)	Month 6	FAS-AR (mean 95%CI): IQOS=54.6 (53.5;55.8) CC=51.8 (50.6;52.9) FAS-EX (mean 95%CI): IQOS=54.6 (53.1;56.2) CC=51.6 (50.4;52.7) [p<0.001*]	↑↑ (Positive)
	Forced expiratory volume in one second (% pred)	Month 6	FAS-AR (mean 95%CI): IQOS=94.4 (93.6;95.1) CC=93.1 (92.4;93.9) FAS-EX (mean 95%CI): IQOS=94.4 (93.4;95.3) CC=93.1 (92.3;93.9) [p=0.008*]	↔↑ (Unclear)
30412	Maximal nicotine concentration (ng/mL)	N/A	CHTP(mean, 95% CI)=6.2950, 5.2610;7.5322 CC(mean, 95%CI)=9.8463, 8.2290;11.7815 Mean ratio=63.9326% (49.6045;82.3991 [95%])	↓ (Negative)
NC102400412	Area under the concentration curve from start of product use to time of last quantifiable concentration (h*ng/mL)	N/A	CHTP(mean, 95% CI)=8.5311, 6.9550;10.4642 CC(mean, 95%CI)=14.2172, 11.5908;17.4388 Mean ratio=60.0052% (44.9517;80.0997 [95%])	↓ (Negative)
† 	monohydroxybutenylmercapturi c acid (pg/mg creat)	Day 5	CHTP (mean, 95%CI)=339.73 (301.82;382.42) CC(mean, 95%CI)=1840.61 (1275.38;2656.32)	↓ (Positive)
c7¢0¢	3-hydroxypropylmercapturic acid (pg/mg creat)	Day 5	CHTP (mean, 95%CI)=494.70 (417.53;586.12) CC(mean, 95%CI)=1187.97 (1026.63;1374.65)	↓ (Positive)
NCT02503254	Carboxyhemoglobin (%)	Day 5	CHTP (mean, 95%CI)=2.7 (2.2; 3.2) CC(mean, 95%CI)=6.4 (5.7; 7.1)	↓ (Positive)
Z	S-phenylmercapturic acid (pg/mg creat)	Day 5	CHTP (mean, 95%CI)=361.48 (289.26; 451.74) CC(mean, 95%CI)=2898.46 (2172.62; 3866.79)	(Positive)
	Carboxyhemoglobin (%)	Week 52	IQOS (mean, 95%CI)=2.59, 2.24;3.01 CC (mean, 95%CI)=4.06, 3.77;4.38 % relative reduction=31.7 (23.3;39.1[95%])	↓ (Positive)
00	8-epi-prostaglandin F2alpha (pg/mg creat)	Week 52	IQOS (mean, 95%CI)=307, 279;338 CC (mean, 95%CI)=327, 307;348 % relative reduction=7.15 (-1.03;14.7[95%])	↔ (Negative)
INC 1 02049330	Total 4-(methylnitrosamino)-1- (3-pyridyl)-1-butanol (pg/mg creat)	Week 52	IQOS (mean, 95%CI)=133, 105;168 CC (mean, 95%CI)=269, 238;305 % relative reduction=46.3 (36.2;54.8[95%])	↓ (Positive)
NC	11-dehydrothromboxane B2 (pg/mg creat)	Week 52	IQOS (mean, 95%CI)=582, 518;654 CC (mean, 95%CI)=586, 538;638 % relative reduction=3.44 (-8.74;14.3[95%])	↔ (Negative)
	White blood cell count (GI/L)	Week 52	IQOS (mean, 95%CI)=6.73, 6.47;6.99 CC (mean, 95%CI)=7.31, 7.07;7.54 LS mean diff=-0.413 (-0.694;-0.131[95%])	↓ (Positive)

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	Soluble intercellular adhesion molecule-1 (ng/mL)	Week 52	IQOS (mean, 95%CI)=246, 230;263 CC (mean, 95%CI)=258, 244;272 % relative reduction=3.11 (0.0231;6.10[95%])	↔ (Negative)
	High-density lipoprotein cholesterol (mg/dL)	Week 52	IQOS (mean, 95%CI)=52.2, 49.5;54.8 CC (mean, 95%CI)=50.6, 48.9;52.3 Mean diff=1.75 (-0.160;3.65[95%])	↔ (Negative)
	Forced expiratory volume in one second (% pred)	Week 52	IQOS (mean, 95%CI)=93.2, 91.1;95.2 CC (mean, 95%CI)=92.3, 90.7;94.0 Mean diff=0.914 (-0.339;2.17[95%])	↔ (Negative)
90229	Maximal nicotine concentration (ng/mL)	N/A	IQOS (mean, 95%CI)=10.70, 8.94;12.8 CC (mean, 95%CI)=12.09, 10.10;14.47 Mean ratio=88.47 (68.64;114.03[95%])	↔ (Positive)
NCT01967706	Area under the concentration curve from start of product use to time of last quantifiable concentration (h*ng/mL)	N/A	IQOS (mean, 95%CI)=23.99, 20.87;27.57 CC (mean, 95%CI)=24.45, 21.27;28.10 Mean ratio=98.13 (80.61;119.46[95%])	↔ (Positive)
88908	Maximal nicotine concentration (ng/mL)	N/A	IQOS (mean, 95%CI)=8.4, 6.8;10.3 CC (mean, 95%CI)=11.9, 9.5;14.9 Mean ratio=70.3% (60.0;82.2[90%])	↓ (Negative)
NCT01780688	Area under the concentration curve from start of product use to time of last quantifiable concentration (h*ng/mL)	N/A	IQOS (mean, 95%CI)=17.7, 15.0;20.8 CC (mean, 95%CI)=22.8, 19.4;26.8 Mean ratio=77.4% (70.5;85.0[90%])	↓ (Negative)
4	monohydroxybutenylmercapturi c acid (pg/mg creat)	Day 5	% reduction IQOS/CC mean (95%CI)=88.5 (84.7–91.4) [p<0.001]	↓ (Positive)
NCT01780714	3-hydroxypropylmercapturic acid (pg/mg creat)	Day 5	% reduction IQOS/CC mean (95%CI)=72.1 (67.4–76.1) [p<0.001]	↓ (Positive)
CT01	Carboxyhemoglobin (%)	Day 5	% reduction IQOS/CC mean (95%CI)=76.7 (74.3–78.9) [p<0.001]	↓ (Positive)
Z	S-phenylmercapturic acid (pg/mg creat)	Day 5	% reduction IQOS/CC mean (95%CI)=93.0 (90.6–94.9) [p<0.001]	(Positive)
898	Maximal nicotine concentration (ng/mL)	N/A	HTP (mean, SD)=1.18±1.13 CC (mean, SD)=7.76±4.65 [p<0.05]	↓ (Negative)
ISRCTN8868 2435	Area under the concentration curve from start of product use to time of last quantifiable concentration (h*ng/mL)	N/A	HTP (mean, SD)=1.07±0.75 CC (mean, SD)=5.97±2.15 [p<0.05]	↓ (Negative)

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	Time to reach nicotine Cmax (min)	N/A	HTP (median, min-max)=9.02, 2.05-31.0 CC (median, min-max)=5.02, 3.90-20.0 [p<0.05]	↑ (Negative)
Nga, 2020	Exhaled Carbon monoxide (ppm)	45 mins post product use	IQOS mean=4.67 CC mean=16.47 (no variance values provided but error bars do not overlap in graph presented) Between product effect significant difference (repeated-measures ANOVA, p<0.001)	↓ (Positive)
	Exhaled Carbon monoxide (ppm)	Immediately after bout 1 (at 5mins) and 2 (at 65mins)	Bout 1, mean (SD): CC=12.1 (3.4) LLTV=not reported [CC sig higher than LLTV, cohens d=2.4] Bout 2, mean (SD): CC= 16.9 (5.8) LLTV=4.5 (2.1) [CC sig higher than LLTV, cohens d=2.9]	↓ (Positive)
	Questionnaire of Smoking Urges	Immediately after bout 1 (at 5mins) and 2 (at 65mins)	"There were no significant differences between any of the conditions immediately following either bout"	\leftrightarrow (Positive)
	Nicotine (ng/mL)	Immediately after bout 1 (at 5mins) and 2 (at 65mins)	Bout 1, mean (SD): CC=24.4 (12.6) LLTV=14.3 (8.1) [CC sig higher than LLTV, cohens d=1.0] Bout 2, mean (SD): CC= 23.7 (14.5) LLTV=16.4 (11.3) [CC higher than LLTV but not significantly]	↓ (Negative)
, 2016	Minnesota Nicotine Withdrawal Scale	Immediately after bout 1 (at 5mins) and 2 (at 65mins)	"There were no significant differences between any of the conditions immediately following either bout."	↔ (Positive)
Lopez,	The Direct Effects of Nicotine Questionnaire	Immediately after bout 1 (at 5mins) and 2 (at 65mins)	"there were no differences between the [CC] and LLTV conditions at that same time point [Bout 1]. There were no significant differences between any of the conditions immediately following bout 2."	↔ (Negative)
	The Direct Effects of Product scale	Immediately after bout 1 (at 5mins) and 2 (at 65mins)	"Was the product satisfying?": "Immediately following bout 1, the mean score for the OB condition of 93.3 (10.51) was significantly higher compared to the scores of 51.2 (30.9) for the LLTV condition (d = 1.8) [] There was a similar pattern following bout 2" "Did the product taste good?": "immediately following bout 1, the mean score for the OB condition of 92.9 (11.4) was significantly higher compared to the score of 43.7 (31.8) for the LLTV condition $[t(14) = 5.2, p < 0.017; d = 2.1]$ "Did the product calm you down?": "immediately following bout 1, the mean score for the OB condition of 68.4 (28.9) was significantly higher compared to the LLTV score of 41.8 (31.2; $[t(14) = 4.1, p < 0.017; d = 0.9]$) [] There were no significant differences between any of the conditions immediately following bout 2"	↓ (Negative)

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7	Augmentation index	No results repor		Not reported
75.	8-epi-prostaglandin F2alpha	No results repor		Not reported
ISRCTN810757 60	Total 4-(methylnitrosami no)-1-(3-pyridyl)-1-butanol (ng/24h)	Day 90	Graph shows levels were equivalent at baseline PP-population mean change baseline-day 90: Glo=-120 CC=-12 Diff (99.94% CI)=-108 (-168;-48) [p<0.0001] CEVal-population mean change baseline-day 90: Glo=-110 CC=-5 Diff (99.94% CI)=-105 (-193; -17)	↓ (Positive)
	Time to reach nicotine Cmax (min)	N/A	Median (range): Glo1.0=4.1, 1.1-45.0 Glo1.1=4.1, 1.2-15.4 CC=6.0, 3.0-9.1	↔ (Positive)
	Maximal nicotine concentration (ng/mL)	N/A	Mean (90%CI): Glo1.0=8.7 (6.93;10.95) Glo1.1=10.9 (8.63;13.70) CC=23.3 (18.46;29.33)	↓ (Negative)
ISRCTN13439529	Area under the concentration curve from start of product use to time of last quantifiable concentration (min*ng/mL)	N/A	Mean (90%CI): Glo1.0=527 (438.7;633.3) Glo1.1=695 (577.6;835.6) CC=1374 (1142.4;1653.1)	↓ (Negative)
RCTN	Intention to use [HTP] Questionnaire	240 min post 1st puff	Mean±SD, median: Glo1.0=2.5 ± 2.67, 2.0 Glo1.1=3.1 ± 2.84, 2.0 CC=9.1 ± 1.37, 10	↓ (Negative)
ISI	Product Liking Questionnaire	3-240min	Mean±SD, median: Glo1.0=720 ± 733, 640 Glo1.1=820 ± 724, 675 CC=2107 ± 403, 2281	↓ (Negative)
	Urge To Smoke questionnaire	5 min post 1st puff	Mean±SD, median: Glo1.0=5.0 \pm 3.33, 5.0 Glo1.1=4.8 \pm 3.27, 5.0 CC=2.6 \pm 3.50, 1.0	↓ (Negative)
	Urge For Product questionnaire	No comparison	to cigarette arm	N/A
	Exhaled Carbon monoxide (ppm)	Day 7	CC(mean)=20.30, Glo(mean)=3.40, IQOS(mean)=3.40, mCC(mean)=20.07, mGlo(mean)=2.80	↓ (Positive)
_	Total 4-(methylnitrosamino)-1- (3-pyridyl)-1-butanol (ng/24h)	Day 7	CC(mean)=197.85, Glo(mean)=128.63, IQOS(mean)=80.35, mCC(mean)=167.02, mGlo(mean)=149.38	↓ (Positive)
)1360, 24988	2-cyanoethylmercapturic acid (ug/24h)	Day 7	CC(mean)=159.04, Glo(mean)=17.84, IQOS(mean)=16.54, mCC(mean)=165.62, mGlo(mean)=21.03	(Positive)
N143(total N-nitrosonornicotine (ng/24h)	Day 7	CC(mean)=15.36, Glo(mean)=5.85, IQOS(mean)=1.06, mCC(mean)=9.62, mGlo(mean)=5.57	↓ (Positive)
ISRCTN14301360/ UMIN000024988	Nicotine equivalents (mg/24h)	Day 7	CC(mean)=8.33, Glo(mean)=6.15, IQOS(mean)=7.58, mCC(mean)=9.77, mGlo(mean)=5.75	(Negative)
H	monohydroxybutenylmercapturi c acid (ng/24h)	Day 7	CC(mean)=770.64, Glo(mean)=49.87, IQOS(mean)=118.38, mCC(mean)=1010.18, mGlo(mean)=98.40	(Positive)
	3-hydroxypropylmercapturic acid (ug/24h)	Day 7	CC(mean)=1448.93, Glo(mean)=568.66, IQOS(mean)=639.21, mCC(mean)=1422.37, mGlo(mean)=656.99	(Positive)

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	total 1-hydroxypyrene (ng/24h)	Day 7	CC(mean)=172.86, Glo(mean)=75.58, IQOS(mean)=50.18, mCC(mean)=195.19, mGlo(mean)=63.46	↓ (Positive)
	S-phenylmercapturic acid (ug/24h)	Day 7	CC(mean)=2.25, Glo(mean)=0.20, IQOS(mean)=0.19 mCC(mean)=2.81, mGlo(mean)=0.20	(Positive)
	o-toluidine (ng/24h)	Day 7	CC(mean)=153.21, Glo(mean)=58.52, IQOS(mean)=54.81, mCC(mean)=119.04, mGlo(mean)=39.39	(Positive)
	4-aminobiphenyl (ng/24h)	Day 7	CC(mean)=10.86, Glo(mean)=2.45, IQOS(mean)=2.25, mCC(mean)=10.44, mGlo(mean)=2.31	↓ (Positive)
	2-aminonaphthalene (ng/24h)	Day 7	CC(mean)=17.80, Glo(mean)=1.74, IQOS(mean)=1.72, mCC(mean)=17.65, mGlo(mean)=1.92	↓ (Positive)
	3-hydroxy-1- methylpropylmercapturic acid (ug/24h)	Day 7	CC(mean)=385.50, Glo(mean)=79.00, IQOS(mean)=79.63, mCC(mean)=362.45, mGlo(mean)=73.23	↓ (Positive)
	2-hydroxyethylmercapturic acid (ug/24h)	Day 7	CC(mean)=5.08, Glo(mean)=2.46, IQOS(mean)=2.60 mCC(mean)=7.13, mGlo(mean)=2.84	(Positive)
	N-acetyl-S-(2-hydroxy-2- carbamoylethyl)cysteine (ug/24h)	Day 7	CC(mean)=17.24, Glo(mean)=15.68, IQOS(mean)=13.75, mCC(mean)=16.40, mGlo(mean)=15.36	↓ (Positive)
	N-acetyl-S-(2-carba- moylethyl)cysteine (ug/24h)	Day 7	CC(mean)=111.65, Glo(mean)=91.75, IQOS(mean)=65.76, mCC(mean)=114.96, mGlo(mean)=88.82	↓ (Positive)
	Nicotine	Not reported		Not reported
DRKS00 012919	Systolic blood pressure (mm Hg)	120 min	Based on graph presenting data throughout study period, SPB at end of exposure was not substantially different between the HTP and CC arms	↔ (Negative)
	Exhaled Carbon monoxide (ppm)	Day 7	CC(mean)=25.3 Glo(mean)=4.4	↓ (Positive)
606	Total 4-(methylnitrosamino)-1- (3-pyridyl)-1-butanol (ng/24h)	Day 7	CC(mean)=289.54 Glo(mean)=195.71	↓ (Positive)
306519	2-cyanoethylmercapturic acid (mg/24h)	Day 7	CC(mean)=0.24 Glo(mean)=0.03	↓ (Positive)
ISRCTN80651909	total N-nitrosonornicotine (ng/24h)	Day 7	CC(mean)=10.85 Glo(mean)=6.10	↓ (Positive)
ISR	Nicotine equivalents (mg/24h)	Day 7	CC(mean)=14.88 Glo(mean)=7.37	↓ (Positive)
	monohydroxybutenylmercapturi c acid (ng/24h)	Day 7	CC(mean)=2552.74 Glo(mean)=240.28	↓ (Positive)

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3-hydroxypropylmercapturic acid (mg/24h)	Day 7	CC(mean)=1.37 Glo(mean)=0.27	↓ (Positive)
total 1-hydroxypyrene (ng/24h)	Day 7	CC(mean)=313.33 Glo(mean)=106.71	(Positive)
S-phenylmercapturic acid (ng/24h)	Day 7	CC(mean)=5572.79 Glo(mean)=231.36	(Positive)
o-toluidine (ng/24h)	Day 7	CC(mean)=146.60 Glo(mean)=38.40	(Positive)
4-aminobiphenyl (ng/24h)	Day 7	CC(mean)=22.36 Glo(mean)=3.36	(Positive)
2-aminonaphthalene (ng/24h)	Day 7	CC(mean)=32.38 Glo(mean)=3.03	(Positive)
N-acetyl-S-(2-hydroxy-2- carbamoylethyl)cysteine (ng/24h)	Day 7	CC(mean)=33554.88 Glo(mean)=24749.07	↓ (Positive)
N-acetyl-S-(2-carba- moylethyl)cysteine (mg/24h)	Day 7	CC(mean)=0.18 Glo(mean)=0.12	↓ (Positive)
3-hydroxy-1- methylpropylmercapturic acid (mg/24h)	Day 7	CC(mean)=0.54 Glo(mean)=0.07	↓ (Positive)
2-hydroxyethylmercapturic acid (ng/24h)	Day 7	CC(mean)=9673.61 Glo(mean)=3954.5	↓ (Positive)
8-epi-prostaglandin F2alpha (no units reported)	Day 7	"8-epi-PGF2α Type III, there was no significant change in all arms except the glo THP arm, which showed a significant decrease in the urinary levels of this BoBE" Without baseline data there is no way to know whether the end of exposure levels differed between study arms.	Unclear
White blood cell count (no units reported)	Day 7	"white blood cell count was significantly reduced between baseline and Day 7 for the glo THP and the prototype EC arms. However, there was no significant change in the nicotine cessation or in the control cigarette arm." Without baseline data there is no way to know whether the end of exposure levels differed between study arms.	Unclear
Nicotine molar metabolic ratio (no units reported)	Day 7	CC(mean)=2.74 Glo(mean)=3.31	↑ (Unclear)

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Exhaled Carbon monoxide units reported)	no Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
Total 4-(methylnitrosamine (3-pyridyl)-1-butanol (no u reported)		"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
2-cyanoethylmercapturic a (no units reported)	cid Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
total N-nitrosonornicotine units reported)	(no Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
monohydroxybutenylmerc c acid (no units reported)	apturi Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group" This is substantiated by the graph presented	↓ (Positive)
3-hydroxypropylmercaptur acid (no units reported)	Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
total 1-hydroxypyrene (no reported)	units Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
S-phenylmercapturic acid units reported)	(no Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
o-toluidine (no units report	ted) Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
4-aminobiphenyl (no units reported)	Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
2-aminonaphthalene (no un reported)	Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
3-hydroxy-1- methylpropylmercapturic a (no units reported)	acid Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)

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	2-hydroxyethylmercapturic acid (no units reported)	Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
	3-hydroxybenzo[a]pyrene (no units reported)	Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
	1-aminonaphthalene (no units reported)	Day 5	"After the five-day study period, the levels of Biomarkers of Exposure (BoEs) of the selected HPHCs observed in the four heated tobacco product groups were significantly reduced to that observed in the CC group"	↓ (Positive)
	Time to reach nicotine Cmax (mins)	N/A	Controlled (mean(SD)): CC=6.71 (5.11) IQOS=5.41 (1.36) Ad lib (mean(SD)): CC=5.84 (1.36) IQOS=6.38 (5.06)	↔ (Positive)
112	Maximal nicotine concentration	Not reported		Not reported
NCT03700112	Baseline adjusted Cmax (ng/mL)	N/A	Controlled (mean(SD)): CC=21.2 (11.7) IQOS=16.1 (7.7) Ad lib (mean(SD)): CC=27.9 (19.6) IQOS=17.4 (7.3)	(Negative)
CT03	Baseline adjusted AUC1hour (hrs*ng/mL)	N/A	Controlled (mean(SD)): CC=7.67 (3.56) IQOS=5.15 (2.32) Ad lib (mean(SD)): CC=9.76 (5.69) IQOS=5.72 (1.88)	(Negative)
Ż	Area under the concentration curve from start of product use to 60 minutes	Not reported		Not reported
	Total 4-(methylnitrosamino)-1- (3-pyridyl)-1-butanol (pg/mg creat)	Day 90	mIQOS (mean, 95%CI)=23.23 (19.34;27.91) CC (mean, 95%CI)=95.03 (77.31;116.82)	↓ (Positive)
70995	monohydroxybutenylmercapturi c acid (pg/mg creat)	Day 90	mIQOS (mean, 95%CI)=141.74 (120.62;166.57) CC (mean, 95%CI)=785.27 (576.82;1069.04)	↓ (Positive)
NCT01970995	3-hydroxypropylmercapturic acid (ng/mg creat)	Day 90	mIQOS (mean, 95%CI)=386.37 (356.30;418.97) CC (mean, 95%CI)=695.58 (602.43;803.13)	↓ (Positive)
N	Carboxyhemoglobin (%)	Day 90	mIQOS (mean, 95%CI)=2.97 (2.88;3.06) CC (mean, 95%CI)=5.73 (5.24;6.25)	↓ (Positive)
	S-phenylmercapturic acid (pg/mg creat)	Day 90	mIQOS (mean, 95%CI)=145.58 (121.67;174.18) CC (mean, 95%CI)=1157.25 (848.59;1578.17)	↓ (Positive)
156	Total 4-(methylnitrosamino)-1- (3-pyridyl)-1-butanol (pg/mg creat)	Day 90	mIQOS (mean, 95%CI)=47.53 (34.80;64.91) CC (mean, 95%CI)=152.11 (108.38;213.47)	↓ (Positive)
NCT01989156	monohydroxybutenylmercapturi c acid (pg/mg creat)	Day 90	mIQOS (mean, 95%CI)=260.98 (205.28;331.79) CC (mean, 95%CI)=1040.71 (677.79;1597.94)	↓ (Positive)
NCT	3-hydroxypropylmercapturic acid (ng/mg creat)	Day 90	mIQOS (mean, 95%CI)=314.05 (281.51;350.34) CC (mean, 95%CI)=606.10 (468.27;784.48)	↓ (Positive)
	Carboxyhemoglobin (%)	Day 90	mIQOS (mean, 95%CI)=2.66 (2.40;2.94)	

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			CC (mean, 95%CI)=5.62 (5.00;6.32)	(Positive)
	S-phenylmercapturic acid		mIQOS (mean, 95%CI)=314.02 (219.66;448.93)	(1 OSILIVE)
	(pg/mg creat)	Day 90	CC (mean, 95%CI)=1218.56 (822.54;1805.25)	(Positive)
	monohydroxybutenylmercapturi		IQOS (mean, 95%CI)=107.39 (97.24;118.60)	(1 OSHIVE)
	c acid (pg/mg creat)	Day 5	CC (mean, 95%CI)=450.19 (300.07;675.42)	(Positive)
982	3-hydroxypropylmercapturic		IQOS (mean, 95%CI)=311.08 (279.59;346.12)	(1 ositive)
NCT01970982	acid (ng/mg creat)	Day 5	CC (mean, 95%CI)=599.67 (511.70;702.76)	(Positive)
)19			IQOS (mean, 95%CI)=2.39 (2.32;2.46)	(1 ositive)
Ĺ.	Carboxyhemoglobin (%)	Day 5	CC (mean, 95%CI)=5.14 (4.66;5.66)	(Positive)
ž	S-phenylmercapturic acid		IQOS (mean, 95%CI)=143.77 (126.08;163.93)	(1 oshive)
	(pg/mg creat)	Day 5	CC (mean, 95%CI)=850.02 (620.40;1164.63)	(Positive)
	monohydroxybutenylmercapturi		IQOS (mean, 95%CI)=192.93 (174.90; 212.83)	(1 OSHIVE)
-,	c acid (pg/mg creat)	Day 5	CC (mean, 95%CI)=2399.40 (1884.60; 3054.83)	(Positive)
932	3-hydroxypropylmercapturic		IQOS (mean, 95%CI)=402.26 (366.55; 441.45)	(1 oshive)
29	acid (ng/mg creat)	Day 5	CC (mean, 95%CI)=931.01 (825.73; 1049.72)	(Positive)
011			IQOS (mean, 95%CI)=1.06 (1.03; 1.08)	(1 dshirte)
NCT01959932	Carboxyhemoglobin (%)	Day 5	CC (mean, 95%CI)=4.51 (4.05; 5.01)	(Positive)
ž	S-phenylmercapturic acid	Day 5	IQOS (mean, 95%CI)=164.45 (144.45; 187.22)	<u> </u>
	(pg/mg creat)		CC (mean, 95%CI)=2922.81 (2362.80; 3615.54)	(Positive)
	Maximal nicotine concentration	27/1	IQOS (geo mean, 95%CI)=14.30 (11.41;17.91) CC (geo mean, 95%CI)=13.82	↔
NCT0195960 7	(ng/mL)	N/A	(11.00;17.35)	(Positive)
5 7	Area under the concentration		IQOS (geo mean, 95%CI)=23.75 (19.74;28.58) CC (geo mean, 95%CI)=24.66	
<u> </u>	curve from start of product use	NT/A	(20.24;30.03)	\leftrightarrow
کِ	to time of last quantifiable	N/A		(Positive)
4	concentration (ng.h/mL)			
∞	Maximal nicotine concentration	N/A	IQOS (mean, 95%CI)=9.60 (7.64;12.07)	\leftrightarrow
_	(ng/mL)	IV/A	CC (mean, 95%CI)=12.34 (10.47;14.54)	(Positive)
NCT0196773 2	Area under the concentration		IQOS (mean, 95%CI)=15.20 (12.01;19.23)	
3	curve from start of product use	N/A	CC (mean, 95%CI)=20.13 (17.72;22.88)	\leftrightarrow
5	to time of last quantifiable	14/71		(Positive)
_	concentration (ng.h/mL)			
NCT0196771 9	Maximal nicotine concentration	N/A	mIQOS (mean, 95%CI)=7.39 (5.68;9.62) CC (mean, 95%CI)=13.02 (10.06;16.85)	\
	(ng/mL)			(Negative)
91	Area under the concentration		mIQOS (mean, 95%CI)=16.56 (12.46;22.01)	
	curve from start of product use	N/A	CC (mean, 95%CI)=29.47 (21.35;40.67)	\leftrightarrow
ž	to time of last quantifiable			(Positive)
	concentration (ng.h/mL)	L		

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	Human Puffing/Smoking Topography (inc. puff count)	During single- use session on Day 5	Group 1 (mean, ±SD) Puff number: IQOS=15.0 (±5.5), CC=17.3(±5.9) Total puff volume: IQOS=730.9mL (±350.4), CC=682.8mL (±224.7) Mean puff volume: IQOS=48.8mL (±17.9), CC=41.3mL (±12.7) Puff duration: IQOS=1.4s (±0.5), CC=1.5s (±0.5) Puff interval: IQOS=17.1s (±20.7), CC=18.8s (±10.6) Session length: IQOS=195.6s (±87.2), CC=289.5s (±85.7) Group 3 (mean, ±SD) Puff number: Glo=15.4 (±7.4), CC=16.0 (±5.6) Total puff volume: Glo=731.3mL (±437.6), CC=596.8mL (±197.1) Mean puff volume: Glo=46.6mL (±16.8), CC=39.3mL (±12.4) Puff duration: Glo=1.6s (±0.5), CC=1.6s (±0.5) Puff interval: Glo=11.1s (±5.8), CC=18.8s (±10.6) Session length: Glo=150.4s (±40.5), CC=269.3s (±88.0)	↓ (Positive) ↑ (Negative) ↑ (Negative) ↓[IQOS] ↔ [Glo] (Positive) ↓ (Negative) ↓ (Positive)
	Daily product consumption	Ambulatory average	IQOS (mean, ±SD)=8.5 (±5.2) Glo (mean, ±SD)=7.0 (±5.5) CC (mean, ±SD)=13.2 (±4.4) [Group 1], 12.6 (±4.7) [Group 3]	↓ (Positive)
s, 2020	Mouth level exposure to NFDPM (mg/session)	During single- use session on Day 5	IQOS (mean, ±SD)=9.6 (±5.0) Glo (mean, ±SD)=4.7 (±2.9) CC (mean, ±SD)=19.0 (±7.7) [Group 1], 16.7 (±7.6) [Group 3]	↓ (Positive)
Jones,	Mouth level exposure to nicotine (mg/session)	During single- use session on Day 5	IQOS (mean, ±SD)=0.98 (±0.51) Glo (mean, ±SD)=0.34 (±0.21) CC (mean, ±SD)=1.55 (±0.63) [Group 1], 1.36 (±0.62) [Group 3]	↓ (Negative)
	Sensory questionnaire (magnitude scale [1-7], 'just right' scale [Low, Just right, High])	During single- use session on Day 5	Group 1 (mean (±SD) magnitude score, just right score) Immediate smoke/aerosol delivery: IQOS=3.7 (± 1.7), Low; CC=5.4 (± 1.3), Just right Draw effort: IQOS=4.1 (± 1.7), High; CC=3.5 (± 1.7), High Mouthful: IQOS=3.8 (± 1.3), Low; CC=4.8 (± 1.0), Just right Irritation: IQOS=3.4 (± 2.0), Just right; CC=2.9 (± 1.8), Just right Intensity of kick/hit: IQOS=3.6 (± 1.7), Just right; CC=3.4 (± 1.8), Just right Taste - likeability: IQOS=3.4 (± 2.0); CC=5.2 (± 1.3) Taste - amount: IQOS=4.2 (± 1.4), Just right; CC=5.0 (± 1.2), Just right Overall likeability: IQOS=3.6 (± 1.9); CC=5.3 (± 1.2) Group 3 (mean (±SD) magnitude score, just right score) Immediate smoke/aerosol delivery: Glo=3.3 (± 1.6), Low; CC=5.0 (± 1.3), Just right Draw effort: Glo=4.9 (± 1.6), High; CC=3.8 (± 1.5), High Mouthful: Glo=3.2 (± 1.3), Low; CC=4.5 (± 1.2), Just right Irritation: Glo=3.6 (± 1.9), Just right; CC=3.3 (± 1.4), Just right	↓ (Negative) ↑ (Negative) ↓ (Negative) ↑↔ (Unclear) ↑↔ (Positive) ↓ (Negative) ↓ (Negative) ↓ (Unclear) ↓ (Unclear) ↓

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			Intensity of kick/hit: Glo=3.9 (± 1.8), Just right; CC=3.8 (± 1.3), Just right Taste - likeability: Glo=2.8 (± 2.0); CC=5.1 (± 1.6) Taste - amount: Glo=4.0 (± 1.8), Just right; CC=4.6 (± 1.3), Just right	(Negative)
			Overall likeability: Glo=3.1 (± 1.9); CC=5.2 (± 1.4)	
	Human Puffing/Smoking Topography (inc. puff count)	During single- use session on day 5	Group 1 (mean, \pm SD) Total puff volume (mL): IQOS=668.1 \pm 322.6, Glo=736.4 \pm 415.8, CC=489.0 \pm 177.7 Mean puff volume (mL): IQOS=63.5 \pm 20.3, Glo=66.7 \pm 23.7, CC=48.9 \pm 14.8 Puff number: IQOS=10.3. \pm 3.6, Glo=10.9 \pm 5.6, CC=10.7 \pm 5.0 Puff duration (s): IQOS=1.8 \pm 0.6, Glo=1.8 \pm 0.6, CC=1.8 \pm 0.6 Puff interval (s): IQOS=8.3 \pm 3.0, Glo=7.4 \pm 2.7, CC=9.7 \pm 3.4 Group 2 (mean, \pm SD) Total puff volume (mL): mGlo=618.2 \pm 389.6, mCC=493.7 \pm 192.4 Mean puff volume (mL): mGlo=62.2 \pm 32.8, mCC=51.1 \pm 16.0 Puff number: mGlo=10.0 \pm 4.5, mCC=10.0 \pm 3.7 Puff duration (s): mGlo=1.8 \pm 0.5, mCC=2.0 \pm 0.5 Puff interval (s): mGlo=8.1 \pm 3.0, mCC=9.9 \pm 3.4	↑ (Negative) ↑ (Negative) ↓[IQOS] ↑[Glo] ↔[mGlo] (Unclear) ↓[mGlo]↔[IQOS/Glo] (Positive) ↓ (Negative)
e, 2018	Daily product consumption	Ambulatory average	IQOS (mean, ±SD)=12.2 ± 6.2 Glo (mean, ±SD)=10.3 ± 5.5 CC (mean, ±SD)=16.0 ± 8.1 mGlo (mean, ±SD)=11.4 ± 5.7 mCC (mean, ±SD)=15.3 ± 6.9	↓ (Positive)
Gee,	Mouth level exposure to NFDPM (mg/stick)	During single- use session on day 5	IQOS (mean, ±SD)=8.4 ± 4.5 Glo (mean, ±SD)=5.2 ± 3.4 CC (mean, ±SD)=13.5 ± 6.2 mGlo (mean, ±SD)=6.2 ± 3.8 mCC (mean, ±SD)=14.8 ± 7.4	↓ (Positive)
	Mouth level exposure to menthol (mg/stick)	During single- use session on day 5	mGlo (mean, \pm SD)=1.4 \pm 0.8 mCC (mean, \pm SD)=1.2 \pm 0.5	↑ (Unclear)
	Mouth level exposure to nicotine (mg/stick)	During single- use session on day 5	IQOS (mean, ±SD)=1.0 ± 0.5 Glo (mean, ±SD)=0.3 ± 0.2 CC (mean, ±SD)=1.3 ± 0.5 mGlo (mean, ±SD)=0.3 ± 0.2 mCC (mean, ±SD)=1.3 ± 0.6	↓ (Negative)
	Mouth insertion depth	Post product use	No comparison to cigarette arm	N/A

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	Pulse wave velocity (m/s)	Acute=post single use Chronic=1 month	Acute: IQOS (mean, ±SD)=10.2 ± 1.7; CC (mean, ±SD)=10.8 ± 2.4 Chronic: IQOS (mean, ±SD)=10.1 ± 1.5; CC (mean, ±SD)=10.2 ± 2.3	↓ (Positive)
NCT03452124	Exhaled Carbon monoxide (ppm)	Acute=post single use Chronic=1 month	Acute: IQOS (mean, ±SD)=14.1±7.3; CC (mean, ±SD)=17.5±7.8 Chronic: IQOS (mean, ±SD)=6.7±6.4; CC (mean, ±SD)=17.4±4.8	↓ (Positive)
NCT0	Perfused boundary region of sublinqual arterial microvessels	N/A	Not reported	N/A
	Global longitudinal strain of left ventricle (%)	1 month	Chronic: IQOS (mean, ±SD)=-20.9±2.5; CC (mean, ±SD)=-20±0.7 GLS was improved in the HNBC compared to the control group at follow-up (diference=2.35%; 95% CI 0.23-4.48, p=0.03)	↑ (Positive)
	Coronary flow reserve (no units)	1 month	Chronic: IQOS (mean, ±SD)=3.5±0.8; CC (mean, ±SD)=2.6±0.2	(Positive)
	Catalase (UI/cm2)	Post exposure to 32 puffs of product	Glo (mean, SD)=12.87, 7.77 CC (mean, SD)=10.01, 3.63	↑ (Positive)
	Malondialdehyde (ng/cm2)	Post exposure to 32 puffs of product	Glo (mean, SD)=46.10, 6.46 CC (mean, SD)=62.80, 12.02	↓ (Positive)
.022	Squalene (μg/cm2)	Post exposure to 32 puffs of product	Glo (mean, SD)=36.97, 24.29 CC (mean, SD)=34.95, 22.54	↑ (Positive)
Dalrymple, 2022	Squalene monohydroperoxide (ng/cm2)	Post exposure to 32 puffs of product	Glo (mean, SD)=73.80, 49.34 CC (mean, SD)=159.45, 67.26	↓ (Positive)
Dalr	Squalene monohydroperoxide/Squalene ratio (ng/μg)	Post exposure to 32 puffs of product	Glo (mean, SD)=2.07, 0.65 CC (mean, SD)=5.19, 1.38	↓ (Positive)
	L* (lightness) (no units)	Post exposure to 32 puffs of product	Glo (mean, SD)=69.30, 3.56 CC (mean, SD)=66.79, 2.57	↑ (Positive)
	a* (green-red) (no units)	Post exposure to 32 puffs of product	Glo (mean, SD)=7.32, 1.88 CC (mean, SD)=8.23, 0.95	↓ (Positive)

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	b* (blue-yellow) (no units)	Post exposure to 32 puffs of product	Glo (mean, SD)=15.72, 2.72 CC (mean, SD)=20.72, 1.91	↓ (Positive)
	Total difference in colour from control (ΔΕ) (no units)	Post exposure to 32 puffs of product	Glo (mean, SD)=2.61, 1.14 CC (mean, SD)=5.39, 1.54	↓ (Positive)
Ioakeimidis, 2021	Augmentation index (%)	30 mins post use	"There were no differences in all baseline measurements between the three sessions." "Compared with sham smoking, cfPWV (Figure 1(b)), baPWV and AIx@75 (Figure 1(c)) increased immediately after the end of tobacco cigarette smoking (by 0.29 m/s, 93 cm/s and 3.3%, respectively) and remained increased after 5 min. Likewise, HNBC smoking induced a significant increase in cfPWV, baPWV and AIx@75 (by 0.30 m/s, 86 cm/s and 3.5%, respectively)."	↓ (Positive)
	Heart rate (bpm)	Post use	"There were no differences in all baseline measurements between the three sessions." "HR increased similarly in both the tobacco cigarette and HNBC sessions (maximum increase by 10 beats/min)"	↔ (Negative)
	Brachial systolic blood pressure (mmHg)	30 mins post use	"There were no differences in all baseline measurements between the three sessions." "Both brachial (Figure 1(a)) and aortic systolic BP increased immediately after the end of smoking by tobacco cigarette (by 11.5 and 10.5 mmHg, p < 0.001 and p < 0.01, respectively) and by HNBC (by 7.5 and 6 mmHg, all p < 0.01)"	↓ (Positive)
	Aortic systolic blood pressure (mmHg)	30 mins post use	"There were no differences in all baseline measurements between the three sessions." "Both brachial (Figure 1(a)) and aortic systolic BP increased immediately after the end of smoking by tobacco cigarette (by 11.5 and 10.5 mmHg, p < 0.001 and p < 0.01, respectively) and by HNBC (by 7.5 and 6 mmHg, all p < 0.01)"	↓ (Positive)
	Carotid–femoral pulse wave velocity (m/s)	30 mins post use	"There were no differences in all baseline measurements between the three sessions." "Compared with sham smoking, cfPWV (Figure 1(b)), baPWV and AIx@75 (Figure 1(c)) increased immediately after the end of tobacco cigarette smoking (by 0.29 m/s, 93 cm/s and 3.3%, respectively) and remained increased after 5 min. Likewise, HNBC smoking induced a significant increase in cfPWV, baPWV and AIx@75 (by 0.30 m/s, 86 cm/s and 3.5%, respectively)."	↓ (Positive)
	Brachial-ankle pulse wave velocity (cm/s)	Post use	"There were no differences in all baseline measurements between the three sessions." "Compared with sham smoking, cfPWV (Figure 1(b)), baPWV and AIx@75 (Figure 1(c)) increased immediately after the end of tobacco cigarette smoking (by 0.29 m/s, 93 cm/s and 3.3%, respectively) and remained increased after 5 min. Likewise, HNBC smoking induced a significant increase in cfPWV, baPWV and AIx@75 (by 0.30 m/s, 86 cm/s and 3.5%, respectively)."	↓ (Positive)

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A wave velocity (cm/s)	10 minutes	IQOS [mean, (SD)]=55.8 (14.2), n=27	+
	post-use	CC [mean, (SD)]=57.9 (15.5), n=27	(Positive)
Diastolic blood pressure	10 minutes	IQOS [mean, (SD)]=71.9 (10.1), n=27	\downarrow
(mmHg)	post-use	CC [mean, (SD)]=75.5 (10), n=27	(Positive)
E wave velocity (cm/s)	10 minutes	IQOS [mean, (SD)]=66.8 (12), n=27	\
	post-use	CC [mean, (SD)]=67.3 (14.1), n=27	(Negative)
E/A ratio (no units)	10 minutes	IQOS [mean, (SD)]=1.2 (0.3), n=27	\leftrightarrow
	post-use	CC [mean, (SD)]=1.2 (0.4), n=27	(Negative)
Em/Am ratio (no units)	10 minutes	IQOS [mean, (SD)]=1.2 (0.5), n=27	\downarrow
	post-use	CC [mean, (SD)]=1.3 (1.0), n=27	(Negative)
Heart water (boson)	10 minutes	IQOS [mean, (SD)]=1.8 (8.7), n=27	<u></u>
Heart rate (bpm)	post-use	CC [mean, (SD)]=82.6 (8.8), n=27	(Positive)
I C	10 minutes	IQOS [mean, (SD)]=38.8 (4.8), n=27	↑
Left atrium diameter (mm)	post-use	CC [mean, (SD)]=38.3 (5.2), n=27	(Negative)
Left ventricle ejection fraction	10 minutes	IQOS [mean, (SD)]=64.5 (3.8), n=27	
(%)	post-use	CC [mean, (SD)]=64.4 (3.9), n=27	(Positive)
Left ventricle global	10 minutes	IQOS [mean, (SD)]=18.3 (3.9), n=27	↑
circumferential strain (%)	post-use	CC [mean, (SD)]=17.5 (3.9), n=27	(Positive)
Left ventricle global	10 minutes	IQOS [mean, (SD)]=17.9 (2.4), n=27	\leftrightarrow
longitudinal strain (%)	post-use	CC [mean, (SD)]=17.9 (2.8), n=27	(Negative)
Left ventricular end-diastolic	10 minutes	IQOS [mean, (SD)]=46.1 (4.1), n=27	<u> </u>
diameter (mm)	post-use	CC [mean, (SD)]=46.3 (4.5), n=27	(Positive)
Peak early diastolic velocity of	10 minutes	IQOS [mean, (SD)]=11.6 (3.6), n=27	↑
the left ventricle (cm/s)	post-use	CC [mean, (SD)]=10.7 (3.8), n=27	(Positive)
Peak late diastolic velocity of	10 minutes	IQOS [mean, (SD)]=9.5 (2.2), n=27	\
the left ventricle (cm/s)	post-use	CC [mean, (SD)]=10 (2.9), n=27	(Positive)
Right atrium diameter (mm)	10 minutes	IQOS [mean, (SD)]=38.2 (4.0), n=27	\
	post-use	CC [mean, (SD)]=38.3 (3.9), n=27	(Positive)
Right ventricle diameter (mm)	10 minutes	IQOS [mean, (SD)]=34.2 (3.2), n=27	\leftrightarrow
	post-use	CC [mean, (SD)]=34.2 (3.3), n=27	(Negative)
Right ventricle free wall strain	10 minutes	IQOS [mean, (SD)]=23.9 (6.2), n=27	<u> </u>
(%)	post-use	CC [mean, (SD)]=21.2 (5.6), n=27	(Positive)
Right ventricle global	10 minutes	IQOS [mean, (SD)]=21.4 (4.1), n=27	↑
longitudinal strain (%)	post-use	CC [mean, (SD)]=19.4 (4.1), n=27	(Positive)
Right ventricle peak early	10 minutes	IQOS [mean, (SD)]=10.7 (2.4), n=27	<u> </u>
diastolic velocity (cm/s)	post-use	CC [mean, (SD)]=10.5 (2.4), n=27	(Positive)

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	Right ventricle peak late	10 minutes	IQOS [mean, (SD)]=15 (4.5), n=27	
	diastolic velocity (cm/s)	post-use	CC [mean, (SD)]=14.5 (3.4), n=27	(Negative)
	Right ventricle systolic	10 minutes	IQOS [mean, (SD)]= 13.1 (2.1), n=27	↑
	myocardial velocity (cm/s)	post-use	CC [mean, (SD)]=12.8 (2.5), n=27	(Negative)
	units) Systolic blood pressure (mmHg)	10 minutes	IQOS [mean, (SD)]= 0.7 (0.2), n=27	\leftrightarrow
		post-use	CC [mean, (SD)]=0.7 (0.2), n=27	(Negative)
		10 minutes	IQOS [mean, (SD)]=114.1 (16.8), n=27	\downarrow
		post-use	CC [mean, (SD)]=120.5 (12.7), n=27	(Positive)
		10 minutes	IQOS [mean, (SD)]=9.8 (2.4), n=27	\uparrow
	the left ventricle (cm/s)	post-use	CC [mean, (SD)]=9.1 (2.3), n=27	(Negative)
	Tricuspid annular plane systolic	10 minutes	IQOS [mean, (SD)]=20.9 (2.5), n=27	\uparrow
	excursion (mm)	post-use	CC [mean, (SD)]=20.2 (2.9), n=27	(Positive)
	Human Puffing/Smoking	During single-	IQOS (median, IQR)=14.0, 13.5-14.0	\uparrow
	Topography (inc. puff count)	use	CC (median, IQR)=13.0, 10.8-16.3	(Negative)
			IQOS (median, IQR)=8.3, 4.5-19.3	
	Maximal nicotine concentration	N/A	CC (median, IQR)=12.9, 7.2-28.6	\downarrow
			Mean maximal nicotine concentration also lower in IQOS group than CC group	(Negative)
			based on graph (Figure 1)	
21	Nicotine	30 minutes	"IQOS delivered about half as much nicotine over 30 minutes (AUC0->30) as a	\downarrow
	Tricotine	50 minutes	cigarette"	(Negative)
Ċ.	Time to reach nicotine Cmax	N/A	IQOS (median, IQR)=4.0, 4.0-6.0	\downarrow
			CC (median, IQR)=6.0, 4.0-8.0	(Positive)
Phillips-Waller, 2021	Urge To Smoke questionnaire	Post product use	"OBC reduced urges to smoke more than IQOS"	↑
				(Negative)
	Area under the concentration curve from start of product use to 60 minutes	N/A	IQOS (median, IQR)=152.0, 91.2-254.5	
			CC (median, IQR)=314, 136.4-465.6	\downarrow
		1,712	"IQOS delivered about half as much nicotine over 30 minutes (AUC0->30) as a	(Negative)
			cigarette"	1
	Nicotine boost effect score	N/A	IQOS (median, IQR)=5.4, 2.6-10.8	↓
		ļ	CC (median, IQR)=12.7, 6.7-26.8	(Negative)
	Questionnaire (Other)	Post product	No comparison to cigarette arm	NE
	(,	use	I 0	NE

^{*} \uparrow = higher in HTP arm; \leftrightarrow = equivocal; \downarrow = lower in HTP arm

Abbreviations: Positive=HTP has positive impact compared to CC; Negative=HTP has negative impact compared to CC; N/A=not applicable; HTP=heated tobacco product; CHTP=carbon HTP; CC=combustible cigarette; [P]NTV=[prototype] novel tobacco vapor; LLTV=loose leaf tobacco vaporiser; create=creatinine; FAS-AR=Full analysis set – as randomised; FAS-EX=Full analysis set – as exposed; Cmax=maximal concentration; mean=arithmetic mean; geo mean=geometric mean