Title
Modelling the impact of a new tobacco product: Review of Philip Morris International's Population Health Impact Model as applied to the IQOS heated tobacco product

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ABSTRACT

Objectives: We review the Population Health Impact Model (PHIM) developed by Philip Morris International (PMI) and used in its application to the US Food and Drug Administration (FDA) to market its heated tobacco product (HTP), IQOS, as a modified risk tobacco product (MRTP). We assess the model against FDA guidelines for MRTP applications and consider more general criteria for evaluating reduced risk tobacco products.

Methods: In assessing the PHIM against FDA guidelines, we consider two key components of the model: the assumptions implicit in the model (outcomes included, relative harm of the new product vs. cigarettes, tobacco-related diseases considered, whether dual or poly-use of the new product is modeled, and what other tobacco products are included) and data used to estimate and validate model parameters (transition rates between nonsmoking, cigarette only smoking, dual use of cigarettes and MRTP, and MRTP only use; and starting tobacco use prevalence).

Results: The PHIM is a dynamic state transition model which models the impact of cigarette and MRTP use on mortality from 4 tobacco-attributable diseases. The PHIM excludes morbidity, underestimates mortality, excludes tobacco products other than cigarettes, does not include FDA-recommended impacts on non-users, and underestimates the impact on other population groups.

Conclusion: The PHIM underestimates the health impact of HTP products and cannot be used to justify a MRTP claim. An assessment of the impact of a potential MRTP on population health should include a comprehensive measure of health impacts, consideration of all groups impacted, and documented and justifiable assumptions regarding model parameters.
WHAT THIS PAPER ADDS:

- Heated tobacco products (HTP), also referred to as heat-not-burn products, are not currently marketed in the United States and their impact on the health of the US population is not known.
- Philip Morris International developed a Population Health Impact Model that they used to estimate the potential impact of marketing a HTP product, IQOS, as a modified risk tobacco product (MRTP) in the US.
- Because the model is used to support a MRTP application, FDA guidelines indicate that it should include the impact of the new product on morbidity and mortality, and the impact on 7 population groups and exposure patterns. However, the model underestimates mortality, omits morbidity measures, excludes impacts on nonusers, and underestimates the impact on other groups. Therefore, the model underestimates the potential impact of IQOS on the population as a whole and does not justify marketing the product as a modified risk tobacco product.
- An assessment of the impact of a new tobacco product on population health should include a comprehensive measure of health impacts, consideration of all groups impacted, and documented and justifiable assumptions regarding model parameters such as the relative harm of the new product compared to existing products and transition rates between tobacco use categories.
INTRODUCTION

Philip Morris International (PMI) submitted an application to the Food and Drug Administration (FDA) to market its heated tobacco product (HTP) IQOS as a modified risk tobacco product (MRTP) in the US, arguing that because the product does not actually burn tobacco, it will have a reduced impact on health compared to cigarettes. PMI used a computational model they developed, the Population Health Impact Model (PHIM),\(^1\) to estimate the potential impact of this IQOS marketing on public health. While the application was denied by the FDA, the proliferation of purported reduced harm products suggests the need for an understanding of how to assess the impact on population health of new tobacco products.

No models specifically consider the health impact of IQOS, but several simulation models analyze the impact of two tobacco products on population health. These models evaluate the impact of a reduced risk tobacco product on population health by comparing a factual scenario (considering cigarette use only) with a counterfactual scenario, in which the new product is introduced. None of the models consider the impact of other tobacco products. Details of the models, the assumptions they are based on, and their findings are summarized in Supplement 1 and Supplement Table 1. Four models compared the health effects of cigarettes with e-cigarettes (or a vaporized nicotine product), measuring health effects either as an index\(^2\) or as mortality.\(^3\) Two of these models reported a net positive impact on health\(^3\)\(^4\)\(^6\) while two reported net population harm.\(^2\)\(^5\) All four research teams assumed that e-cigarettes were safer than cigarettes by factors ranging from 5% to 30%, but they differed in their assumptions about the impact of e-cigarettes on cigarette smoking initiation and cessation. Three studies analyzed the impact of introducing a nonspecified MRTP on cigarette smoking and mortality. Each study reported a potential reduction in mortality,\(^7\)\(^8\) though one study indicated that mortality could
increase if the MRTP were 50% as risky as cigarettes and 50% of initiates were never smokers. One study evaluated the impact of promoting use of the smokeless product snus on a health index, and concluded that promoting snus as a safer product than cigarettes is not likely to result in population health benefits.

These models illustrate how different assumptions about what is included in the model as well as the data sources for estimating transition rates and tobacco use prevalence lead to varying conclusions about the net impact of a new product. These model characteristics will be reviewed for the PHIM.

PMI’s multiple tobacco product model, the PHIM, was refined for its application for IQOS. This paper reviews the FDA guidelines for MRTP applications and assesses whether the PHIM as used in the IQOS MRTP application meets the criteria the FDA has developed to determine whether or not the impact of IQOS on population health justifies the introduction of the product as a MRTP. We also consider more generally the criteria for assessing the impact of a new tobacco product on population health.

METHODS

We evaluate the PHIM as published and as submitted for marketing IQOS as a MRTP against FDA guidelines for MRTP applications. In our evaluation, we consider two key components of the model: the assumptions implicit in the model (outcomes included, relative harm of the new product vs. cigarettes, tobacco-related diseases considered, whether dual or poly-use of the new product is modeled, and what other tobacco products are included) and data used to estimate and validate model parameters (transition rates between nonsmoking, cigarette only smoking, dual use of cigarettes and MRTP, and MRTP only use; and starting tobacco use prevalence).
The FDA issued draft guidelines for MRTP applications in March 2012. The guidelines specify that “scientific studies submitted by the applicant “should contain an overall assessment of the potential effect that the marketing of the product as proposed may have on tobacco-related morbidity and mortality”. (page 21) The guidelines further recommend that the potential impact on mortality and morbidity be assessed for 7 population groups and exposure patterns. (page 22)

RESULTS

The PMI PHIM

The PHIM, developed by PMI researchers and their collaborators, is described briefly here and in more detail in Supplement 2. The PHIM is a dynamic state transition model which models the impact of cigarette and MRTP use on mortality. It follows a cohort aged 15 and older for 20 years. The PHIM consists of a prevalence component (“P-component”) and an epidemiological risk component (“E-component”). The P-component models changes in the distribution of cigarette and/or MRTP use occurring in a hypothetical population over a defined period. The model compared a null (i.e. no MRTP) scenario and an MRTP scenario. (p. 88) For each scenario, transition probabilities for initiation, re-initiation, and cessation of smoking and of product switching (including dual cigarette/MRTP use) are estimated from historical cigarette smoking prevalence data, and pre-market Perception and Behavioral Assessment studies conducted by PMI. The E-component uses the tobacco use patterns from the P-component along with estimates of the relative risk (RR) of death for lung cancer, ischemic heart disease (IHD), stroke, and chronic obstructive pulmonary disease (COPD) to estimate mortality using published estimates of RR for smoking and assumptions about how much less risky MRTP use is compared to smoking.
Sensitivity analyses vary assumptions about initiation and reinitiation of tobacco use; transition rates between smoking, MRTP, and dual use; time frames; and the RR of the MRTP vs. cigarettes.

**Comparison of IQOS MRTP application with FDA guidelines**

**Impact of IQOS on morbidity.** The PHIM does not include any measure of morbidity, such as incident or prevalent cases of tobacco-related illness. One way of quantifying the impact of morbidity is through healthcare costs, which incorporate the severity and time course of illness, and would include hospitalizations, outpatient care, medications, and other services. No estimates of healthcare costs are made in the PHIM.

**Impact of IQOS on mortality.** The PHIM considers mortality from four diseases caused by smoking – lung cancer, IHD, stroke, and COPD.

The base case in the IQOS MRTP application assumes that compared to cigarettes, sole MRTP use is 80% less risky and dual use of MRTP and cigarettes is 40% less risky than cigarette smoking alone. The RR of death for dual use of cigarettes and IQOS is assumed to be the midpoint of the risk of cigarette smoking and the risk of IQOS use.\(^1\) (page 19) To simulate the mortality impact on the US population, the model uses smoking prevalence from 1990 projected through 2010.

**Impact of IQOS on different types of individuals.** We next assess how the PHIM treats the 7 population groups and exposure patterns recommended for consideration by the FDA.\(^1\) (page 22) More detailed descriptions are contained in Supplement Table 2.

1) **Tobacco users who switch from other commercially marketed tobacco products to the proposed product.** The PHIM considers switching only from cigarettes. PMI acknowledges that
other tobacco products are not considered in their model, arguing that there is no evidence to indicate that IQOS users will switch from other tobacco products.\textsuperscript{12} (page 7)

2) Tobacco users and non-users who, after adopting the proposed product, switch to or switch back to other tobacco products that may present higher levels of individual health risk. The PHIM assumes that each month 0.1\% of IQOS users will switch to cigarette smoking, but that after a year of IQOS use virtually no users will become cigarette smokers or dual users. They also assume that 10\% of dual IQOS/cigarette smokers will become sole cigarette smokers each month.\textsuperscript{12} (page 14, Table 4)

3) Tobacco users who opt to use the proposed product rather than cease tobacco use altogether. PMI indicates that this group was “considered by a specific analysis in which current conventional cigarette smokers who would otherwise have switched to MRTP or to dual use, quit instead”.\textsuperscript{13} (Module 7.4, page 5) PMI indicates that “here, the reduction in deaths associated with MRTP introduction was estimated to be about 11 times greater in males and about 13 times greater in females than that for the basic analysis”.\textsuperscript{13} (Module 7.4, page 5)

4) Tobacco users who opt to use the proposed product rather than an FDA-approved tobacco cessation medication. PMI indicates that this is “outside the present scope of the model”.\textsuperscript{13} (Module 7.4, page 5)

5) Non-users who initiate tobacco use with the proposed product, such as youth, never users, former users. The PHIM assumes that uptake of the IQOS HTP will be limited among youth because of the relatively high cost. It assumes that the percent of never-smokers who will initiate tobacco use with IQOS each month ranges from .05\% to .08\% (after 25 years), and that the rate drops with age, with no one initiating use after age 35.\textsuperscript{12} (Module 6.5, page 13, Table 3) The model assumes that reinitiation rates of former smokers with IQOS range from .01\% for
youth aged 15-19 to .08% of older adults (aged 75-79) after more than 25 years.\textsuperscript{12} (Module 6.5, page 13, Table 3)

6) \textit{Tobacco users who use the product in conjunction with other tobacco products.} The PHIM assumes that few smokers or IQOS users will become dual users\textsuperscript{12} (Module 6.5, page 14, Table 4) and that fewer than .02% of never tobacco users and fewer than .04% of former smokers will become dual users.\textsuperscript{12} (Module 6.5, page 13, Table 3)

7) \textit{Non-users who experience health risks from the product.} Risk to non-users is not considered in the PHIM.

\textbf{DISCUSSION}

The PHIM is similar in structure to many of the published models reviewed, which are all dynamic in nature and model state transitions in tobacco use over time, with the exception of one steady state model.\textsuperscript{2} The PHIM focuses on mortality as the outcome measure as do all but 2 models which included a health effects index.\textsuperscript{2,10} The PHIM models the population aged 15 and older, an improvement over some of the models which focus on a subgroup of the population. It follows the population for 20 years, which is reasonable for MRTP application purposes, and is line with the published models, which use varying time horizons from 10 to 84 years.

However, the PHIM analysis of IQOS has some important limitations that are apparent in reviewing the model against FDA recommendations. Morbidity-related outcomes are omitted, mortality is underestimated, transition rates used in the model are based on PMI perception studies, and the model uses data for the US in 1990 as a starting point. The role of other tobacco products such as e-cigarettes as well as impact on non-users is not considered. Thus, the analysis of IQOS does not fully satisfy FDA guidelines for MRTP applications, and results in an overestimation of the benefit of IQOS on population health.
Morbidity is ignored
The PHIM does not include any measures of morbidity, such as tobacco-related disease incidence or tobacco-attributable healthcare costs, though this is an FDA requirement. Morbidity costs are more than half of total costs of cigarette smoking for high income countries,\textsuperscript{15} so this omission is potentially serious.

Mortality is underestimated
The clinical results presented for US adults to justify the lower RR of mortality for IQOS vs. cigarette use do not show statistically significant improvements in the biomarkers of harm that PMI assessed in actual people who used HTP (with a single exception).\textsuperscript{16} This contradicts the assertion of reduced harm, and does not justify the 70\% to 90\% reductions in risk assumed in the PHIM. The RR of mortality for IQOS compared to cigarettes is a critical parameter in the model and a smaller reduction in harm should be used in the analyses.

The RRs of mortality from smoking used in the PHIM are based on multi-country studies rather than those published by the 2014 US Surgeon General and based on US cohorts.\textsuperscript{17} PMI’s sensitivity analyses indicate that the proportion of smoking-attributable deaths from the 4 causes for men would increase 15\% (2005-09) if based on the RRs from the Surgeon General report,\textsuperscript{11} with less of a change for women. The Surgeon General estimates, which are more current and vetted through a more thorough process of independent peer review than the PHIM estimates, are more appropriate and should be used in these analyses.

The PHIM assumes that the RR of dual use of IQOS and cigarettes is the midpoint of the 2 RRs. However, there is some evidence that dual users of cigarettes and e-cigarettes have greater risks of negative health outcomes than sole cigarette users,\textsuperscript{18} suggesting that there could
be greater risks for dual users of IQOS and cigarettes and that the PHIM model may underestimate the number of deaths attributable to dual use.

The inclusion of only 4 smoking-attributable diseases in the PHIM further reduces the estimates of mortality from IQOS vs. cigarette use. At least 22 causes of death for adults\textsuperscript{19} and 4 causes of death for infants\textsuperscript{20} have been causally linked to cigarette smoking. PMI acknowledges that the “overall estimates of deaths saved due to the introduction of IQOS would have to be increased about 50% to give an estimate for all smoking-related diseases combined”.\textsuperscript{12} (p. 41)

Given that mortality is the main measure of population health used in the PHIM, the use of low RRs and the inclusion of only 4 causes of death will result in an overestimate of the benefit of IQOS introduction as an MRTP which will greatly impact the results.

**Assumptions about transition rates are not well justified**

The PHIM uses transition probabilities for smoking dating back to 1986 for 12 countries. PMI does report an adjustment for poor model forecast performance through 2005 but does not report the methodology or provide documentation of the model predictions against historical data. Other models use more recent data, document the methodology, and report the predictive performance against historical data. For example, Warner and Mendez\textsuperscript{5} validate their model to US data through 2015 and Levy et al.\textsuperscript{3} calibrate their model using US data through 2010. The absence of an explanation of methodology and documentation for the PHIM predictive performance is a serious weakness because poor forecasts of status quo and alternative scenarios may bias the results.

Transition probabilities for IQOS initiation, reinitiation, cessation, and product switching are based on PMI perception surveys. The only empirical data available are from Italy, but these data report on ever use and are thus not comparable to the PHIM estimates.\textsuperscript{21} Youth have
initiated tobacco use with e-cigarettes at high rates,\textsuperscript{22} and may find the IQOS product to be similarly appealing. Flavors, electronic features, and perceptions of harm are factors that are important determinants of adolescent decisions regarding tobacco use, and IQOS is likely to appeal to them on all these characteristics.\textsuperscript{23} The PHIM assumption that youth uptake will be limited because of the relatively high cost ignores the use of coupons to reduce prices, a common tobacco industry pricing strategy, and also ignores shared use among users. Recent estimates suggest that the prevalence of sharing e-cigarette devices among adolescents over the previous 30 days exceeds 70%.\textsuperscript{24}

The PHIM makes optimistic assumptions about cigarette smoking cessation rates associated with IQOS use, assuming that .4–1.5% of smokers will quit smoking each month due to IQOS use,\textsuperscript{12} (Module 6.5, Table 3) a relatively high rate in light of evidence that many IQOS users continue to smoke cigarettes, including PMI’s own finding that 36% of Japanese IQOS users use another tobacco product.\textsuperscript{25} Recent research has produced evidence for the US that, with the current regulatory environment and smoking behaviors, e-cigarettes do not increase smoking cessation in the general population greater than what would have occurred without them.\textsuperscript{26} Furthermore, the potential effectiveness of e-cigarettes in aiding smoking cessation may depend greatly on the level of the smoker’s nicotine dependence.\textsuperscript{27} This is also likely to impact the effectiveness of IQOS in cessation, but is not acknowledged in the PHIM.

The potential gateway effect of IQOS, is not fully considered. There is evidence for youth and young adults that e-cigarette use increases subsequent uptake of cigarette smoking.\textsuperscript{28} PMI indicates in its application that IQOS mimics cigarette smoking better than e-cigarettes or vaping because of more rapid nicotine delivery, suggesting that IQOS may be much more effective at addicting youth and young adults to nicotine as well as increasing transitions to
cigarette smoking. A net increase in nicotine addiction and cigarette uptake among adolescents and young adults is a realistic possibility that the PHIM does not consider.

Transition rates are one of the key parameters in the model and their correct estimation is critical to the results.

The model uses the 1990 US population and smoking prevalence as the starting point for the simulations

The PHIM simulates the health impact on the population starting with a baseline population and smoking prevalence representative of the US in 1990.\textsuperscript{12} (Module 6.5.2.2) It is not clear why 1990 data was used, when smoking prevalence was much greater than in more recent years; data for 2015 were readily available at the time of the analyses. Other published models use more recent prevalence data from 2000,\textsuperscript{8} 2006,\textsuperscript{10} 2011,\textsuperscript{6} and 2016.\textsuperscript{3} The use of 1990 prevalence is likely to lead to higher than actual smoking-attributable costs and higher expected benefits from IQOS.

The PHIM ignores other tobacco products, such as e-cigarettes.

The population health results would be different if the PHIM comparison were between IQOS and a lower risk product such as e-cigarettes. There are reasons to expect that e-cigarette users may find IQOS to be a tempting and attractive product, and ignoring the role of e-cigarette use in a model of the population health impact of IQOS will lead to an incomplete analysis.

The PHIM assumes very low rates of transition to dual use, contrary to empirical evidence from other countries showing that many of those individuals who use IQOS will continue to use their previous product. In Japan, where IQOS products are now available, over one-third of IQOS users are poly-users, most of whom also smoke cigarettes.\textsuperscript{25} Dual use of electronic tobacco products (HTP products including IQOS, Glo, and Ploom Tech, or non-
nicotine e-cigarettes) and combustible cigarettes was reported by 3.4% of Japanese internet
survey respondents in 2017. Thus, actual evidence of dual IQOS and cigarette use indicates
that the assumptions of dual use rates in the PHIM are too low.

**Impact of IQOS on non-users is not considered**

Ignoring the impact on non-users who experience health risks from IQOS is not reasonable.
Empirical evidence already exists for secondhand exposure from HTP aerosol. A Greek study
found that nicotine levels for IQOS aerosol were greater than those in e-cigarettes at low puff
duration, though lower than tobacco cigarettes. Another study using an animal model that
exposed rats to cigarette smoke and IQOS aerosol at levels that were relevant to real world
human exposure levels found that both exposures resulted in similar vascular impairment.

There is also direct evidence of negative health impacts from exposing human nonusers to HTP
aerosol. In Japan, 49% of never-tobacco users and 41% of former tobacco users exposed to
secondhand HTP aerosol reported symptoms including general illness, eye discomfort, or a sore
throat.

Children are particularly likely to be impacted by exposure to HTP products. They may
suffer negative health effects when exposed to their parents’ secondhand aerosol, as they are
when exposed to secondhand cigarette smoke. A Canadian study found that children suffered
respiratory effects from exposure and digestive effects of ingestion of e-cigarettes. Women who
use IQOS while pregnant may cause lifelong health impacts for their children, as is the case for
women who smoke cigarettes or use snuff while pregnant. Another potential risk from
IQOS use is fires and explosions, such as those that occur with e-cigarettes. Ignoring the health
impact of IQOS on nonusers overestimates the benefit of IQOS as a MRTP. While this impact
may be of a smaller magnitude than the impact on users of IQOS or cigarettes, the impact on non-users is recommended by the FDA for consideration.

**Conclusion**

The PHIM has a structure not unlike other simulation models reviewed. However, because it is used to justify the marketing of a tobacco product as a MRTP, it must satisfy FDA guidelines that other models are not subject to. The FDA is likely to receive a number of applications for MRTP orders in the coming years, and it is important that reasonable criteria be established for reviewing them. Future analyses of the impact of new tobacco products used for social decision making such as regulatory actions should consider all relevant and substantial social effects. This includes both morbidity and mortality that arise from a comprehensive list of tobacco-attributable diseases. Model-based estimates need to carefully document methods for estimating key parameters such as transition rates and to validate model’s predictive performance. Also, the effects of policy on all populations that will be affected should be included in the analyses, including non-tobacco users who will suffer health effects. These recommendations are relevant for the evaluation of new tobacco products as well as potential harm reduction products more generally.

PMI, through its analysis of IQOS using the PHIM, has not shown that this product would “significantly reduce harm and the risk of tobacco-related disease to individual tobacco users; and benefit the health of the population as a whole taking into account both users of tobacco products and persons who do not currently use tobacco products”. 14 (page 3) As new tobacco products are introduced into US and worldwide markets, particularly those that purport to be less harmful than currently used products, models of population health impacts will play an
important role. The PMI PHIM as applied to the marketing of IQOS as a MRTP illustrates some of the potential pitfalls of analysis that should be avoided.
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COMPETING INTERESTS

None of the authors have any conflicts to declare.

CONTRIBUTORSHIP

Dr. Max drafted the paper and Dr. Lightwood took the lead on statistical review in major revisions. All authors contributed to the analyses and reviewed and edited the manuscript.


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