Regulation of Flavors in Tobacco Products: A Proposed Rule by the Food and Drug Administration
Docket No. FDA-2017-N-6565  83 FR 12294

Please find this comment on the Advanced Notice of Proposed Rule-Making for Regulation of Flavors in Tobacco Products1. We are grateful for the opportunity to comment at this point in the process.

At this point, we recommend that further consideration of FDA rule-making on flavors is confined to combustible tobacco products, albeit with due concern for unintended consequences. The challenges of rule-making and the public health test for flavored non-combustibles such as vaping products are far greater than for combustibles. This is because it is impossible to rule out significant harm-reduction benefits from use of non-combustible nicotine products by both adults and youth as alternatives to smoking. No such benefits exist with combustible products and the issues are therefore more straightforward.

If flavors increase the appeal of harm reduction transitions and pathways, then intervening to reduce flavor-related appeal may cause harm. We do not believe that, at this point, FDA can reliably distinguish between harms and benefits that arise from flavors in non-combustible nicotine products, or provide assurance that its interventions would not cause more harm than good. This applies to both adult and youth populations. We see no ethical or legal rationale to exclude harm reduction benefits to young people from policy analysis simply because we would prefer them not to use nicotine products at all.

Further, we urge FDA to consider the importance of the appeal of non-combustible products as part of its wider nicotine strategy and its expressed intent to move nicotine users away from combustible products. Without the availability of appealing legally-available non-combustible products like e-cigarettes, there is a danger that consumers will seek the nicotine products they want in the black or gray market or start making their own2.

In support of this position, we make the following observations.

1. The policy issues arising with rule-making on flavored combustible products are very different to those related to non-combustible products. With combustibles, if it can be established that flavored

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1 U.S. Food & Drug Administration, Regulation of Flavors in Tobacco Products: A Proposed Rule by the Food and Drug Administration, 83 FR 12294, 21 March 2018 [link]

2 See Comment by Attorney General Miller (Iowa) and 17 others: Tobacco Product Standard for Nicotine Level of Combusted Cigarettes, 11 July 2018 [link] Tracking number: 1k2-947r-gewy
products cause a material increase in consumption or facilitate initiation, then there is a case for rule-making to reduce harm. In that event, the case against rule-making to ban flavored combustible products would rest on whether a ban would be unacceptably discriminatory in some way, whether rule making would work and create the intended result or users would mostly switch to unflavored tobacco products, or whether it would cause unintended consequences, including illicit trade and other forms of countervailing response. There is a plausible route to rule-making for flavored combustibles, albeit with some significant evidential challenges. We have no further comment on rule-making for flavored combustible products.

2. Providing an evidential basis for rule-making on flavored non-combustible products is a far more formidable challenge. This is because, in some situations, use of these products is net beneficial to health. No such complication applies to combustible products. When flavored non-combustible products contribute to smoking cessation, when they substitute for smoking or reduce smoking to low levels, and when they prevent initiation on combustible products there is a potential health benefit from harm reduction that should be acknowledged. These effects can apply to both adults and to youth. The harm reduction benefits arise because non-combustible products are much less risky than smoking products.

3. When non-combustible products substitute completely or mostly for combustible products, there will be a health benefit, and, based on what is known about such products, the reduction in individual risk is likely to be substantial (one to two orders of magnitude). However, if they lead to nicotine use in people who would never have otherwise have used nicotine, the risk to health should be assumed non-trivial (even if not yet identified), but likely to be quite low in absolute terms. Long term health impacts remain the subject of speculation and no material adverse health effects have so far been detected at a significant scale, and projections based on greatly reduced toxicity suggest much lower risk.

4. There is only a serious risk from uptake of non-combustible products, if: (1) vaping causes nicotine use that would not have otherwise happened; and (2) this leads to smoking via a ‘gateway effect’; (3) vaping does not allow a subsequent reversal of the smoking back to vaping before significant damage accumulates. There is so far no compelling evidence of a gateway effect, and some


6 Tobacco Advisory Group of the Royal College of Physicians (London), Nicotine without smoke: tobacco harm reduction. 28 April 2016 [link]


evidence that vaping products may be an ‘exit’ from smoking\textsuperscript{9,10}. Though many gateway claims have been made by motivated academics, all suffer from the inherent and unresolvable weakness of uncorrected residual confounding for common liabilities and other methodological challenges that have yet to be overcome\textsuperscript{11}. These fundamental problems have rendered all gateway claims to date unreliable.

5. Establishing the balance of benefits and harms to give a net positive or negative impact will be exceedingly difficult. However, unless some surprising new evidence emerges, the balance is unlikely to come out in favor of net harm. For young people, this is because regular vaping is highly concentrated in smokers\textsuperscript{12} and young people who vape report tobacco harm reduction motivations for vaping (i.e. to reduce harm from smoking)\textsuperscript{13,14}. Furthermore, while there has been a rapid rise in youth vaping there has also been a rapid decline in smoking, consistent with vaping displacing smoking at population level. Although such observations alone cannot prove a causal link, they cannot rule it out either. Given the rapid reduction in both youth\textsuperscript{15} and adult\textsuperscript{16} smoking over the period in which vaping has risen, FDA would need a high standard of contrary evidence to base policy on the assumption that vaping is a net cause of harm to young people or to adults or that there is no causal relationship between the decline of cigarette use and rise of e-cigarettes.

6. Even if FDA could establish that vaping was a cause of harm to either adults or to youth (and as discussed above, we do not believe FDA can do this) it would require a further evidential challenge to attribute the harmful effects to one or more flavors or flavor descriptors. FDA would need to rely on an implicit claim that there is a material additional uptake of vaping that is caused by particular flavors or flavor descriptors. At one level, this is a banal claim: if the products tasted disgusting or tasted of nothing, then few would use them. It follows, that, in total, flavors are integral to the viability of vaping products. It is not, therefore, a surprise that young people report use of e-


\textsuperscript{14} Shiffman S. Sembower MA. PATH Data: Harm Reduction is Teens’ Top Reason for Using e-cigarettes, Poster SRNT, Florence March 2017 [link]


\textsuperscript{16} National Center for Health Statistics, National Health Interview Survey, Early releases [link]. Figure 8.1. Prevalence of current cigarette smoking among adults aged 18 and over: United States, 2006–2017.
cigarettes “because they come in flavors I like”\textsuperscript{17}. It would be more concerning if young people vaped but \textit{did not like} the flavors, suggesting they were in a strong grip of dependency. But given that flavors are integral to vaping products, a ban on all non-tobacco flavors would amount to a \textit{de facto} ban on the product itself, except in its tobacco-flavored form. There is no logic to banning most vaping products and no logic to offering an exemption for tobacco flavors, which are still artificial and added to nicotine liquids. It would be particularly misguided to leave tobacco flavors as the only choice because many vapers are trying to escape the tobacco experience altogether, and non-tobacco flavors are becoming more prominent as the initial product that smokers initially try and then ultimately migrate to\textsuperscript{18}. Broadly based rule-making banning most or all flavors in non-combustibles (other than tobacco) would run counter to FDA’s broader nicotine strategy, in which satisfactory alternatives to cigarettes are a necessary component of actions to reduce nicotine in cigarettes.

7. If a ban on all flavors cannot be justified, the question then is whether there is a subset of flavors or flavor descriptors that are problematic and could be the subject of rule-making. It is difficult to identify what these flavors are and whether they are a \textit{cause} of uptake of vaping. It cannot be assumed that childish branding or tradenames make a product ‘kiddie-appealing’. FDA’s policy should not be made on the basis of naive assessments of the motivations of adolescents, many of whom are trying to graduate to adulthood or reject child-like innocence. It is perfectly possible that other features of products drive youth use, for example, styling, convenience, nicotine delivery, and whether it has become fashionable. Or it may be that flavors with more edgy or humorous names capture the attention of young people. Though FDA has conducted high profile enforcement against what it considers to be child-orientated packaging\textsuperscript{19} it has not provided any evidence that these products are in fact disproportionately attractive to adolescents or stand out from the thousands of available flavors as successful inducements to vaping.

8. Even if a particular flavor is found to be appealing to young people, it requires a further step in the chain of reasoning to establish that this is a \textit{cause} of changed behavior (i.e. it caused the uptake of vaping) or whether it just reflects a \textit{preference} among those who vape and who would vape anyway for other reasons. FDA has not shown how it will establish causal links between particular flavors or flavor descriptors and net additional vaping, and it is a non-trivial problem to demonstrate this relationship. A further issue is to determine how to specify a flavor or flavor descriptor in regulations in a way that is legally precise and the effect of the measure clear – and not vulnerable to easy circumvention. The danger would be of overly broad classifications that unlawfully prohibit products that are not part of the problem, as defined by FDA, of youth appeal. Should a rule concentrate on descriptors or somehow characterizing flavor sensations?


\textsuperscript{18} Russell C, McKeganey N, Dickson T, Nides M. Changing patterns of first e-cigarette flavor used and current flavors used by 20,836 adult frequent e-cigarette users in the USA. \textit{Harm Reduct J.} BioMed Central; 2018 Jun 28;15(1):33. \[link\]

\textsuperscript{19} U.S. Food & Drug Administration. FDA, FTC take action against companies misleading kids with e-liquids that resemble children’s juice boxes, candies and cookies. Press release. 1 May 2018 \[link\]
9. The final challenge for FDA is to show that whatever intervention it makes will actually have the intended effect of reducing harm and not trigger harmful unintended consequences. Unintended consequences could include users: making their own flavored liquids; buying from overseas internet suppliers; using other flavors in the same way; relapsing to smoking; continuing to smoke instead of switching; and adopting different risk-taking behaviors such as vaping cannabinoids. FDA has no intervention research we are aware of, though it could carefully study the effects of the flavor ban in San Francisco once it has been in place long enough to allow for observable comparisons.

10. FDA has begun to argue that harm reduction benefits should not be recognized for nicotine users under the age of 18, asserting that these users should not be using nicotine at all. According to Commissioner Gottlieb:

   *Even if kids are using ENDS instead of cigarettes -- and that migration in part accounts for the decline in youth cigarette use -- that's still not an acceptable trade.*

We do not agree that FDA should or can ignore harm reduction benefits to young people who would otherwise smoke. Nothing in the public health test for FDA rulemaking exempts the health of young people from consideration. We recognize FDA's aspiration that young people should not use nicotine in any form and we respect the sentiment. However, a responsible regulator can only proceed on the basis that young people do engage in risky behaviors and do use nicotine, often in its most dangerous form, smoking, whatever our preferences. Early risk-taking with any tobacco or nicotine product, such as an e-cigarette, may result from social or emotional rewards from trying a product, including peer approval or mood enhancement. Thus, eliminating all experimentation may not be a realistic goal, just as it has not been for cigarettes. This is, objectively, the case in reality – 19.6 percent of high school age students and 5.6 percent of middle school students use tobacco/nicotine products in some form. It is, therefore, a legitimate responsibility of regulators to mitigate harms that arise to these 3.1 million at-risk adolescent nicotine users, of which 2.3 million are using combustible tobacco products. FDA should not intervene in a way that obstructs harm reduction or increases actual harm to young people simply because FDA starts with a prior but unrealistic belief they should behave differently to how they do in reality, a point made forcefully and convincingly by Professor Lynn Kozlowski at a recent conference.

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21 Gottlieb S. Commissioner of Food & Drugs, FDA’s Nicotine and Tobacco Regulation and the Key Role of Regulatory Science (speech), 18 June 2018 [link]

22 Tobacco Control Act, Section 907(a)(3) [link]


11. To summarize, the chain of reasoning required to justify rule-making to prohibit particular flavors, flavor categories or flavor descriptors in non-combustible products is extremely challenging, with the real possibility that FDA intervention could cause harm both to adults and young people if it make misjudgments about the (1) effects of vaping on health, and (2) the effect of flavors on vaping. FDA would need to show that vaping itself is a source of net harm (this is unlikely) and show that particular flavors or descriptors were increasing uptake and contributing to harm (this is difficult). Finally, it would need to show its proposed intervention would be proportionate and effective, and not prone to excessive unintended consequences (for this there is no credible evidence). The FDA does not have a reliable case at any point in this chain of reasoning.

In December 2017, General Miller raised many of these methodological issues with FDA Commissioner Gottlieb in advance of publication of the ANPRM so that they could be considered in detail in the published ANPRM. We do not believe the concerns raised have been addressed in the ANPRM. We have included the letter to Dr. Gottlieb and accompanying briefing as Appendix 1 and 2 respectively. We request that these communications are considered as part of this comment.

We hope that these views contribute to appropriate and effective FDA policy-making in the field of nicotine and tobacco.

Yours faithfully,

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Appendix 1: Letter to Dr. Gottlieb from AG Miller, 20 December, 2017

Dear Dr. Gottlieb

Re: a framework for the considering the appeal of flavors in nicotine products

Following FDA’s announcement of a comprehensive nicotine strategy on 28 July, we write regarding one aspect of the strategy: the possible rule-making with respect to flavored nicotine products.

We are concerned that some prominent commentary about flavors in low-risk tobacco and nicotine products, including that provided in the Surgeon General’s 2016 report, is overly simplistic in presuming that the primary purpose and consequence of offering flavors is to recruit current non-users, especially youth, to nicotine use. A proper assessment requires a deeper analysis, and must at least consider the possibility that these low-risk products can function as alternatives to combustibles, not only for adults, but also as a diversion from youth smoking uptake and as support for youth smoking cessation. In that case, the attractiveness and appeal of such non-combustibles may be a positive factor in reducing the use of the far more harmful products, such as cigarettes. This harm-reduction benefit may apply for both adult and adolescent users.

Any justification for an intervention must show that a rule is appropriate for the protection of public health and that it is reasonable to conclude that harms will outweigh benefits. To show this is likely would require a long chain of reasoning, supported by credible data. Is a flavor attractive? Is it differentially attractive to youth, versus adults? If it is differentially attractive, does it change behavior? If it changes behavior, is the change harmful or beneficial? How would an intervention affect behavior? Would an intervention reduce harm or reduce benefits in youth and adult populations? What are the potential mechanisms for unintended consequences?

The attached memo discusses the analysis that is necessary to show that harms (or benefits) arise from flavored nicotine products. We hope that the ANPRM on flavors will reflect these issues at the point of publication. This will provide a more realistic foundation for public comment and help to raise the level of public and political debate about this important and controversial issue.

Yours sincerely

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Appendix 2: Briefing provided to Dr. Gottlieb, 20 December 2017

Assessing potential harms and benefits arising from flavors in nicotine products

Following the 28 July announcement of FDA’s new nicotine strategy, publication of several Advanced Notices of Proposed Rulemaking (ANPRMs) is expected within months. These will provide a more detailed articulation of the strategy. This memo discusses significant issues for consideration in the drafting of the forthcoming ANPRM on the role of flavored tobacco/nicotine products in attracting youth.

In articulating its strategy, FDA should be mindful of the real-world complexity governing potential harms and benefits of flavors. To that end, this memo sets out a possible framework for assessing harm and benefits as a series of questions below.

1. **Is the flavor used in a combustible or non-combustible product?**

Very different considerations are required depending on whether or not there is combustion. Flavored non-combustible products offer a ‘harm reduction’ pathway to smokers (or users who would otherwise smoke), and the appeal of such products may thereby create a benefit. No such benefit applies in the case of combustibles – and a completely different approach is required to analyze public health impacts and to define appropriate policy. Combustible and non-combustible flavored products should never be lumped together in policy considerations, given the pronounced variation in risk and the opportunity to substantially reduce health risks to people who would otherwise smoke.

The rest of this memo concerns flavors in non-combustible tobacco/nicotine products.

2. **Is the cause for concern an actual flavor or the way it is described (or both)?**

The first step is to identify the actual cause for concern and hence the possible subject of rule-making. There are three potential options for regulation of flavors: (1) any properties of the flavor chemical that are harmful to health; (2) characteristics of the flavor itself (i.e. its sensory properties) that make the product inherently more palatable or appealing to younger people; (3) characteristics of the flavor descriptor (i.e. descriptive words or imagery – including trademarks or brand values from other products). We believe the first of these should beyond the scope of this ANPRM, but could managed through technical standards as required. For concerns about youth appeal, any rule-making initiative in this area requires a clear system for defining which flavor characteristics or descriptors are subject to regulation, and how a rule will define what is in scope and what is out of scope.

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3  Are all flavors, whole flavor categories or specific flavors the cause for concern?

At one level, it is obvious that flavors play a role in all use of vaping products. If the products taste bad or are flavorless then few people will use them. But flavors are integral to vaping products – all products are flavored in some way, even if with a tobacco flavor. Supposedly ‘unflavored’ cigarettes are not flavorless, but taste of the thousands of chemicals in tobacco smoke. Every orally consumed tobacco/nicotine product is flavored in some way. Eliminating an essential component of vaping products (the flavor) would amount to a de facto prohibition of vaping products. Such action would run counter to FDA’s new nicotine strategy, which stresses the importance of the availability of low-risk nicotine products as alternatives to combustible cigarettes. It follows that the question is how to identify a subset of flavors, with well-described selection criteria, that present concerns above and beyond simply making vaping products viable.

4  How will the subset of flavors that have a particular role in attracting youth be identified?

Commissioner Gottlieb stated: “I have real concerns about kids’ use of e-cigarettes […] especially those products marketed with obviously kid-appealing flavors”. But what constitutes an obviously kid-appealing flavor? It may be obvious what constitutes a flavor with childlike branding, but it is not obvious that such childlike flavors appeal to the adolescent population at risk. It is just as likely that adolescents are concerned with reinforcing their adult identity and will prefer flavors or branding reflecting adult values.

One option to identify youth-attracting flavors would be to focus on those that have the greatest proportion of sales to younger people. However, unless preferences are uniform across all age-bands, then there will always be a category that has higher youth uptake. How pronounced should the bias towards youth sales be before the flavor or category becomes a matter of concern? It is likely that adults would use more tobacco flavor, as most adult vapers will be current or former smokers. To account for this, should any assessment of youth-adult biases in flavor preferences be assessed net of tobacco flavor use? How disproportional to adult appeal/use must youth appeal/use be for a flavor to be considered a concern?

Much advocacy focusses on trademark names, such as Gummy Bear or Cotton Candy. Just because these flavors can be found on the internet somewhere, it does not mean they have a noticeable effect on population behavior. How should appeal of such flavors be characterized?

5  Does a flavor preference create a change in behavior to increase e-cigarette use?

While there might be flavors that are more or less attractive to youth, it requires an additional step to show that these flavors exert such a powerful attraction that they cause additional use of a product where there would otherwise be no use. At least one experiment suggests that flavors exert negligible

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3 U.S. Food and Drug Administration, Remarks by Scott Gottlieb, M.D. Commissioner of Food and Drug Administration: Protecting American Families: Comprehensive Approach to Nicotine and Tobacco, July 28, 2017. [link]
attraction on non-users. In this study, when teenage subjects were asked to rate their interest in using e-cigarettes on a scale of 0-10 when offered in a list of flavors, they reported minimal interest, reaching an average interest score of only 0.41 out of 10.

It is possible for a person’s decision to try vaping to be made for different reasons (to bond with friends, to try something other than smoking) and the choice of flavor is a secondary or lower consideration—much as the decision to go out for a meal with friends may not be caused by desire for a particular item on the menu. For example, Ambrose et al. offer a widely cited analysis of PATH survey data in which young people who already vape are asked their reasons for product use. This includes the option: “(It) comes in flavors I like,” for each tobacco/nicotine product. But an affirmative answer to that question will have been a trivially obvious response for many—who would use a product with a flavor they did not like? The question does not identify specific flavors of concern, so it is referring to an integral feature of the product, without which the product would have no appeal. Yet this study is frequently cited as justification for intervening to restrict flavors to protect youth.

6 What is the behavior of concern and what is a distraction?

If flavors are playing a role in changing behavior, it is important to be clear what defines a harmful risk behavior worthy of significant intervention and possible trade-offs with other objectives such as adult smoking cessation. Data suggests much adolescent e-cigarette use is experimental and occasional and, as such, poses minimal risk. The National Youth Tobacco Survey for 2014 showed that 74 percent of high school students who were using e-cigarettes used them on less than ten days in the month preceding the survey, 45 percent on only 1-2 days, with less than 10 percent being daily users. Furthermore, the more regular e-cigarette use is strongly concentrated in smokers. As Ken Warner puts it: Non-smoking high school students are highly unlikely to use e-cigarettes; among those who do, most used them only on 1–2 of the past 30 days. The headline prevalence figures for adolescent e-cigarette use are based on any use in the past 30 days. As a result, the headline figures conceal and merge very different behaviors: regular use and experimentation. Regular e-cigarette users are mostly smokers and already at risk.

A further complication is that many adolescents vapers report mostly using e-cigarettes without nicotine and therefore without the key dependence-forming agent. The Monitoring the Future Survey found

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4 Shiffman S, Sembower MA, Pillitteri JL, Gerlach KK, Gitchell JG. The impact of flavor descriptors on nonsmoking teens’ and adult smokers’ interest in electronic cigarettes. *Nicotine Tob Res* 2015; published online Jan 7 [link].


6 Campaign for Tobacco Free Kids and others, The Flavor Trap: How Tobacco Companies Are Luring Kids with Candy-Flavored E-Cigarettes and Cigars, 15 March 2017 [link]


that only 20 percent of 12th graders reported using e-cigarettes with nicotine\textsuperscript{10}. Again, the youth e-cigarette statistics that drive public and political concern are ‘contaminated’ by the inclusion of very different behaviors.

When gauging the scale of potentially problem vaping behavior, we should focus on regular or daily use of nicotine-based products – a small subset of the total. Even among this group, it cannot be assumed that the vaping behavior is problematic – it may be or become an alternative to smoking. This is the subject of the following section.

7 Would youth uptake of e-cigarettes caused by flavors be harmful or beneficial to health?

If it is assumed that: (1) it is possible to identify flavors that are attractive to youth and (2) to show that these flavors change behavior – increase regular nicotine use or cause initiation - it is then necessary to establish (3) whether the change in behavior is harmful or beneficial. What if the behavior change prompted by an appealing flavor was to divert a young person from smoking to vaping? Given that regular vaping is concentrated in smokers, this is a distinct possibility.

A reanalysis\textsuperscript{11} of the PATH data considered by Ambrose et al\textsuperscript{5}, showed that harm-reduction (to self and others) motivation was the most important reported reason for using e-cigarettes, cited by 88% of the young people surveyed. Moreover, almost all the youth who cited the availability of flavors as a motive for e-cigarettes use also cited harm reduction as a motive. The authors remind us that “Teens commonly endorse multiple reasons for using e-cigarettes, rendering the analysis of motives complex”. It is quite possible to conclude from this data that palatable or even enjoyable e-cigarette flavors assist with realizing the primary motivations to reduce harm or quit smoking. In other words, the flavors contribute to a health benefit in youth.

This raises the challenge that harm-reduction motivations may apply to e-cigarette use by young people under 18 years and their use of e-cigarettes may be beneficial. There is also the plausible hypothesis that, whatever the motivation, teenage vaping has played a contributory role in the rapid decline in teenage smoking witnessed in the United States since 2010\textsuperscript{12}\textsuperscript{13}. In this case, enhanced appeal of the e-cigarette products may be supporting the displacement of cigarette initiation or consumption with e-cigarette use, which is a much lower risk behavior. Before flavors are denounced as increasing teenage e-cigarette use, it is important to have a sense of the counterfactual: what would have happened in the absence of e-cigarettes? Would young vapers simply have smoked? That e-cigarettes can substitute for

\textsuperscript{10} Miech R, Patrick ME, et al. What are kids vaping? Results from a national survey of US adolescents. Tob Control; 2016 [link].

\textsuperscript{11} Shiffman S., Sembower MA, PATH Data: Harm Reduction is Teens' Top Reason for Using e-cigarettes. Poster SRNT 2017, Pinney Associates [link].


smoking among youth is supported by convergent results of independent analyses showing that regulations limiting access to e-cigarettes increase youth smoking\textsuperscript{14, 15}.

8 How are trade-offs between potential harms and potential benefits to youth addressed?

It is possible that some youth use of e-cigarettes occurs among youth who would not otherwise have used tobacco or nicotine. In this case, there is a behavioral pathway leading to a potential net harm to health, if trial converts into persistent long-term use. It is also possible that some youth e-cigarette use will be an alternative to smoking, in which case there is a pathway leading to a health benefit. It is possible to construct a far more complex model than this two-pathway example\textsuperscript{16}. Leaving aside adults at this stage, the net health effect on youth will be a function of how many young people follow each of these pathways and what the net harm or benefit to them would be in each pathway. Kozlowski and Warner\textsuperscript{17} warn that fears that e-cigarettes will serve as a net ‘gateway’ to smoking are exaggerated and could undermine the much larger potential for discouraging smoking in the whole population.

Given that it is unlikely that vaping risk exceeds five percent of smoking\textsuperscript{18}, and given that regular e-cigarette use is concentrated in young people who smoke or have a high propensity to smoke\textsuperscript{19}, then it is highly unlikely that this calculation of trade-offs will resolve showing net harm, and it is much more likely it will show net benefit. Simulation modeling with sensitivity analyses that examine all the state and transition pathways shows that the gateway effect would have to be implausibly large to increase the net public health harm\textsuperscript{20}.

9 How will beneficial impacts for adults be reconciled with any potential impacts on youth?

A 2017 assessment suggest that e-cigarettes are likely having a positive (i.e. downward) effect on adult smoking prevalence, which has been falling rapidly\textsuperscript{21}, via an increased smoking cessation rate\textsuperscript{22}.

\begin{thebibliography}{99}
\bibitem{Friedman} Friedman AS. How does Electronic Cigarette Access affect Adolescent Smoking? \textit{J Health Econ} Published Online First: October 2015. [link]
\bibitem{Pesko} Pesko MF, Hughes JM, Faisal FS. The influence of electronic cigarette age purchasing restrictions on adolescent tobacco and marijuana use. \textit{Prev Med (Baltim)}, February 2016 [link]
\bibitem{Kozlowski} Kozlowski LT, Warner KE. Adolescents and e-cigarettes: Objects of concern may appear larger than they are. \textit{Drug Alcohol Depend.} 2017 May;174(1 May 2017):209–14. [link][PDF]
\bibitem{Royal} For an expert approximation see: Royal College of Physicians (London), \textit{Nicotine without smoke: tobacco harm reduction}. 28 April 2016 [link] "Although it is not possible to precisely quantify the long-term health risks associated with e-cigarettes, the available data suggest that they are unlikely to exceed 5% of those associated with smoked tobacco products, and may well be substantially lower than this figure". (Section 5.5 page 87)
\bibitem{Data} National Center for Health Statistics, National Health Interview Survey [link], Sample Adult Core component. Figure 8.1. Prevalence of current cigarette smoking among adults aged 18 and over: United States, 1997–2016. [data]
\end{thebibliography}
Conclusion The substantial increase in e-cigarette use among US adult smokers was associated with a statistically significant increase in the smoking cessation rate at the population level. These findings need to be weighed carefully in regulatory policy making regarding e-cigarettes and in planning tobacco control interventions.

Further, we know that adults make extensive use of non-tobacco flavors, including fruit and candy, even though these may be considered childish, or even ‘kid-appealing’. One study found 68 percent of American adult e-cigarette users had used non-tobacco flavors in the past 30 days. Of these, 45 percent had used fruit, 44 per cent menthol or mint, and 26 per cent candy, chocolate or other sweet flavor.\(^{23}\)

We argue above that there are unlikely to be net harms to youth arising from e-cigarette use. However, if it were somehow shown there are net harms to youth, then how should these be traded off against potential substantial gains in the adult population? Again this would require assessment of behavior-change pathways induced by the flavored product, in adults as well as youth. Some evidence\(^{24}\) suggests that the availability of non-tobacco flavors helps some adult smokers transition completely away from smoking and to the much safer practice vaping. As above, it is likely that benefits to people already smoking, or at high risk of smoking, would greatly outweigh risk from additional uptake of vaping.

10 What impact would a rulemaking intervention by FDA have?

In this complex landscape of multiple behavioral pathways how will it be possible to assess unintended harmful consequences of a policy designed to reduce the appeal of e-cigarettes? In fact, that question should be applied to several FDA interventions, including the Real Cost campaign, which will now target e-cigarettes\(^{25}\) or the regulatory burdens created by the deeming rule\(^{26}\).

It is unclear how FDA could design interventions that only address (minor) harms without compromising the likely (substantial) benefits. It would first need to know the disposition of harms and benefits attributable to flavors (as discussed in 2-9 above). Then it would need to assess how a flavor-related intervention would modify the behavior, and the patterns of harms and benefits. Then it would need to be confident that its intervention would reduce harm rather than increasing it. This is exactly the sort of assessment, analysis, and modeling that FDA demands of companies applying to market new tobacco products or make modified-risk claims for products. We have seen little to suggest FDA or the advocacy community urging action on flavors is remotely close to being able to make that assessment.

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\(^{25}\) Food and Drug Administration, FDA to expand public education campaign to focus on prevention of youth e-cigarette use, news release, 8 August 2017 [link]

\(^{26}\) For a description of unintended consequence and limited benefits of the deeming rule, see: Brief of amici curiae of Clive Bates and fifteen other in support of plaintiffs’ motion for summary judgement, 5 August 2016 [link].
Regrettably, the 2016 U.S. Surgeon General’s report on youth and e-cigarettes\textsuperscript{27} did not engage with the complexities set out above, and therefore cannot provide helpful scientific orientation to policymakers.