Continuous-time system identification of a smoking cessation intervention

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Accepted author version posted online: 17 Dec 2013. Published online: 05 Feb 2014.

To cite this article: Kevin P. Timms, Daniel E. Rivera, Linda M. Collins & Megan E. Piper, International Journal of Control (2014): Continuous-time system identification of a smoking cessation intervention, International Journal of Control, DOI: 10.1080/00207179.2013.874080

To link to this article: http://dx.doi.org/10.1080/00207179.2013.874080

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Continuous-time system identification of a smoking cessation intervention

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(Received 28 February 2013; accepted 6 December 2013)

Cigarette smoking is a major global public health issue and the leading cause of preventable death in the United States. Toward a goal of designing better smoking cessation treatments, system identification techniques are applied to intervention data to describe smoking cessation as a process of behaviour change. System identification problems that draw from two modelling paradigms in quantitative psychology (statistical mediation and self-regulation) are considered, consisting of a series of continuous-time estimation problems. A continuous-time dynamic modelling approach is employed to describe the response of craving and smoking rates during a quit attempt, as captured in data from a smoking cessation clinical trial. The use of continuous-time models provide benefits of parsimony, ease of interpretation, and the opportunity to work with uneven or missing data.

Keywords: continuous-time identification; behavioural science; statistical mediation; smoking cessation; self-regulation

1. Introduction

Cigarette smoking is a major global public health issue. Approximately 10 million annual global deaths are expected to result from smoking by 2020 (Fish & Bartholomew, 2007), and the global smoking population is expected to surpass 1.7 billion by 2025 (Erhardt, 2009). In the United States, cigarette smoking is the leading cause of premature death and $157B in economic loss is attributed to tobacco use annually (Centers for Disease Control and Prevention, 2010; Killeen, 2011). Forty years of decreases in smoking rates have recently stalled in the United States, where one in five adults is an active smoker (Centers for Disease Control and Prevention, 2012). This smoking rate persists despite the fact that nearly 70% of smokers have expressed a desire to quit (Centers for Disease Control and Prevention, 2011). Largely due to the chronic, relapsing nature of cigarette smoking, over 90% of unaided attempts to quit smoking fail (Fiore et al., 2008).

Interventions play an important role in smoking prevention and cessation. Generally speaking, interventions for behavioural health disorders seek to reduce unhealthy behaviours and promote healthy ones through prevention or treatment, and are used to address many public health concerns in addition to smoking such as other substance abuse, obesity, sexually transmitted infections, and cancer screening, and can be pharmacological or behavioural in nature (Baker et al., 2011; Collins, 2012). Traditionally, these interventions are ‘fixed’, meaning they are not systematically operationalised, and the composition and dosage of an intervention component is given to all individuals receiving the intervention and do not vary over time (Collins, Murphy, & Bierman, 2004). The effectiveness of existing fixed smoking cessation interventions is limited. Counseling alone, for example, has a reported success rate below 15% (Fiore et al., 2008; Fish & Bartholomew, 2007). Pharmacological interventions (e.g., nicotine replacement therapies such as Nicorette®) have individual one-year abstinence rates below 35% (Fish & Bartholomew, 2007), which may be lower at longer term follow-up (Irvin & Brandon, 2000; Irvin, Hendricks, & Brandon, 2003). Such low success rates are particularly troubling given the gravity of cigarette smoking as a public health issue.

To address the limitations of fixed interventions, recent efforts in behavioural health have centred around development of so-called ‘adaptive’ interventions, where treatment components and dosage vary according to participant response (Collins et al., 2004). These interventions consist of closed-loop dynamical systems and may be more effective behavioural health interventions as they essentially seek to optimally adapt to the changing needs of a patient (Nandola & Rivera, 2013; Rivera, Pew, & Collins, 2007). Control systems engineering principles offer an appealing
framework for developing algorithms that implement these optimised, time-varying smoking cessation interventions. However, the impact of using control engineering concepts in the design of time-varying smoking cessation interventions is tied to the reliability of the smoking behaviour change models upon which the algorithms are based (Nandola & Rivera, 2013; Riley et al., 2011; Rivera, 2012).

The development of reliable models has been greatly enhanced by increased access to intensive longitudinal data (ILD). ILD in behavioural settings is loosely defined as quantitative or qualitative measurements recorded frequently over time and is more readily available due to increasing use of mobile and computerised technologies in behavioural trials (Walls & Schafer, 2006). ILD facilitates the dynamic modelling of behaviour and generally provides a means for improved analysis of inter- and intraindividual variability (Collins, 2006). In the context of cigarette smoking, ILD offers an opportunity to study the dynamics of smoking behaviour change (e.g., daily smoking rate, average craving level) during a quit attempt. Whereas traditional quantitative modelling methodologies from behavioural science (e.g., structural equation models, SEMs) are static in nature (Bollen, 1989), dynamical systems modelling and system identification offer a framework for more comprehensive characterisation of dynamic behavioural relationships and how smoking cessation interventions affect these dynamics (Timms, Rivera, Collins, & Piper, 2012). Recently, similar models have been used for improved evaluation of gestational weight gain and fibromyalgia interventions; these models also offer an appealing basis for development of optimised, time-varying interventions that draw from control systems engineering principles (Deshpande, Nandola, Rivera, & Younger, 2011; Dong et al., 2013; Nandola & Rivera, 2013; Rivera, 2012). In terms of smoking cessation, an intervention based in controller design methods could feature feedback in order to systematically assign medication dosages based on patient reports of withdrawal symptoms, for example (Timms, Rivera, Collins, & Piper, 2013). Given the gravity of cigarette smoking as a public health issue and the modest effectiveness of even the most efficacious treatments available, an improved ability to inform and evaluate behavioural health interventions warrants development of dynamic models of smoking cessation.

In this article, smoking cessation is described as a process of behaviour change, and this process is represented via continuous-time models. A continuous-time approach is particularly appealing in behavioural health settings (Rivera, 2012; Timms et al., 2012). Notably, for low-order dynamics (which appears to be the case for many dynamic behaviours; Deshpande et al., 2011; Rivera, 2012; Timms et al., 2012; Timms, Rivera, Collins, & Piper, in press), continuous-time models estimated from discrete-time data result in parsimonious expressions through which important dynamic features are more easily discerned. Consequently, insight into behavioural phenomena and intervention effects may be more easily interpretable with continuous-time models. As will be demonstrated, continuous-time models that capture inverse response in craving during a quit attempt can be easily identified with a right half plane zero term, the estimated value of which may shed light on underlying dynamic phenomena (Timms et al., 2012, 2013). Similarly, the parameters in continuous-time models are more meaningful in terms of understanding a process, as such models more transparently describe the continuous nature of actual physical systems of interest. An a priori understanding of a system can also be more easily preserved with continuous-time models, whereas discrete-time models of a second-order system, for example, may introduce additional parameters due to sampling. Furthermore, discrete-time models estimated at fixed sampling rates may not be representative of process dynamics observed under different sampling rates (Garnier, Wang, & Young, 2008). This may be an important consideration, given the range of time scales of interest to behavioural scientists (e.g., short term, long term, non-standard time periods such as pubertal time scales; Collins et al., 2004). Additionally, missing data and irregular sampling intervals are characteristic of self-reported behavioural health experiments, whether intentional or not; this supports the appeal of a continuous-time modelling approach, which will not have parameters that are a function of the sampling time, and therefore inherently manage the issue of non-ideal data measurement. Consequently, continuous-time models of discrete data collected at a sufficiently fast sampling rate can produce models that accurately represent the overall dynamics of behavioural phenomena, while avoiding challenges associated with missing data and inconsistent sampling intervals in discrete-time modelling (Garnier et al., 2008).

Behavioural scientists often rely on behaviour change theories to guide intervention design, evaluation, and delivery (Riley et al., 2011). In the context of cigarette smoking, the concepts of statistical mediation and self-regulation have been of particular interest (McCarthy et al., 2008a; Piper et al., 2008; Timms et al., 2012, in press; Velicer, Redding, Richmond, Greeley, & Swift, 1992; Walls & Rivera, 2009). Statistical mediation is a modelling paradigm central to the social and behavioural sciences, describing a multivariate causal relationship in which an independent variable affects a mediator variable and an outcome variable, with the mediator also affecting the outcome (MacKinnon, 2008). Self-regulation theory within smoking considers a process in which nicotine levels, behavioural state, or emotional state set points are regulated by smoking activity (Solomon & Corbit, 1974; Timms et al., 2013, in press; Velicer et al., 1992; Walls & Rivera, 2009). However, the utility of models that describe these behaviour change theories for the purposes of development of optimised, time-varying interventions has been limited; this is largely a consequence of the static nature of traditional behavioural science models and the difficulties historically associated
with intensive collection of behavioural data (Riley et al., 2011). This article employs a modelling framework that leverages ILD and continuous-time system identification in order to describe smoking cessation as a mediational and self-regulatory process.

This article is organised as follows. First, a clinical trial of bupropion and counseling as aids to smoking cessation is outlined; this study was conducted at the University of Wisconsin Center for Tobacco Research and Intervention (UW-CTR) and funded by the Trans-disciplinary Tobacco Use Research Centers (TTURC; McCarthy et al., 2008b). Behavioural signals for treatment group averages and two single subject examples from the clinical trial are then presented. The features of the signals of interest in this article – Craving, average daily craving level, and Cigsmoked, total daily smoking – are discussed in general terms. Statistical mediation is first presented in conceptual terms and the connection to dynamic model development is then outlined. The iterative procedure used for model estimation is then described and the resulting empirical dynamic mediation models are discussed. Next, self-regulatory smoking cessation models are presented, estimated, and compared. In examination of both the mediation and self-regulation models, parameter estimates and model simulations are analysed for treatment group averages. Following this, examples of single subject models are briefly discussed. Finally, conclusions and recommendations are presented.

### 2. Smoking cessation overview

Dynamic models are obtained in a secondary analysis of a TTURC-funded study conducted by the UW-CTR. In this double-blinded, placebo-controlled, randomised clinical trial, 101 subjects received both active bupropion and counseling as treatment (the ‘AC’ group), 101 received active bupropion and no counseling (‘ANc’), 100 received a placebo and counseling (‘PC’), and 101 received a placebo and no counseling (‘PNc’). Participants receiving bupropion took 150 mg per day starting one week prior to the quit date and 300 mg per day from four days prior to quit to eight weeks post-quit. Bupropion SR (Zyban SR) is commonly prescribed as a smoking cessation treatment (Fiore et al., 2008; Fish & Bartholomew, 2007), although the exact mechanism that makes it an effective smoking treatment is debated (Horst & Preskorn, 1998; Warner & Shoaib, 2005). Generally, bupropion is thought to interfere with nicotine dependence mechanisms (Warner & Shoaib, 2005), and has been shown to alleviate withdrawal symptoms, including craving (McCarthy et al., 2008b). In lieu of active bupropion, the PC and PNc groups took placebo medication. Subjects receiving counseling completed two pre-quit counseling sessions, one quit-date session, and five sessions over the following four weeks post-quit. Sessions focused on preparation, coping, motivation, and relapse prevention. In lieu of counseling, the ANc and PNc groups spoke with staff about medication use adherence and received general encouragement (McCarthy et al., 2008b).

For the two weeks prior to and four weeks immediately following the target quit date, participants were instructed to complete assessments through personal digital assistants (PDAs) each day immediately after waking up, before going to bed, and four to seven times throughout the day as prompted by the PDA at pseudo-random intervals. These self-reports generally collected data on smoking frequency and withdrawal symptoms. Although data from each set of reports or a combination of reports could be used to model the smoking process, the Evening Report (ER) is the focus of this article’s efforts. The ER featured questions on a 10-point Likert scale covering topics such as withdrawal symptoms (McCarthy et al., 2008b), positive affect (generally, the degree to which an individual feels enthusiastic and alert), negative affect (generally, the degree to which an individual feels anger, disgust, guilt, fear, and nervousness; Watson, Clark, & Tellegen, 1988), and motivation to abtain. Table 1 provides a selection of items from the ER (McCarthy et al., 2008b). The relationship between Craving

<table>
<thead>
<tr>
<th>Code</th>
<th>Question</th>
<th>Scale</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urge</td>
<td>Since last ER on average – Bothered by urges?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Cigsmoked</td>
<td>Since last ER on average – Cigarettes on my mind?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Thinksmk</td>
<td>Since last ER on average – Thinking about smoking a lot?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Bother</td>
<td>Since last ER on average – Bothered by desire to smoke?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Enthus</td>
<td>Since last ER on average – Enthusiastic?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Tense</td>
<td>Since last ER on average – Tense or anxious?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Sad</td>
<td>Since last ER on average – Sad or depressed?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Anger</td>
<td>Since last ER on average – Bothered by anger/irritability?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Excellent</td>
<td>Since last ER on average – Concentration was excellent?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Happy</td>
<td>Since last ER on average – Happy and content?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Food</td>
<td>Since last ER on average – Thinking about food a lot?</td>
<td>1-11, No!!...Yes!!</td>
</tr>
<tr>
<td>Confidence</td>
<td>Since last ER on average – Confidence in ability to quit?</td>
<td>1-11, Low!!...High!!</td>
</tr>
<tr>
<td>Motive</td>
<td>Since last ER on average – Motivation to quit/stay quit?</td>
<td>1-11, Low!!...High!!</td>
</tr>
<tr>
<td>Cigsmoked</td>
<td>Total number of cigarettes smoked since the last ER?</td>
<td>0-99</td>
</tr>
</tbody>
</table>
and Cigsmked variables is the focus of this article, as was done in a statistical study of the same ILD by McCarthy et al. (2008a). Craving is defined as the sum of Urge, Cigsmind, Thinksmk, and Bother. Cigsmked is the total number of cigarettes smoked per day.

Both nomothetic (group level) and idiographic (single participant) models are of general interest to behavioural scientists. In this article, treatment group average models are the primary focus. To produce the group average signal, each report item was averaged across all members in a group for each relative day over the week prior to and four weeks immediately following the target quit date. This filtering that occurs by averaging the data across all single subject data points in a group, and putting the time series in deviation variable form, was the only data pre-processing done prior to group average model estimation. While the continuous-time approach employed here can effectively model data with missing samples or non-constant sampling, missing ER data for the two single subject examples was interpolated for straightforward use of standard MATLAB estimation routines. Interpolation consisted of averaging adjacent measured values or extending the adjacent measured value to the appropriate boundary. For the single subject example from the AC group, eight days of data points were imputed. Seven days of data points were imputed for the single subject example from the PNc group. Although single subject data sets are often noisier than corresponding group average data, no filtering was done on these data prior to idiographic model estimation. Figure 1 depicts the Craving and Cigsmked ILD for two group averages (solid blue, AC; dashed green, PNc) and two single subject examples (dash-dot magenta, AC; dotted red, PNc).

As seen in Figure 1, the group average Craving signals feature quit-induced inverse response. With a continuous-time modelling approach, a priori knowledge that the groups’ Craving signals features a right half plane zero is more easily preserved (Garnier et al., 2008). The group average Cigsmked signals feature a dramatic quit-day drop, followed by a relatively small and slow resumption of smoking. The single subject data sets display greater variability. In Figure 1, the PNc single subject does not feature a net reduction in craving. The AC subject has little resumption in smoking – reflecting a successful quit attempt – while the PNc subject features significant resumption to approximately pre-quit levels.

3. Statistical mediation modelling

The concept of statistical mediation is a prominent model of change in the social and behavioural sciences. As previously described, mediation defines a causal relationship in which an independent variable, $X$, affects a mediator, $M$, and an outcome, $Y$, with $M$ also contributing to $Y$ (MacKinnon, 2008). Behavioural scientists use path diagrams to depict this type of process (Bollen, 1989; MacKinnon, 2008). A mediational model path diagram – not to be confused with a block diagram – is depicted in Figure 2(a): $a$, $b$, and $c'$ represent gains for the $X$ to $M$, $M$ to $Y$, and $X$ to $Y$ pathways, respectively (MacKinnon, 2008). SEM representations of mediation are found in Equations (1) and (2):

$$M = \beta_0 + a X + e_1,$$

$$Y = \beta_0 + c' X + b M + e_2.$$  (1) (2)

Historically, Equations (1) and (2) have characterised mediation thought to be captured within cross-sectional studies; $X$ typically codes the presence or absence of an intervention and $M$ and $Y$ data are collected at a small number of time points (MacKinnon, 2008).

A dynamical framework is developed in this manuscript according to a more general definition of mediation described in Collins, Graham, and Flaherty (1998). Collins et al. (1998) underscores a temporal relationship between $X$, $M$, and $Y$, where a change in an independent variable at some time is said to result in lagged changes in the mediator and outcome (Collins et al., 1998). The SEMs in Equations (1) and (2) still apply under the Collins et al. (1998) definition, but the variables are a function of specific, discrete times.

Prior work by Navarro-Barrientos, Rivera, and Collins (2011) established how path diagrams in SEM correspond to steady-state process models; from these, fluid analogies can be constructed that lead to dynamical systems amenable to estimation via system identification methods. Drawing
from techniques used in production inventory management in supply chains, fluid analogies describe dynamic behaviours according to a structural relationship defined by a behavioural model (Schwartz, Wang, & Rivera, 2006). The fluid analogy in Figure 2(b) depicts a physical system analogous to behaviour change according to a mediational mechanism. Here, the independent variable corresponds to the exogenous input to the system, and the endogenous quantities in the path diagram (M and Y in Figure 2(a)) are represented as inventories. Dynamic, mediated behaviour change models are obtained when each inventory is considered in the context of a general conservation principle:

\[
\text{Accumulation} = \text{Sum of Inflows} - \text{Sum of Outflows.} \tag{3}
\]

A system of basic, first-order differential equations results from the application of Equation (3) to Figure 2(b):

\[
\tau_1 \frac{dM}{dt} = aX(t - \theta_1) - M(t) + e_1(t), \tag{4}
\]

\[
\tau_2 \frac{dY}{dt} = c'X(t - \theta_2) + bM(t - \theta_2) - Y(t) + e_2(t). \tag{5}
\]

where the derivative terms describe the changing levels of the inventory, \(a, b\) and \(c'\) are the system gains, \(\tau_1\) and \(\tau_2\) are the inventory time constants, and \(\theta_1, \theta_2\) and \(\theta_3\) are the time delays. It can be shown that at steady-state, Equations (4) and (5) simplify to the structural models in Equations (1) and (2).

Higher order differential equations could be used to describe more complex dynamic behaviour while still adhering to relationships depicted in Figure 2(b). While this simple fluid analogy reflects structural relationships defined by Equations (1) and (2), the resulting differential equations are relatively restrictive. Specifically, the outcome inventory dynamics are bound by a single time constant, despite the fact that the inventory accepts contributions from both the independent variable input and the mediator inventory outlet. This restriction is not necessary in a behavioural setting. Figure 3(a) is a less restrictive fluid analogy that describes mediated behaviour change. In this analogy, each pathway from Figure 2(a) is represented by an inventory, each with its own characteristic dynamics. Figure 3(b) is the corresponding block diagram and highlights the fact that \(Y\) is the result of two processes, where \(Y_D\) accounts for the outcome change that is a direct result of the input variable change, and \(Y_I\) accounts for the outcome change that is an indirect result of the input variable change via the mediator variable change. Figure 3(b) also highlights the parallel-cascade nature of time-varying behaviours in a mediational relationship. Equations (6) and (7) are the corresponding Laplace-domain models:

\[
M(s) = P_a(s) X(s) + d_1(s), \tag{6}
\]

\[
Y(s) = P_c(s) X(s) + P_b(s) M(s) + d_2(s). \tag{7}
\]

In accordance with McCarthy et al. (2008a), this article primarily treats \textit{Craving} and \textit{Cigsmked} as the mediator and outcome, respectively. The independent variable input, \textit{Quit}, is modelled as a unit step occurring on the quit date and corresponds to a transition from not attempting to quit smoking to attempting to quit. \(d_1\) and \(d_2\) in Equations (6) and (7) represent process disturbances, as opposed to measurement noise. In this framework, they represent un-modelled factors that influence the mediator and outcome. In this context, \(d_1\) represents factors other than the initiation of a quit attempt that contribute to, or mitigate, \textit{Craving}, such as negative or positive life events (i.e., changes in \textit{Stress}); \(d_2\) represents factors other than the initiation of a quit attempt or changing \textit{Craving} levels that influence \textit{Cigsmked}, such as the accessibility or inaccessibility of cigarettes. If these process disturbances are measured and uncorrelated with the inputs to \(P_a(s), P_b(s),\) and \(P_c(s),\) they could be explicitly modelled (Ljung, 1999). In behavioural health settings, explicitly modelling disturbances is challenging, as additional influences that are truly independent and exogenous
Figure 3. (a) Generalised fluid analogy for a mediated behavioural intervention developed from production inventory management models in supply chains. (b) Block diagram of statistical mediation.

are often not obvious. Consequently, accurate measurement and estimation of reliable models presents significant practical and related estimation issues. In the following, it is assumed that process disturbances are uncorrelated with the inputs and not of significant influence. The group averaging likely effectively filters out uncontrolled disturbances, suggesting these assumptions are reasonable. Ultimately, reliable identification and characterisation of possible process disturbances would require novel identification experiments. With novel clinical trial data, measurements of hypothesised disturbances could be included in estimation of Equations (6) and (7) and validated (e.g., through cross-validation); future validation of a time-varying disturbance could significantly contribute to development of an improved—engineering-based or otherwise —smoking intervention, as it would ultimately offer tobacco treatment practitioners an additional degree-of-freedom on which to intervene.

In fitting the Laplace-domain models in Equations (6) and (7) to the Craving and Cigsmked signals, a prediction-error approach is used to estimate continuous-time linear models from sampled data (Ljung, 2009). Model estimation initially employed the lowest order equation structure possible (a gain), as preliminary visual inspection of the data indicated low-order dynamic phenomena; transfer function structures with gradually increased complexity were evaluated as necessary according to the following iterative estimation and validation procedure:
(1) Estimation of $P_a(s)$ as a single-input single-output (SISO) system with Quit as the input and Craving as the output according to a given low-order transfer function structure.

(2) Simultaneous estimation of $P_a(s)$ and $P_c(s)$ as a multi-input single-output (MISO) system with Craving and Quit as the inputs and Cigsmked as the output according to given low-order transfer function structures for $P_a(s)$ and $P_c(s)$.

(3) Simulation of the Craving and Cigsmked responses to Quit according to the estimated $P_a(s)$, $P_b(s)$, and $P_c(s)$ expressions.

(4) Evaluation of Craving and Cigsmked goodness-of-fit on a 0% to 100% scale, calculated according to the following criterion:

$$\text{Fit} \% = 100 \left(1 - \frac{||y(t) - \hat{y}(t)||_2}{||y(t) - \bar{y}||_2}\right), \quad (8)$$

where $y(t)$ is the data to which the model is fit, $\hat{y}(t)$ is the simulated output, and $\bar{y}$ is the average of all $y$ values.

This procedure was implemented in MATLAB through a custom graphical user interface (GUI) built for flexible model estimation. Using the GUI, the four steps were repeated for different combinations of $P_a(s)$, $P_b(s)$, and $P_c(s)$ transfer function structures with parameter estimation relying on the pem command from the System Identification Toolbox in MATLAB. To use this routine, the input and output data were defined as an iddata object and the structure of the model to be estimated was defined as an idproc model object. For specification of the idproc model structure, the process models notation was used in which a single-output continuous-time model transfer function structure is specified and can feature one to three poles, an integrator, a zero, and a time-delay (Ljung & Singh, 2012). The idproc and process models functionality employs an indirect continuous-time estimation approach in which discrete-time estimation methods are first used before the resulting discrete-time representation is transformed into the equivalent continuous-time model. This two-step approach has the advantage of drawing from established discrete-time estimation methods to produce consistent and statistically efficient parameter estimates, and still results in continuous-time models that parsimoniously represent complex behaviours and can more easily be interpreted (Garnier et al., 2008). In the iterative four-step procedure used here, a set of model estimates were ultimately identified as appropriate representations of the behavioural dynamics for each group average and the single subject examples according to goodness-of-fit values, a concern for model parsimony, evaluation of parameter realisability, and through simulation.

Table 2 contains the parameter estimates, settling times (in days), and goodness-of-fit values for the mediation models; ILD and model outputs are shown in Figures 4 and 5. The iterative estimation procedure’s employment of low-order structures where complexity was increased only as necessary, the associated high goodness-of-fit values, and the corresponding simulations, which appear to accurately model the dynamic features observed in the ILD, suggest the following transfer function structures adequately represent cessation as a Craving-mediated process:

$$P_a(s) = \frac{a(\tau_a s + 1)}{(\tau_1 s + 1)}, \quad (9)$$

$$P_b(s) = \frac{b}{(\tau_3 s + 1)}, \quad (10)$$

$$P_c(s) = c'. \quad (11)$$

The estimated models feature high-fit percentages according to Equation (8) for the group averages.

<table>
<thead>
<tr>
<th>Treatment data set</th>
<th>AC Avg</th>
<th>ANc Avg</th>
<th>PC Avg</th>
<th>PNe Avg</th>
<th>AC Sgl</th>
<th>PNe Sgl</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mediator fit [%]</td>
<td>87.77</td>
<td>78.88</td>
<td>77.80</td>
<td>64.72</td>
<td>69.44</td>
<td>44.80</td>
</tr>
<tr>
<td>Outcome fit [%]</td>
<td>89.17</td>
<td>83.06</td>
<td>91.49</td>
<td>84.38</td>
<td>77.09</td>
<td>58.98</td>
</tr>
<tr>
<td>$ab$</td>
<td>0.94</td>
<td>1.96</td>
<td>1.45</td>
<td>1.17</td>
<td>-0.24</td>
<td>19.34</td>
</tr>
<tr>
<td>Mediator settling time [days]</td>
<td>35.69</td>
<td>35.91</td>
<td>35.82</td>
<td>35.90</td>
<td>26.34</td>
<td>33.87</td>
</tr>
<tr>
<td>Outcome settling time [days]</td>
<td>34.56</td>
<td>35.26</td>
<td>35.60</td>
<td>35.29</td>
<td>10.64</td>
<td>33.86</td>
</tr>
<tr>
<td>$a$</td>
<td>-11.10</td>
<td>-8.38</td>
<td>-7.12</td>
<td>-3.90</td>
<td>-20.38</td>
<td>3.10</td>
</tr>
<tr>
<td>$\tau_a$</td>
<td>-2.28</td>
<td>-4.60</td>
<td>-14.18</td>
<td>-24.21</td>
<td>-4.23</td>
<td>100.00</td>
</tr>
<tr>
<td>$\tau_1$</td>
<td>7.74</td>
<td>10.99</td>
<td>18.34</td>
<td>17.13</td>
<td>6.01</td>
<td>16.47</td>
</tr>
<tr>
<td>$b$</td>
<td>-0.08</td>
<td>0.23</td>
<td>-0.20</td>
<td>-0.30</td>
<td>0.01</td>
<td>6.25</td>
</tr>
<tr>
<td>$\tau_3$</td>
<td>4.59</td>
<td>2.89</td>
<td>0.42</td>
<td>1.89</td>
<td>1.29</td>
<td>95.53</td>
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</table>
order of the structures indicate that over-parameterisation is not taking place. In general, the high signal-to-noise ratios of the group average data sets are conducive to model estimation with high goodness-of-fit values, regardless of the transfer function structure. The lower mediator fit value for the PNc group supports this assertion, as this group’s Craving appears to have a lower signal-to-noise ratio than its counterparts.

The net decrease in Craving is greatest in the AC group, smallest in the PNc group, and follows a logical relationship to treatment condition (a is −11.10 for AC, −8.38 for ANc, −7.12 for PC, and −3.90 for PNc). As indicated by the negative system zeroes, the group average craving signals feature pronounced inverse response. It is known that a zero term in a dynamical systems model can result from two lower order sub-processes in parallel. Because inverse response in Craving results from a unit step (Quit), it can be deduced that $P_a(s)$ reflects parallel sub-processes in competition with each other. The first sub-process has a positive gain and faster speed of response than the second, negative-gained sub-process. The positive-gained sub-process corresponds to the immediate, quit-induced increase in Craving while the negative-gained subprocess corresponds to the post-quit settling of Craving to below baseline levels. For the case that $P_a(s)$ is described by the differential equation structure in Equation (9), $P_a(s) = P_a(s) + P_{o1}(s)$; $P_{o1}(s) = K_{o1}$ and $P_{o2}(s) = K_{o2}/(\tau_1s + 1)$, where the time constant is equal to that for the overall $P_a(s)$ function. It follows that $K_{o1} = -a\tau_2/\tau_1$ and $K_{o2} = a - K_{o1}$, where $a$ is the $P_a(s)$ gain, $\tau_o$ is the $P_a(s)$ zero, and $\tau_1$ is the $P_a(s)$ time constant. This notion of competing parallel processes within the overall smoking cessation process agrees with the observation that quitting smoking involves delayed and immediate gratification motives (executive and impulsive neurological processes, respectively) that compete during a quit attempt (Bickel et al., 2007). Such insight into the nature of these underlying sub-processes highlights the utility of a continuous-time system identification approach, as the implications of the $P_a(s)$ transfer function’s first order with zero structure were easily identified.

For the group average Cigsmked models, there is a dramatic quit-date drop in smoking followed by a relatively small and slow resumption. This dramatic quit-date smoking reduction is modelled by $P_a(s)$. Considering the treatment group averages, the magnitude of the initial drop is largest for the AC group, a 15.01 cigarette per day decrease, and smallest for the PNc group, a 10.42 cigarette per day decrease. For each model corresponding to the parameter estimates in Table 2, the direct contribution of the quit attempt
to Cigsmked is immediate, and the \( Y_D \) signal in Figure 3 acts as a step of magnitude \( c' \). The resumption of smoking is modelled by the mediated pathway, specifically \( P_3(s) \). For the group averages, the speed of resumption is small, as the \( P_3(s) \) time constant, \( \tau_3 \), is under five days for the groups. The speed of resumption of smoking does not strictly adhere to an expected relationship with respect to treatment condition: the AC group features the largest \( \tau_3 \) and the PC group has the smallest. The magnitude of resumption during a quit attempt is quantified with \( ab \), and is relatively small for the groups. Comparing \( ab \) and \( ab + c' \) values, the mediated pathway and net outcome gains, respectively, it is apparent that the mediated pathway’s contribution to the net effect of the quit attempt is consistently small for all of the group averages. Interestingly, the mediated pathways’ relative contribution to the outcome does not follow a natural progression in terms of relationship to treatment; the mediated pathway’s contribution to the outcome is 6.3% for the AC group, 14.9% for ANc, 10.7% for PC, and 13.0% for PNc.

As seen in Figure 5, the AC single subject example appears to successfully quit smoking. The estimated mediation model for this subject is consistent with this observation. Specifically, the magnitude of resumption is near zero \((ab \text{ equal to } -0.24)\), and the speed at which quit success is achieved is significantly faster than that of the group average counterparts, as the outcome settling time is 10.64 days. Conversely, the PNC single subject example appears to fully relapse (comparatively large \( b \) estimate) and does not feature inverse response (positive values for both the \( a \) and \( \tau_a \) values).

The estimated single subject models are generally less accurate, particularly the PNC subject models. This can be attributed to greater variance in the single subject data. The greater degree of variance is evident in Figure 1: both subjects’ baseline signals are very noisy, feature Craving signals with lower signal-to-noise ratios, and the PNC subject’s resumption also shows greater variance. These data quality issues are typical when considering single subject data and consequently pose a significant challenge to optimisation of smoking interventions, given that patient-specific models would ideally act as the basis for development of personalised smoking cessation treatments. As parameters in continuous-time models are not a function of the sampling time, a continuous-time system identification approach is appealing in terms of managing data quality issues such as missing data and non-constant measurement intervals – both common characteristics of self-reported behavioural data (McCarty et al., 2008b; Timms et al., 2012). Discussed in more detail in Section 5, future estimation of reliable single subject smoking cessation models may benefit from data collected in clinical trials designed with system identification in mind (Deshpande, Rivera, & Younger, 2012; Ljung, 1999; Rivera, 2012).

Using an iterative estimation procedure similar to that previously described, models for the ‘reverse’ mediation scenario were also estimated, where Cigsmked was the mediator and Craving the outcome (models not shown). This similarly resulted in models with high goodness-of-fit values and corresponding simulations that appeared to accurately represent the Craving and Cigsmked dynamics. This suggests that a Craving-Cigsmked inter-relationship captured in the clinical trial ILD is not fully described by a single mediation model. This significantly motivated identification of an alternative description of the smoking cessation process that accounts for this inter-relationship in a parsimonious manner.

4. Self-regulation model

One of the greatest opportunities afforded by ILD is the ability to study self-regulating and self-exciting phenomena (negative and positive feedback, respectively) within behaviour change processes (Collins, 2006). Regulatory behaviours have historically been of significant interest in terms of characterising addiction generally and smoking behaviours specifically (Carver & Scheier, 1998; Solomon, 1977; Solomon & Corbit, 1974; Velicer et al., 1992). The Nicotine Regulation Model, for example, proposes that smoking is done in an attempt to maintain a smoker’s blood nicotine set point, where deviations from this set point are the result of environmental conditions (e.g., cigarettes not readily available; Velicer et al. 1992). More complicated but conceptually similar mechanisms propose that cigarettes are smoked in order to regulate emotional states or an Urge set point (Solomon, 1977; Solomon & Corbit, 1974; Timms et al., 2013, in press; Velicer et al., 1992; Walls & Rivera, 2009); these emotional states may be directly affected by environmental factors unrelated to the act of smoking (Velicer et al., 1992). However, the ability of such theories to inform effective treatment strategies, such as time-varying adaptive interventions, has been limited. This is largely due to the historically significant challenges associated with effective measurement and mathematical modelling of behavioural dynamics (Riley et al., 2011).

Figure 6 depicts a block diagram of the smoking cessation process that features self-regulation. Generally, Figure 6 suggests that cigarette smoking is done in order to maintain a Craving set point and an attempt to quit smoking is a disturbance on this self-regulatory relationship. Essentially, this block diagram reflects an intuitive process: over time, an increasing desire to smoke leads to smoking activity, which then reduces that desire in the short-term. Specifically, Figure 6 describes a feedback loop in which a biochemical or psychological controller, \( C(s) \), responds to the deviation, \( e \), between a craving set point, \( r_{crav} \), and the actual measured craving signal \( e = r_{crav} - Craving \). Cigsmked is a sum of the outputs from \( C(s) \), the craving self-regulator, and \( P_D(s) \), the effect of the quit attempt; Cigsmked then acts as an input to \( P(s) \), producing
Craving. The associated closed-loop transfer function structures are as follows:

\[
Craving = \left( \frac{PC}{1+PC} \right) r_{\text{crav}} + \left( \frac{PP_d}{1+PC} \right) \text{Quit}. \tag{12}
\]

\[
\text{Cigsmked} = \left( \frac{C}{1+PC} \right) r_{\text{crav}} + \left( \frac{P_d}{1+PC} \right) \text{Quit}. \tag{13}
\]

As the output of \( P(s) \) is \( \text{Craving} \), this function will require a system zero, which stems from the sum of two sub-processes in parallel. Figure 6 depicts these underlying, competing processes. Mapping the self-regulatory relationship in Figure 6 to a generalised description of self-regulating behaviours described in Carver and Scheier (1998), \( P(s) \) and \( C(s) \) correspond to ‘Effect on Environment’ and ‘Behaviour’ processes, respectively.

As in the case of the mediation models, estimating the self-regulation models employed a prediction-error approach to obtain continuous-time linear models from sampled data (Ljung, 2009). The system identification procedure is similar to that previously described:

1. Estimation of \( P(s) \) as a SISO system with \( \text{Cigsmked} \) as the input and \( \text{Craving} \) as the output according to a given low-order transfer function structure.
2. Simultaneous estimation of \( C(s) \) and \( P_d(s) \) as a MISO system with \( e = r_{\text{crav}} - \text{Craving} \) and \( \text{Quit} \) as the inputs and \( \text{Cigsmked} \) as the output according to giving low-order transfer function structures for \( C(s) \) and \( P_d(s) \).
3. Simulation of the \( \text{Craving} \) and \( \text{Cigsmked} \) responses to \( \text{Quit} \) according to the estimated \( P(s) \), \( C(s) \), and \( P_d(s) \) expressions.
4. Calculation of \( \text{Craving} \) and \( \text{Cigsmked} \) goodness-of-fit according to the criterion in Equation (8).

This procedure was implemented in MATLAB through the previously described custom GUI that relied on the pem routine, an idproc model object, and the process models notation for estimation (Ljung & Singh, 2012). The four steps were repeated for different combinations of transfer function structures, beginning with the lowest order (gain-only) equation structure, with structural complexity increased as necessary. Similarly, various craving set points were examined: baseline \( \text{Craving} \) level, a linearly decreasing \( \text{Craving} \) function, and absolute \( \text{Craving} \) equal to zero. In assessing the group average and single subject candidate model estimates for goodness-of-fit, model parsimony, parameter realisability, and through simulation, the appropriate \( r_{\text{crav}} \) value was found to equal the baseline \( \text{Craving} \) level, and the following transfer function structures were found to appropriately represent the observed cessation dynamics:

\[
P(s) = \frac{K_1 (\tau_{a_s} s + 1)}{(\tau_1 s + 1)}, \tag{14}
\]

\[
P_d(s) = K_d, \tag{15}
\]

\[
C(s) = \frac{K_c}{(\tau_c s + 1)}. \tag{16}
\]

The parameter estimates and goodness-of-fit percentages are tabulated for the treatment group averages and the single subjects in Table 3. The corresponding model outputs are depicted in Figures 4 and 5.

As evident in Table 3 and Figures 4 and 5, high goodness-of-fit values and high fidelity simulations are obtained with low-order transfer function structures. As before, the negative system zero, \( \tau_{a_s} \), indicates \( P(s) \) represents the sum of two competing sub-processes in parallel. Whereas the group average mediation models suggest the sub-process with the faster speed of response is positive-gained, it is now the negative-gained sub-process that has an immediate speed of response \( P_1(s) = K_{p_1} \), where \( K_{p_1} < 0 \); conversely, it is the positive-gained function that has the slower speed of response \( P_2(s) = K_{p_2}/(\tau_{p_2} s + 1) \), where \( K_{p_2} > 0 \) and \( \tau_{p_2} \) equals \( \tau_1 \) from Equation (14) and Table 3. The negative value of \( K_{p_1} \) corresponds to the initial increase in \( \text{Craving} \) that results from the quit-induced, step-like initial decrease in \( \text{Cigsmked} \). The positive value of \( K_{p_2} \) corresponds to the settling of \( \text{Craving} \) to below baseline levels that results from the dramatic reduction in the group average \( \text{Cigsmked} \) signals. As the group average estimates in Table 3 for \( K_1 \) are positive, it follows that the magnitude of \( K_{p_1} \) is greater than that of \( K_{p_1} \). As previously described, the sum of two sub-processes agrees with the concept of dual impulsive and executive neurological processes that compete during a quit attempt (Bickel et al., 2007). Altogether, the craving reduction per unit decrease in daily cigarettes smoked is larger for the active drug groups versus the placebo groups: \( K_1 \) equal to 0.77 for the AC group, 0.74 for ANc, 0.50 for PC, and 0.52 for PNc.
Table 3. Self-regulation model parameter estimates and goodness-of-fit values.

<table>
<thead>
<tr>
<th>Treatment data set</th>
<th>AC Avg</th>
<th>ANc Avg</th>
<th>PC Avg</th>
<th>PNC Avg</th>
<th>AC Sgl</th>
<th>PNC Sgl</th>
</tr>
</thead>
<tbody>
<tr>
<td>Craving fit [%]</td>
<td>87.33</td>
<td>77.65</td>
<td>77.51</td>
<td>62.25</td>
<td>66.90</td>
<td>57.59</td>
</tr>
<tr>
<td>Cigsmked fit [%]</td>
<td>89.16</td>
<td>83.03</td>
<td>91.44</td>
<td>84.12</td>
<td>77.09</td>
<td>62.99</td>
</tr>
<tr>
<td>$P(s) K_1$</td>
<td>0.77</td>
<td>0.74</td>
<td>0.50</td>
<td>0.52</td>
<td>1.57</td>
<td>2.21</td>
</tr>
<tr>
<td>$P(s) \tau$</td>
<td>-1.99</td>
<td>-3.76</td>
<td>-14.34</td>
<td>-21.90</td>
<td>-3.05</td>
<td>3.45</td>
</tr>
<tr>
<td>$P(s) \tau_1$</td>
<td>8.22</td>
<td>14.23</td>
<td>18.70</td>
<td>26.75</td>
<td>6.88</td>
<td>10.76</td>
</tr>
<tr>
<td>$C(s) K_c$</td>
<td>0.08</td>
<td>0.23</td>
<td>0.20</td>
<td>0.30</td>
<td>-0.01</td>
<td>-6.25</td>
</tr>
<tr>
<td>$C(s) \tau_c$</td>
<td>4.59</td>
<td>2.89</td>
<td>0.42</td>
<td>1.89</td>
<td>1.29</td>
<td>95.53</td>
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be accounted for by classic mediation. This is significant as a Craving-Cigsmked inter-relationship was originally suggested by the fact that both the mediation models presented and the reverse mediation models (not shown) have high goodness-of-fit values and high fidelity simulations. Altogether, smoking cessation behaviour change is more appropriately and parsimoniously represented as a self-regulatory process as opposed to a mediational relationship. Furthermore, very poor models result from estimation of reverse self-regulation models, where \( P(s) \) is said to accept Craving as the input, producing Cigsmked as the output, etc. This supports the case that Figure 6 is a more appropriate representation of the Craving and Cigsmked relationship (i.e., Craving is the variable being regulated).

5. Conclusions and recommendations

Drawing from ILD collected from a clinical trial of bupropion and counseling as aids to smoking cessation, system identification models were developed to describe smoking cessation as mediational and self-regulatory processes. Ultimately, these models differ in how they each describe the resumption of smoking: for mediation, resumption is the result of daily changes in Craving (via \( P_s(s) \)); for self-regulation, resumption is the result of a craving self-regulator. Analysis of both sets of models highlight the utility of continuous-time system identification in behavioural health settings. Specifically, continuous-time system identification facilitates estimation of parsimonious expressions that accurately represent complex dynamic features within the smoking cessation process. Furthermore, these continuous-time expressions are conducive to straightforward identification and interpretation of the dynamics – in this article, shedding light on the nature of the two competing sub-processes that together form Craving and that the craving self-regulator is a proportional-with-filter controller on average. Altogether, the models developed suggest that self-regulation more appropriately describes the process of smoking cessation. Regardless of structure, parameter values estimated from the group average data, which are signals with high signal-to-noise ratios, suggest both bupropion and counseling have some effect on craving and reduction of smoking behaviour.

This article has effectively shown that system identification is useful in analysis of smoking cessation interventions and for comprehensively describing the process of smoking cessation. The dynamical modelling strategy used here could be further applied to the clinical trial data examined here in order to study alternate signal relationships and behavioural mechanisms. Notably, participants in the UW-CTRI clinical trial completed up to seven self-reports at pseudo-random intervals throughout the day (McCarthy et al., 2008b). These reports assessed environmental factors as well as behavioural states as experienced by the participant at the time the report was completed (whereas the ER focused on a participant’s average behavioural state over the previous 24 hours). Behavioural data collected in this fashion is said to reflect the influence of an individual’s natural environment, and may measure such influences. Continuous-time models estimated using data from these reports could provide some insight into environmental sources of inter-individual variability, and consequently may act as the basis for more reliable idiographic model estimation. However, the generalisability of such models would still be limited due to the secondary nature of the analysis. In the future, more informative single subject data sets – and ultimately more reliable patient-specific smoking cessation models—could be obtained through a novel smoking cessation clinical trial that draws from experimental design techniques in system identification. Such a trial may vary intervention dosage over time (e.g., bupropion dose, counseling frequency), use self-reports more conducive to measurement of nuanced behavioural dynamics (e.g., Craving assessed on a 0 to 100 point scale), or feature a longer self-reporting protocol. However, the medical, practical and ethical concerns associated with human subjects would also have to be addressed simultaneously. A smoking cessation clinical trial designed to produce more informative single subject data sets would involve experimental design strategies similar to those described in Deshpande et al. (2012) and Deshpande and Rivera (2013). Specifically, Deshpande et al. (2012) and Deshpande and Rivera (2013) propose optimisation-based approaches for the design of periodic, deterministic input signals that facilitate cross-validation, constraint handling, and altogether ‘patient-friendly’ operation.

Ultimately, self-regulatory models similar to those estimated here could inform novel treatment strategies (Nandola & Rivera, 2013; Riley et al., 2011; Rivera, 2012; Timms et al., 2013). Although accurate models of patient-specific behavioural dynamics would ideally be used to personalise interventions, a more practical intervention design approach may rely on a self-regulatory model described in the block diagram in Figure 6, but is parameterised to reflect a representative failed quit attempt – full resumption in Cigsmked, inverse-free Craving, and no net change in Cigsmked or Craving. A representative model of a failed quit attempt could be similar to the PNC single subject model examined here. Timms et al. (2013) presents such an example of a representative model of quit attempt failure. For designing an adaptive smoking cessation intervention, such a representative model could be used in conjunction with controller-design principles to develop an algorithm that defines intervention adjustment (e.g., medication dosage increases), based on a patient’s baseline conditions, self-reported smoking and withdrawal symptoms (e.g., daily Craving and Cigsmked reports), and environmental or other disturbances (e.g., Stress). A Hybrid Model Predictive Control (HMPC) approach is appealing as it can simultaneously manage manipulated
variables that are on discrete scales (e.g., discrete medication dosage) and constraints (e.g., medication toxicity levels) in an optimal manner (Nandola & Rivera, 2013; Rivera, 2012; Timms et al., 2013). Figure 7 depicts the general form of this control scheme. An optimised, adaptive smoking intervention designed in an HMPC framework could also include features of other well-known control approaches. For example, variables that are effectively non-time-varying but may be relevant to the cessation process, such as the presence of a genetic variant in a patient’s nicotine metabolism genes (Chen et al., 2012), could act as scheduling variables in an intervention featuring gain-scheduled Model Predictive Control (Chisci, Falugi, & Zappa, 2003). Finally, event-based control offers controller capabilities that may be appropriate for intervention design. Specifically, event-based control concepts offer a way to mitigate relapse that may otherwise result from time-varying cues to smoke (e.g., proximity to smokers) or other disturbances (Pawlowski, Guzman, Normey-Rico, & Berenguel, 2012), and would be particularly appealing for interventions that draw from models of cessation dynamics on shorter time scales (e.g., within-day dynamic models).

Acknowledgements
The content is solely the responsibility of the authors and does not necessarily represent the views of the NIH.

Funding
This work was supported by an award from the American Heart Association, a National Research Service Award from the National Institute on Drug Abuse at the National Institutes of 950 Health [grant number F31 DA035035], and the Office of Behavioural and Social Sciences Research and the National Institute on Drug Abuse at the National Institutes of Health [grant number K25 DA021173], [grant number R21 DA024266], [grant number P50 DA10075].

Figure 7. Block diagram of a Hybrid Model Predictive Control approach to design of an optimal, adaptive smoking cessation intervention.

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