Chapter #

Nonparametric Modeling and Model-Based Control of the Insulin-Glucose System∗

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1. Introduction

Diabetes represents a major threat to public health with alarmingly rising trends of incidence and severity in recent years, as it appears to correlate closely with emerging patterns of nutrition/diet and behavior/exercise worldwide. The concentration of blood glucose in healthy human subjects is about 90 mg/dl and defines the state of normoglycemia. Significant and prolonged deviations from this level may give rise to numerous pathologies with serious and extensive clinical impact that is increasingly recognized by current medical practice. When blood glucose concentration falls under 60 mg/dl, we have the acute and very dangerous state of hypoglycemia that may lead to brain damage or even death if prolonged. On the other hand, when blood glucose concentration rises above 120 mg/dl for prolonged periods of time, we are faced with the detrimental state of hyperglycemia that may cause a host of long-term health problems (e.g. neuropathies, kidney failure, loss of vision etc.). The severity of the latter clinical effects is increasingly recognized as medical science advances and diabetes is revealed as a major lurking threat to public health with long-term repercussions.

Prolonged hyperglycemia is usually caused by defects in insulin production, insulin action (sensitivity) or both (Carson et al., 1983). Although blood glucose concentration depends also on the action of several other hormones (e.g. epinephrine, norepinephrine, glucagon, cortisol), the exact quantitative nature of this dependence remains poorly understood and the effects of insulin are considered the most important. So traditionally, the scientific community has focused on the study of this causal relationship (with infused insulin being the “input” and blood glucose being the “output” of a system representing this functional relationship), using mathematical modeling as the means of quantifying it. Needless to say, the employed mathematical model plays a critical role in

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achieving (or not) the goal of effective glucose control. In addition, blood glucose concentration depends on many factors other than hormones, such as nutrition/diet, metabolism, endocrine cycles, exercise, stress, mental activity etc. The complexity of these effects cannot be modeled explicitly in a practical context at the present time and, thus, the aggregate effect of all these factors is usually represented for modeling purposes as a stochastic “disturbance” that is additive to the blood glucose level (or its rate of change).

Numerous studies have been conducted over the last 40 years to examine the feasibility of continuous blood glucose concentration control with insulin infusions. Since the achievement of effective glucose control depends on the quantitative understanding of the relationship between infused insulin and blood glucose, much effort has been devoted over the last 30 years to the development of reliable mathematical and computational models (Bergman et al., 1981; Cobelli et al., 1982; Sorensen, 1985; Tresp et al., 1999; Hovorka et al., 2002; Van Herpe et al., 2006; Markakis et al., 2008a; Mitsis et al., in press). Starting with the visionary works of Kadish (Kadish, 1964), Pfeiffer et al. on the “artificial beta cell” (Pfeiffer et al., 1974), Albisser et al. on the “artificial pancreas” (Albisser et al., 1974) and Clemens et al. on the “biostator” (Clemens et al., 1974), the efforts for online glucose regulation through insulin infusions have ranged from the use of relatively simple linear control methods (Salzsieder et al., 1985; Fischer et al., 1990; Chee et al., 2003a; Hernjak & Doyle, 2005) to more sophisticated approaches including optimal control (Swan, 1982; Fisher & Teo, 1989; Ollerton, 1989), adaptive control (Fischer et al., 1987; Candas & Radziuk, 1994), robust control (Kienitz & Yoneyama, 1993; Parker et al., 2000), switching control (Chee et al., 2005; Markakis et al., in press) and artificial neural networks (Frank et al., 1998; Trajanoski & Wach, 1998). However, the majority of recent publications have concentrated on applying model-based control strategies (Salzsieder et al., 1990; Parker et al., 1999; Lynch & Bequette, 2002; Rubb & Parker, 2003; Hovorka et al., 2004; Hernjak & Doyle, 2005; Dua et al., 2006; Van Herpe et al., 2007; Markakis et al., 2008b) for reasons that are elaborated below.

These studies have had the common objective of regulating blood glucose levels in diabetics with appropriate insulin infusions, with the ultimate goal of an automated closed-loop glucose regulation (the holy grail of “artificial pancreas”). Due to the inevitable difficulties introduced by the complexity of the problem and the limitations of proper instrumentation or methodology, the original grand goal has often been substituted by the more modest goal of “diabetes management” (Harvey et al., 1986; Berger et al., 1990; Deutsch et al., 1990; Salzsieder et al., 1990) and the use of man-in-the-loop control strategies with partial subject participation such as meal announcement (Goriya et al., 1988; Fisher, 1991; Brunetti et al., 1993; Hejlesen et al., 1997; Shimoda et al., 1997; Chee et al., 2003b).

In spite of the immense effort and the considerable resources that have been dedicated to this task, the results so far have been modest, with many studies contributing to our better understanding of this problem but failing to produce an effective solution with potential clinical utility and applicability. Technological limitations have always been a major issue, but recent advancements in the technology of long-term glucose sensors and insulin micro-pumps (Laser & Santiago, 2004; Klonoff, 2005) removed some of these past roadblocks and presented us with new opportunities in terms of measuring, analyzing and controlling blood glucose concentration with on-line insulin infusions.

It is our view that the lack of a widely accepted model of the insulin-glucose system (that is accurate under realistic operating conditions) represents at this time the main obstacle in achieving the stated goal. We note that almost all efforts to date for modeling the
insulin-glucose system (and consequently, for developing control strategies based on these models) have followed the “parametric” or “compartmental” route, which postulates a specific model structure (in the form of a set of differential/difference and algebraic equations) based on specific hypotheses regarding the underlying physiological mechanisms, in accordance with existing knowledge and current scientific understanding. The unknown parameters of the postulated model are subsequently estimated from the data, usually through least-squares or Bayesian fitting (Sorenson, 1980). Although this approach retains physiological relevance and interpretability of the obtained model, it presents the major limitation of being constrained a priori and, therefore, being subject to possible biases that may narrow the range of its applicability. This constraint becomes even more critical in light of the intrinsic complexity of physiological systems that includes the presence of nonlinearities, nonstationarities and patient-specific dynamics.

We propose that this modeling challenge be addressed by the so-called “nonparametric” approach, which employs models of the general form of Volterra functional expansions and their many variants (Marmarelis, 2004). The main advantage of this generic model form is that it remains valid for a very broad class of systems and covers most physiological systems under realistic operating conditions. The unknown quantities in these nonparametric models are the “Volterra kernels” (or their equivalent representations that are discussed below), which are estimated by use of the available data. Thus, there is no need for a priori postulation of a specific model and no problems with potential modeling biases. The estimated nonparametric models are “true to the data” and capable of predicting the system output for all possible inputs. The latter attribute of “universal predictor” makes them suitable for the purpose of model-based control of complex physiological systems, for which accurate parametric models are not available under broad operating conditions.

This book chapter begins with a brief presentation of the nonparametric modeling approach and its comparative advantages to the traditional parametric modeling approaches, continues with the presentation of a nonparametric model of the insulin-glucose system and concludes with the demonstration of the feasibility of incorporating such a model in a model-based control strategy for the regulation of blood glucose.

2. Nonparametric Modeling

The modeling of many physiological systems has been pursued in the context of the general Volterra-Wiener approach, which is also termed nonparametric modeling. This approach views the system as a “black box” that is defined by its specific inputs and outputs and does not require any prior assumptions about the model structure. As mentioned before, the nonparametric approach is generally applicable to all nonlinear dynamic systems with finite memory and contains unknown kernel functions that are estimated in practice by use of the available input-output data. Although the seminal Wiener formulation of this problem required the use of long data-records of white-noise inputs (Marmarelis & Marmarelis, 1978), this requirement has been removed and nonparametric modelling is now feasible with arbitrary input-output data of modest length (Marmarelis, 2004). In this formulation, the dynamic relationship between the input \( i(t) \) and output \( g(t) \) of a causal, nonlinear system of order \( Q \) and memory \( M \) is described in discrete-time by the following general/canonical expression of the output in terms of a hierarchical series of discrete multiple convolutions of the input:
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\[ g(n) = \sum_{q=0}^{Q} \sum_{m_1=0}^{M} \ldots \sum_{m_q=0}^{M} k_q(m_1, \ldots, m_q)i(n-m_1) \ldots i(n-m_q) = \]

\[ k_0 + \sum_{m=0}^{M} k_1(m)i(n-m) + \sum_{m_1=0}^{M} \sum_{m_2=0}^{M} k_2(m_1, m_2)i(n-m_1)i(n-m_2) + \ldots \]

where the \( q \)th convolution term corresponds to the effects of the \( q \)th order nonlinearities of the causal input-output relationship and involves the Volterra kernel \( k_q(m_1, \ldots, m_q) \), which characterizes fully the \( q \)th order nonlinear properties of the system. The linear component of the model/system corresponds to first convolution term and the respective first order kernel \( k_1(m) \) corresponds to the traditional impulse response function of a linear system. The general model of Eq. (1) can approximate any causal and stable system with finite memory to a desired accuracy for appropriate values of \( Q \) (Boyd & Chua, 1984). This approach has been employed extensively for modeling physiological systems because of their intrinsic complexity (Marmarelis, 2004).

Figure 1: The architecture of the Laguerre-Volterra network (LVN) that yields efficient approximations of nonparametric Volterra models in a robust manner using short data-records under realistic operating conditions (see text for description).
Among the various methods that have been developed for the estimation of the
discrete Volterra kernels from input-output data, we select the method utilizing a Volterra-
equivalent network in the form of a Laguerre-Volterra Network (LVN), which has been
found to be efficient for the accurate representation of high-order systems in the presence
of noise using short input-output records (Mitsis & Marmarelis, 2002). Therefore, it is well
suited to the present application that typically relies on relatively short input-output records
and is characterized by considerable measurement errors and systemic noise. The LVN
model consists of an input layer of a Laguerre filter-bank and a hidden layer of \( K \) hidden
units with polynomial activation functions (Figure 1). At each discrete time \( n \), the input
signal \( i(n) \) is convolved with the Laguerre filters and the filter-bank outputs are
subsequently transformed by the hidden units, the outputs of which form additively the
model output. The unknown parameters of the LVN are the in-bound weights and the
coefficients of the polynomial activation functions of the hidden units, along with the
Laguerre parameter of the filter-bank and the output offset. These parameters are estimated
from input-output data through an iterative procedure based on gradient descent. The filter-
bank outputs \( v_j \) are the convolutions of the input \( i(l) \) with the respective impulse response function \( b_j \) of the \( j \)-th order discrete-time Laguerre function:

\[
b_j(m) = \alpha^{(m-j)/2} (1 - \alpha)^{j/2} \sum_{i=0}^{m-j} \left( -1 \right)^i \binom{m-j}{i} \alpha^{i-j} (1 - \alpha)^i\]

(2)

where the Laguerre parameter \( \alpha \) in Eq. (2) lies between 0 and 1 and determines the rate of
exponential decay of the Laguerre functions. As indicated in Figure 1, the weighted sums \( u_i \)
of the filter-bank outputs \( v_j \) are subsequently transformed into \( z_i \) by the hidden units
through polynomial transformations:

\[
u_i(n) = \sum_{j=0}^{L-1} w_{k,j} v_j(n)
\]

(3)

\[
z_i(n) = \sum_{k=1}^{Q} c_{q,k} u_k^q(n)
\]

(4)

The model output \( g(n) \) is formed as the summation of the hidden-unit outputs \( z_k \) and a
constant offset value \( g_0 \):

\[
g(n) = \sum_{k=1}^{L} z_k(n) + g_0 = \sum_{k=1}^{L} \sum_{q=1}^{Q} c_{q,k} u_k^q(n) + g_0
\]

(5)

where \( L \) is the number of functions in the filter-bank, \( K \) is the number of hidden units, \( Q \) is
the nonlinear order of the model and \( w_{k,j} \) and \( c_{q,k} \) are the in-bound weights and the
polynomial coefficients of the hidden units respectively. The input and output time-series
data are used to estimate the LVN model parameters (\( w_{k,j}, c_{q,k} \), the offset \( g_0 \) and the Laguerre
parameter \( \alpha \)) with an iterative gradient-descent algorithm as (Mitsis & Marmarelis, 2002):

\[
\delta^{(r+1)} = \delta^{(r)} + \gamma_s e^{(r)}(n) \sum_{k=1}^{L} f_k^{(r)}(u_k^{(r)}(n)) \sum_{j=0}^{Q} w_{k,j} [v_j(n-1) + v_{j+1}(n)]
\]

(6)

\[
w_{k,j}^{(r+1)} = w_{k,j}^{(r)} + \gamma_s e_j^{(r)}(n) f_k^{(r)}(u_k^{(r)}(n)) v_j(n)
\]

(7)
where $\delta$ is the square root of the Laguerre parameter $\alpha$; $\gamma$, $\beta$, and $\gamma$ are positive learning constants, $f$ denotes the polynomial activation function of Eq. (4), $r$ denotes the iteration index and $e^{(r)}(n)$ and $f^{(r)}(u_k(n))$ are the output error and the derivative of the polynomial activation function of the $k^{th}$ hidden unit evaluated at the $r^{th}$ iteration, respectively.

The equivalent Volterra kernels can be obtained in terms of the LVN parameters as:

$$ k_n(m_1,...,m_n) = \sum_{k=1}^{K} \sum_{m_{k-h}}^{m_{k-h}} \sum_{m_{k-h}}^{m_{k-h}} c_{m,k} \sum_{j=0}^{\frac{q}{m_h}} w_{k,h} \cdot b_{k,h}(m_1)\cdot b_{k,h}(m_n) $$

which indicates that the Volterra kernels are implicitly expanded in terms of the Laguerre basis and the LVN represents a parsimonious way of parameterizing the general nonparametric Volterra model (Marmarelis, 1993; Marmarelis, 1997; Mitsis & Marmarelis, 2002; Marmarelis, 2004).

The structural parameters of the LVN model ($L,K,Q$) are selected on the basis of the normalized mean-square error (NMSE) of the output prediction achieved by the model, defined as the sum of squares of the model residuals divided by the sum of squares of the de-meaned true output. The statistical significance of the NMSE reduction achieved for model structures of increased order/complexity is assessed by comparing the percentage NMSE reduction with the $alpha$-percentile value of a chi-square distribution with $p$ degrees of freedom ($p$ is the increase of the number of free parameters in the more complex model) at a significance level $alpha$, typically set at 0.05.

The LVN representation is just one of the many possible Volterra-equivalent networks (Marmarelis & Zhao, 1997) and is also equivalent to a variant of the general Wiener-Bose model, termed the Principal Dynamic Modes (PDM) model. The PDM model consists of a set of parallel branches, each one of which is the cascade of a linear dynamic filter (PDM) followed by a static, polynomial nonlinearity (Marmarelis, 1997). This leads to model representations that are more parsimonious and facilitate physiological interpretation, since the resulting number of PDMs has been found to be small (2 or 3) in actual applications so far. The PDM model is formulated next for a finite memory, stable, discrete-time SISO system with input $i$ and output $g$. The input signal $i(n)$ is convolved with each of the PDMs $p_k$ and the PDM outputs $u_k(n)$ are subsequently transformed by the respective polynomial nonlinearities $f_k$ to produce the model-predicted blood glucose output as:

$$ g(n) = g_b + f_1[u_1(n)] + ... + f_K[u_K(n)] =$$

$$ = g_b + f_1[p_1(n)\ast i(n)] + ... + f_K[p_K(n)\ast i(n)] $$

where $g_b$ is the basal value of $g$ and the asterisk denotes convolution. Note the similarity between the expressions of Eq. (5) and Eq. (10), with the only difference being the basis of functions used for the implicit expansion of the Volterra kernels (i.e., the Laguerre basis versus the PDMs) that makes the PDM representation more parsimonious – if the PDMs of the system can be found.
3. A Nonparametric Model of the Insulin-to-Glucose Causal Relationship

In the current section, we present and briefly analyze a PDM model of the insulin-glucose system (Figure 2), which is a slightly modified version of a model that appeared in (Marmarelis, 2004). This PDM model has been obtained from analysis of infused insulin–blood glucose data from a Type 1 diabetic over an eight-hour period. In the subsequent computational study it will be treated as the putative model of the actual system, in order to examine the efficacy of the proposed model-predictive control strategy. It should be emphasized that this model is subject-specific and valid only for the specific type of fast-acting insulin analog that was used in this particular measurement. Different types of insulin analogs are expected to yield different models for different subjects (Howey et al., 1994). The PDM model employed in each case must be estimated with data obtained from the specific patient with the particular type of infused insulin. Furthermore, this model is expected to be generally time-varying and, thus, it must be adapted over time at intervals consistent with the insulin infusion schedule.

Firstly, let us give a succinct mathematical description of the PDM model of Figure 2: the input \( i(n) \), which represents the concentration of infused insulin at discrete time \( n \) (not the rate of infusion as in many computational studies), is transformed by the upper (\( h_1 \)) and lower (\( h_2 \)) branches through convolution to generate the PDM outputs \( v_1(n) \) and \( v_2(n) \). Subsequently, \( v_1(n) \) and \( v_2(n) \) are mapped by the cubic nonlinearities \( f_1 \) and \( f_2 \) respectively; their sum, \( f_1(v_1) + f_2(v_2) \), represents the time-varying deviation of blood glucose concentration from its basal value \( g_0 \). To sum up, the blood glucose concentration at each discrete time \( n \) is given by:

**Figure 2:** The putative PDM model of the insulin–glucose system used in this computational study (see text for description of its individual components).
(11)

where \( g = 90 \text{ mg/dl} \) is a typical basal value of blood glucose concentration and \( D(n) \) represents the aforementioned “disturbance” term that incorporates all the other systemic and extraneous influences on blood glucose (described in detail later).

Remarkably, the two branches of the model of Figure 2 appear to correspond to the two main physiological mechanisms by which insulin affects blood glucose according to the literature, even though no prior knowledge of this was used during its derivation. The first mechanism (modeled by the upper PDM branch) is termed “glucolepsis” and reduces the blood glucose level due to higher glucose uptake by the cells (and storage of excess glucose in the liver and adipose tissues) facilitated by the insulin action. The second mechanism (modeled by the lower PDM branch) is termed “glucogenesis” and increases the blood glucose level through production or release of glucose by internal organs (e.g., converting glycogen stored in the liver), which is triggered by the elevated plasma insulin. It is evident from the corresponding PDMs in Figure 2 that the latter mechanism of glucogenesis is somewhat slower and can be viewed as a counterbalancing mechanism of “biological negative feedback” to the former mechanism of glucolepsis. Since the dynamics of the two mechanisms and the associated nonlinearities are different, they do not cancel each other but partake in an intricate act of dynamic counter-balancing that provides the desired physiological regulation. Note also that both nonlinearities shown in the PDM model of Figure 2 are supralinear (i.e., their respective outputs change more than linearly relative to a change in their inputs) and of significant curvature (i.e., second derivative); intuitively, this justifies why linear control methods, based on linearizations of the system, will not suffice and, thus, underlines the importance of considering a nonlinear control strategy in order to achieve satisfactory regulation of blood glucose.

The glucogenic branch corresponds to the combination of all factors that counteract to hypoglycaemia and can be triggered by the concentration of insulin; although their existence is an undisputed fact (even the early model of Sorensen (Sorensen, 1985) attests to that), to the best of our knowledge, none of the existing models in the literature exhibits a strong glucogenic component. This emphasizes the importance of being “true to the data” and the dangers from imposing a certain structure \textit{a priori}. Another consequence is that including a significant glucogenic factor complicates the dynamics and much more care should be taken in the design of a controller.

Unlike the extensive use of parametric models for the insulin-glucose system, there are very few cases to date where the nonparametric approach has been followed e.g., the Volterra model in (Florian & Parker, 2002) which is, however, distinctly different from the nonparametric model of Figure 2. A PDM model of the functional relation between spontaneous variations of blood insulin and glucose in dog was presented by Marmarelis et al. (Marmarelis et al., 2002) and exhibits some similarities to the model presented above. Driven by the fact that the Minimal Model (Bergman et al., 1981) and its many variations over the last 25 years is by far the most widely used model of the insulin-glucose system, the equivalent nonparametric model was derived computationally and analytically (i.e., the Volterra kernels were expressed in terms of the parameters of the Minimal Model) and was shown to differ significantly from the model of Figure 2 (Mitsis & Marmarelis, 2007). To emphasize the important point that the class of systems representable by the Minimal Model
and its many variations (including those with pancreatic insulin secretion) can be also represented accurately by an equivalent nonparametric model, although the opposite is not generally true, we have performed an extensive computational study comparing the parametric and nonparametric approaches (Mitsis et al., in press).

4. Model-Based Control of Blood Glucose

In this section we formulate the problem of on-line blood glucose regulation and propose a model predictive control strategy, following closely the development in (Markakis et al., 2008b). A model-based controller of blood glucose in a nonparametric setting has also been proposed by Rubb & Parker (Rubb & Parker, 2003); however, both the model and the formulation of the problem are quite different than the ones presented here.

4.1 Closed-Loop System of Blood Glucose Regulation

The block diagram of the proposed closed-loop control system for on-line regulation of blood glucose is shown in Figure 3. The PDM model presented in Section 3 plays the role of the real system in our simulations and defines the deviation of blood glucose from its basal value, in response to a given sequence of insulin infusions \( i(n) \). The glucose basal value \( g_0 \) and the glucose disturbance \( D(n) \) are superimposed on it to form the total value of blood glucose \( g(n) \). Measurements of the latter are obtained in practice through commercially-available continuous glucose monitors (CGMs) that generate data-samples every 3 to 10 min (depending on the specific CGM). In the present work, the simulated CGM is assumed to make a glucose measurement every 5 min. Since the accuracy of these CGM measurements varies from 10% to 20% in mean absolute deviation by most accounts, we add to the simulated glucose data Gaussian “measurement noise” \( N(n) \) of 15% (in mean absolute deviation) in order to emulate a realistic situation. Moreover, the (short) time lag in the concentration of glucose between blood and interstitial fluids is modeled as a pure delay of 5 minutes in the measurement of \( g(n) \). A digital, model-based controller is used to compute the control input \( i(n) \) to the system, based on the measured error signal \( e(n) \).
(the difference between the targeted value of blood glucose concentration $g_t$ and the measured blood glucose $g_m(n)$). The objective of the controller is to attenuate the effects of the disturbance signal and keep $g(n)$ within bounds defined by the normoglycaemic region. Usually the targeted value of blood glucose $g_t$ is set equal (or close) to the basal value $g_0$ and a conservative definition of the normoglycaemic region is from 70 to 110 mg/dl.

### 4.2 Glucose Disturbance

It is desirable to model the glucose disturbance signal $D$ in a way that is consistent with the accumulated qualitative knowledge in a realistic context and similar to actual observations in clinical trials - e.g. see the patterns of glucose fluctuations shown in (Chee et al., 2003b; Hovorka et al., 2004). Thus, we have defined the glucose disturbance signal through a combination of deterministic and stochastic components:

- **Terms of the exponential form $n \cdot \exp(-0.19 \cdot n)$**, which represent roughly the metabolic effects of Lehmann-Deutsch meals (Lehmann & Deutsch, 1992) on blood glucose of diabetics. The timing of each meal is fixed and its effect on glucose concentration has the form of a negative gamma-like curve, whose peak-time is at 80 minutes and peak amplitude is 100 mg/dl for breakfast, 350 mg/dl for lunch and 250 mg/dl for dinner.

- **Terms of the exponential form $n \cdot \exp(-0.15 \cdot n)$**, which represent random effects due to factors such as exercise or strong emotions. The appearance of these terms is modeled with a Bernoulli arrival process with parameter $p=0.2$ and their effect on glucose concentration has again the form of a negative gamma-like function with peak-time of approximately 35 minutes and peak amplitude uniformly distributed in [-10, 30] mg/dl.

- **Two sinusoidal terms of the form $\alpha_i \cdot \sin(\omega_i \cdot n + \phi_i)$** with specified amplitudes and frequencies ($\alpha_i$ and $\omega_i$) and random phase $\phi_i$ uniformly distributed within the range [-$\pi/2$, $\pi/2$]. These terms represent circadian rhythms (Lee et al., 1992; Van Cauter et al., 1992) with periods 8 and 24 hours and amplitudes around 10 mg/dl.

- **A constant term $B$** which is uniformly distributed within the range [50, 80] and represents a random bias of the subject-specific basal glucose from the nominal value of $g_0$ that many diabetics seem to exhibit.

An illustrative example of the combined effect of these disturbance factors on glucose fluctuations can be seen in Figure 4.
Figure 4: Typical effect of glucose disturbance on the levels of blood glucose over a period of 24 hours.

The structure of the glucose disturbance signal described above is not known to the controller. However, in order to apply Model Predictive Control (MPC - the specific form of model-based control employed here) it would be desirable to predict the future values of the glucose disturbance term $D(n)$ within some error bounds, so that we can obtain reasonable predictions of the future values of blood glucose concentration over a finite horizon. To achieve this, we hypothesize that the glucose disturbance signal $D$ can be considered as the output of an Auto-Regressive (AR) model:

$$D(n) = D \cdot a + w(n)$$  \hspace{1cm} (12)

where $D = [D(n-1) \ D(n-2) \ ... \ D(n-K)]$, $a = [a_1 \ a_2 \ ... \ a_K]^T$ is the vector of coefficients of the AR model, $w(n)$ is an unknown “innovation process” (usually viewed as a white sequence), and $K$ is the order of the AR model. At each discrete-time instant $n$, the prediction task consists of estimating the coefficient vector $a$, which in turn allows the estimation of the future values of glucose disturbance: according to the Certainty Equivalence Control (Bertsekas, 2005) we can use the estimated disturbance values as if they were actual values, in order to compute the glucose disturbance over the desired future horizon, using the AR model sequentially. The estimation of the coefficient vector can be performed with least-squares methods (Sorenson, 1980). Note, however, that we cannot know a priori whether the AR model is suitable for capturing the glucose disturbance presented above or if the least-squares criterion is appropriate in the AR context. What is most pertinent is the lack of correlation among the residuals. For this reason, we also compute the autocorrelation of the residuals and seek to make its values for all non-zero lags statistically insignificant, a fact indicating that all structured or correlated information in the glucose disturbance signal has
been captured by the AR model. A critical part of this procedure is the determination of the best AR model order $K$ at every discrete-time instant - in the present study this is achieved using the Akaike Information Criterion (Akaike, 1974).

### 4.3 Model - Based Control of Blood Glucose

Here we outline the concept of Model Predictive Control (MPC), which is at the core of the proposed control algorithm. Having knowledge of the nonlinear model and all the past input-output pairs, the goal of MPC is to determine the control input value $i(n)$ at every time instant $n$, so that the following cost function is minimized:

$$J(n) = [g(n+p|n) - g_t]^T \cdot \Gamma_y \cdot [g(n+p|n) - g_t] + \Gamma_U \cdot i(n)^2$$

(13)

where $g(n+p|n)$ is the vector of predicted output values over a future horizon of $p$ steps using the model and the past input values, $\Gamma_y$ is a diagonal matrix of weighting coefficients assigning greater importance to the near-future predictions and $\Gamma_U$ a scalar that determines how “expensive” is the control input. We also impose a “physiological” constraint to the above optimization problem in order to avoid large deviations of plasma insulin from its basal value and, consequently, the risk of hypoglycaemia: we limit the magnitude of $i(n)$ to a maximum of 1.5 mU/L. The procedure is repeated at the next time step to compute $i(n+1)$ and so on -- details on MPC and relevant control issues can be found in (Camacho & Bordons, 2007; Bertsekas, 2005).

In our simulations, we considered a prediction horizon of 40 min ($p = 8$ samples) and exponential weighting $\Gamma_y$ with a time constant of 50 min. As measures of precaution against hypoglycaemia, we used a target value for blood glucose that is greater than the reference value ($g_t = 105$ mg/dl) and also applied asymmetric weighting to the predicted output vector, as in (Hernjak & Doyle, 2005), whereby we penalized 10 times more the deviations of the vector $g(n+p|n)$ that are below $g_t$. The scalar $\Gamma_U$ was set to 0 throughout our simulations.

### 4.4 Results

Throughout this section we assume that MPC has perfect knowledge of the nonlinear PDM model. Figure 5 presents MPC in action: the top panel shows the blood glucose levels without any control, apart from the basal insulin infusion (blue line), called also the “No-Control” case, and after MPC action (green line). The mean value (MV), standard deviation (SD) and the percentage of time that glucose is found outside the normoglycaemic region of 70-110 mg/dl (PTO) are reported between the panels for MPC and “No-Control”. The bottom panel shows the infused insulin profile determined by the MPC. Figure 6 presents the autocorrelation function of the estimated innovation process $w$: the fact that its values for all non-zero time-lags are statistically insignificant (smaller than the confidence bounds determined by the null hypothesis that the residuals are uncorrelated with zero mean) implies that the structure of the glucose disturbance signal is captured by the AR-Model. This result is important, considering that we have included a significant amount of stochasticity in the disturbance signal. In Figure 7 we show how the order of the AR model varies with time, as determined by the AIC, for the simulation case of Figure 5.
Figure 5: Model Predictive Control of blood glucose concentration: The top panel shows the blood glucose levels corresponding to the general stochastic disturbance signal, with basal insulin infusion only (blue line) and after MPC action (green line). The mean value (MV), standard deviation (SD) and percentage of time that the glucose is found outside the normoglycaemic region of 70-110 mg/dl (PTO) are reported between the panels for MPC and without control action. The bottom panel shows the insulin profile determined by the MPC.

Figure 6: Estimate of the autocorrelation function of the AR model residuals for the simulation run of Figure 5.
Figure 7: The time-variations of the AR model order (as determined by AIC) for the simulation run of Figure 5.

Figure 8 provides further insight into how the attenuation of glucose disturbance is achieved by MPC: the controller determines the precise amount of insulin to be infused, given the various constraints, so that the time-varying sum of the outputs of glucolepsis (blue line) and glucogenesis (green line) cancel the stochastic disturbance (red line) in order to maintain normoglycaemia. A comment, however, must be made on the large values of the various signals of Figure 8: the PDM model presented in Section 3 aims primarily to capture the input-to-output dynamics of the system under consideration and not its internal structure (like parametric models do). So, even though the PDMs of Figure 2 seem intuitive and can be interpreted physiologically, we cannot expect that all the signals included there will make physiological sense (apart from the input and the output signals of course).

Finally, in order to average out the effects of stochasticity in glucose disturbance upon the results of closed-loop regulation of blood glucose, we report in Table 1 the average performance achieved by MPC over 20 independent simulation runs of 48 hours each. The evaluation of performance is done by comparing the standard indices (mean value, standard deviation, percent of time outside the normoglycaemic region) for the MPC and the "No-Control" case. The total number of hypoglycaemic events is also reported in the last row, since it is critical for patient safety. The results shown in this Table and in the Figures above indicate that MPC can regulate blood glucose quite well (as attested by the significant improvement in all measured indices) and, at the same time, does not endanger the patient.
Figure 8: MPC preserves normoglycaemia by cancelling out the effects of glucose disturbance (red line), the glucoleptic branch (blue line) and the glucogenic (green line) branch.

<table>
<thead>
<tr>
<th></th>
<th>NO CONTROL</th>
<th>MPC</th>
</tr>
</thead>
<tbody>
<tr>
<td>MV</td>
<td>182.6</td>
<td>111.5</td>
</tr>
<tr>
<td>SD</td>
<td>89</td>
<td>42</td>
</tr>
<tr>
<td>PTO</td>
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<td>25</td>
</tr>
<tr>
<td>HYPO</td>
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<td>0</td>
</tr>
</tbody>
</table>

Table 1: Averages of 20 independent simulation runs of 48 hours each. Presented are the mean value (MV) and the standard deviation (SD) of glucose fluctuations, the percentage of time that glucose is found outside the normoglycaemic region 70-110 mg/dl (PTO) and the number of hypoglycaemic events, for the cases of no control action and MPC.

5. Discussion

This chapter is dedicated to the potential application of nonparametric modeling for model-based control of blood glucose through automated insulin infusions and seeks to:

- Briefly outline the nonparametric modeling methodology and present a data-based nonparametric model, in the form of Principal Dynamic Modes (PDM), of the dynamics between infused insulin and blood glucose concentration. This model form provides an accurate, parsimonious and interpretable representation of this causal relationship for a specific patient and was obtained using a relatively short
data-record. The estimation of nonparametric models (like the one presented here) is robust in the presence of noise and/or measurement errors and not liable to model misspecification errors that are possible (or even likely) in the case of hypothesis-based parametric or compartmental models. More information on the performance of nonparametric models in the context of the insulin-glucose system can be found in (Mitsis et al., in press).

- Show the efficacy of utilizing PDM models in Model Predictive Control (MPC) strategies for on-line regulation of blood glucose. The results of our computational study suggest that a closed-loop, PDM - MPC strategy can regulate blood glucose well in the presence of stochastic and cyclical glucose disturbances, even when the data are corrupted by measurement errors and systemic noise, without risking dangerous hypoglycaemic events.

- Suggest an effective way for predicting stochastic glucose disturbances through an Auto-Regressive (AR) model, whose order is determined adaptively by use of the Akaike Information Criterion (AIC) or other equivalent statistical criteria. It is shown that this AR model is able to capture the basic structure of the glucose disturbance signal, even when it is corrupted by noise. This simple approach offers an attractive alternative to more complicated techniques that have been previously proposed -- e.g. utilizing a Kalman filter (Lynch & Bequette, 2002).

A comment is warranted regarding the procedure of insulin infusions, either intravenously or subcutaneously. Various studies have shown that in the case of fast acting, intravenously infused insulin the time-lag between the time of infusion and the onset of its effect on blood glucose is not significant, e.g. see (Hovorka, 2005) and references within. However, in the case of subcutaneously infused insulin, the considerably longer time-lag may compromise the efficacy of closed-loop regulation of blood glucose. Although this issue remains an open problem, the contribution of this study is that it demonstrates that the dynamic effects of infused insulin on blood glucose concentration are “controllable” under the stipulated conditions, which seem realistic. Nonetheless, additional methodological improvements are possible, if the circumstances require them, which also depend on future technical advancements in glucose sensing and micro-pump technology, as well as the synthesis of even faster-acting insulin analogs.

There are numerous directions for future research, including improved methods for prediction of the glucose disturbance and the adaptability of the PDM model to the time-varying characteristics of the insulin-to-glucose relationship. From the control point of view, a critical issue remains the possibility of plant-model mismatch and its effect on the proposed MPC strategy (since the presented MPC results rely on the assumption that the controller has knowledge of an accurate PDM model). Last, but not least, it is obvious that the clinical validation of the proposed control strategy, based on nonparametric models, is the ultimate and necessary validation step in adopting this approach.
6. References


Nonparametric Modeling and Model-Based Control of the Insulin-Glucose System


Mitsis, G.; Markakis, M. & Marmarelis, V. (in press). Non-parametric versus parametric modeling of the dynamic effects of infused insulin on plasma glucose. *IEEE Transactions on Biomedical Engineering*


