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## Therapy-driven Deep Glucose Forecasting

Eleonora Maria Aiello<sup>a</sup>, Giuseppe Lisanti<sup>b</sup>, Lalo Magni<sup>c</sup>, Mirto Musci<sup>a</sup>, Chiara Toffanin<sup>a,\*</sup>

<sup>a</sup> University of Pavia, Department of Electrical, Computer and Biomedical Engineering, via Ferrata 5, Pavia 27100, Italy.
 <sup>b</sup> University of Bologna, Department of Computer Science and Engineering, Mura Anteo Zamboni 7, Bologna 40126, Italy.
 <sup>c</sup> University of Pavia, Department of Civil and Architecture Engineering, via Ferrata 5, Pavia 27100, Italy.

#### Abstract

The automatic regulation of blood glucose for Type 1 diabetes patients is the main goal of the artificial pancreas, a closed-loop system that exploits continue glucose monitor data to define an optimal insulin therapy. One of the most successful approach for artificial pancreas is the model predictive control, which exhibits promising result on both virtual and real patients. The performance of such controller is highly dependant on the reliability of the glucose-insulin model used for prediction purpose, which is usually implemented with classic mathematical models. In this paper we propose Deep Glucose Forecasting, a deep learning approach for forecasting glucose levels, based on a Long-Short Term Memory implementation. Our solution takes in input the previous values obtained through continue glucose monitor, the carbohydrate intake, the suggested insulin therapy and forecasts the interstitial glucose level of the patient. The proposed architecture has been trained on 100 virtual adult patients of the UVA/Padova simulator, and tested on both virtual and real patients. The proposed solution outperforms classical general-population models and reaches performance comparable to classical personalized models when fine-tuning is exploited on real patients. *Keywords:* diabetes, forecasting, prediction, deep learning, LSTM

### 1. Introduction

Type 1 Diabetes (T1D) is a chronic metabolic disease characterized by high Blood Glucose (BG) level, known as hyperglycaemia. Hyperglycaemia can cause long-term complications including damages to blood vessels, eyes, kidneys, and nerves and it is caused by the dysfunction of pancreatic  $\beta$ -cells responsible for the production of insulin. This hormone regulates the BG concentration by allowing cells and tissues to absorb glucose from the bloodstream. T1D patients need exogenous insulin injections to keep the glucose concentration in the euglycemic range. Their goal is to minimize diabetes complications related to hyperglycemia and simultaneously avoid hypoglycemia, a condition that could be caused by excessive insulin administration.

<sup>\*</sup>Corresponding author

*Email addresses:* eleonoramaria.aiello01@universitadipavia.it (Eleonora Maria Aiello), giuseppe.lisanti@unibo.it (Giuseppe Lisanti), lalo.magni@unipv.it (Lalo Magni), mirto.musci@unipv.it (Mirto Musci), chiara.toffanin@unipv.it (Chiara Toffanin)

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The automatic regulation of the BG concentration for people affected by T1D through exogenous insulin administrations [1, 2, 3] is the main purpose of the so-called artificial pancreas. The artificial pancreas is a closed-loop system that exploits the glucose measurements obtained via Continuous Glucose Monitor (CGM) to compute and automatically deliver the proper amount of insulin via subcutaneous insulin pump. The core of the artificial pancreas is the control algorithm that defines the optimal insulin amount to infuse. The Model Predictive Control (MPC) resulted one the most promising approach to this problem in the last years, obtaining successful results both *in silico* and *in vivo* [4, 5, 6, 7, 8, 9, 10]. The MPC approach exploits a glucose-insulin model to forecast the BG values in order to compute the optimal insulin therapy. For this reason, the predictive performance of the model plays a key role in the overall control performance.

Recently, a branch of the research was moved towards new identification techniques in order to have more effective models to be used for both the control algorithms and the safety systems. A complete review can be found in [11]. Thanks to the availability of a huge amount of data collected during long-period trials in free-living conditions, new data-driven approaches have been studied with promising results [12] [13]. These approaches have received an increasing attention in the last few years mainly because of the remarkable performance obtained in several research fields [14] [15]. Depending on the task at hand (i.e., classification, regression, prediction, etc.) the aim of these approaches is to learn a model directly from the data by minimizing a loss function between the output of the model and the expected value. Among these solutions, recurrent neural networks represent a family of deep learning architectures which have been explicitly designed to model the evolution over time of a phenomenon. In particular, given an input composed of a sequence of observations from a signal, such as the BG level in our scenario, these models try to predict its future value or values.

- The main goal of this work is the development of a forecasting model able to predict the future BG of a patient subject to several possible insulin treatments in order to define the optimal future insulin therapy for that patient. In this perspective, we propose a deep learning architecture for sequence-to-sequence prediction which is able to forecast the BG level of T1D patients. Our architecture is composed of two models, one observing the CGM measurements, insulin injections and carbohydrate intakes up to a given time t and
- <sup>35</sup> a second model that receives as input the future insulin that will be administered to the patient and the future carbohydrates that he/she will assume. Both models are composed of stacked Long-Short Term Memory (LSTM) [16] networks. The output of the two models is combined and given as input to a fully connected (FC) layer which is used to predict the future values of the Interstitial Glucose (IG), considering a fixed prediction horizon. Training is performed in a supervised fashion on a subset of identities, separated
- <sup>40</sup> from those that will be considered as test, in order to obtain a model which is able to generalize to new unseen data. The proposed architecture obtains state-of-the-art performance on both *in silico* and *in vivo* data, considering several prediction horizons.

#### 2. Related Works

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Several works treated the model identification problem in the last years by exploiting *in silico* **17**, **18**, **19**, **20**, **21**, **22**, **23**, **24**, **25**, **26** and *in vivo* **12**, **13**, **27** data. For a comprehensive literature review please refer to **11**, **28**. The *in silico* data used for model identification **17**, **18**, **22**, **24** were obtained through realistic closed-loop clinical protocols simulated via the UVA/Padova simulator **129**, **30** in order to produce a sufficient input-output excitation for identification purpose. On the other hand, *in vivo* data were collected during either short and controlled trials on hospitalized patients **18** and trials outside the hospital environment in **50** free-living conditions **4**, **5**, **6**, **7**, **8**, **31**.

The availability of such huge amount of data increased the interest in new data-driven techniques to identify a better glucose-insulin model to include in a MPC. Classical model can be either nonlinear or linear: the nonlinear models usually allow for a better approximation of the underlying dynamic, but being based on partial differential equations, they are extremely costly from a computational point-of-view. On the other hand, the linear model approximation is less precise, but their computational load is limited.

Recently, several MPC algorithms have been developed using different type of models. Among them, the MPC proposed in [32] used a nonlinear models whose parameters were re-estimated at each control step in order to take into account the daily variability of the patient. This MPC was tested in several clinical trials [33, 34, 35, 36, 37] with promising results. The Zone-MPC [38] was based on the third-order, discrete-time, linear time-invariant model proposed in [39]. Although the simple structure of this model,

the Zone-MPC obtained good results *in vivo* on both adults and adolescents [40] [41]. The MPC described in [42] (MPC-P) was synthesized on the basis of the average linear time-invariant metabolic model computed by averaging the model parameters of the virtual population of the UVA/Padova simulator. The MPC-P was tested in several clinical trials [31] [43] obtaining good results thanks to capability of the simulator to represent all the diabetic population and not only a restricted subgroup of individuals. However, a model with better predictive capabilities would increase the overall control performance. In this paper we

design a nonlinear model based on deep learning techniques to be included in the MPC-P and we analyze its prediction capabilities. The proposed model is then compared to the linearized average model (AVG) currently used in the MPC-P.

#### 70 2.1. Solutions based on Neural Networks

Recently, a few solutions exploiting deep learning techniques for glucose level prediction in diabetic patients have been proposed [44, 45, 46, 47, 48]. In our knowledge, the first solution exploiting neural networks for modeling the BG metabolism of a T1D patient was proposed in [49]. In particular, the authors tries to predict the glucose level of a diabetic patient by training a Recurrent Neural Network (RNN) architecture which receives insulin levels, meals and level of exercise as inputs, alongside current and

previous estimates of BG. However, the data used for both train and test is acquired from a single patient and this may result in a lack of generalization for the final model. Similarly, Allam et. al. [44], propose to use CGM signals to train a RNN for predicting future values of the glucose concentration, considering several prediction horizons. Again, the data used for both train and test comes from the same population, which may result in a model that hardly generalize to new unseen data.

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LSTM networks achieve state-of-the-art performance in modeling several time-dependent phenomena. For this reason, the authors of [45] propose to exploit LSTM in a model which takes CGM values, insulin dosages and carbohydrate intake as inputs and tries to predict the glucose level at prediction horizons of 30 and 60 minutes. Data incoming from four patients acquired using different CGM devices has been used in both train and test. Unfortunately, the training needed to be repeated multiple times, mainly because of initialization issues which left the optimization stuck in a bad local optima.

An LSTM-based architecture has also been exploited in [46]. In this case, the model is trained on the measurements provided by CGM systems, and used to predict a singular value after a pre-defined prediction horizon. The output is modeled as a univariate Gaussian distribution, so as to be able to follow the uncertainty of the prediction. The LSTM dimension is set to 128 and it is trained on the Ohio T1DM Dataset [50], considering the first 80% of the glucose level as training data for each patient, and validating on the last 20%. A more complex architecture was designed by Sun et. al. [47]. In particular, they propose to use a sequential model with one LSTM layer, one bidirectional LSTM layer and several fully connected layers to predict BG levels for different prediction horizons. The model is trained on the CGM measurements of both *in silico* and *in vivo* data coming from 20 real patients.

Convolutional RNNs have also been exploited to predict the BG level [48]. The concatenated time series of glucose level, carbohydrate and insulin is firstly preprocessed by a deep convolutional networks, so that the recurrent LSTM layers accepts these features instead of the CGM measurements directly. The model is trained on *in silico* data consisting on a small sample of 10 adult T1D subjects simulated using the UVA/Padova simulator.

#### 3. Method

Glucose concentration depends mainly on the injected insulin and carbohydrate intake, which have opposite effects on glucose levels. Indeed, it is well-known from physiology that an increase in insulin results in a decrease in glucose concentration, while a meal intake produces a glucose rise. Of course, the future evolution of the BG is also influenced by its current value and it is consistent with its trend.

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Thus, the inputs of the model proposed in this work is composed by three measurable signals sampled at a given  $T_s$ : the injected insulin (ins) recorded by subcutaneous insulin pump, the carbohydrate amount (cho) inserted manually by the patient, and the glucose concentration (cgm) measured by the CGM sensor.

The output of the model is the interstitial fluid glucose concentration (ig). Specifically, the signal cgm is the interstitial (i.e. subcutaneous) glucose concentration measured by a CGM device and affected by measurement noise, while ig is the real interstitial glucose. Each signal is rescaled, independently, using the minimum and maximum values and then subdivided in samples of fixed size, depending on the prediction horizon (ph) in analysis. These sub-samples constitute the training and testing data for our model, as detailed in Section  $\mathbf{4}$ .

Denoting the current time with  $t_0$  and given  $ph \in PH$ , where  $PH = [5, 10, \dots, 60]$  is the set of the

considered prediction horizons, let's define the following signals:

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$$\begin{aligned} \overleftarrow{\mathbf{cgm}}(t_0, ph) &= \left[ \operatorname{cgm}(t_0 - ph), \operatorname{cgm}(t_0 - ph + 1), \cdots, \operatorname{cgm}(t_0 - 1) \right], \\ \overleftarrow{\mathbf{ins}}(t_0, ph) &= \left[ \operatorname{ins}(t_0 - ph), \operatorname{ins}(t_0 - ph + 1), \cdots, \operatorname{ins}(t_0 - 1) \right], \\ \overleftarrow{\mathbf{cho}}(t_0, ph) &= \left[ \operatorname{cho}(t_0 - ph), \operatorname{cho}(t_0 - ph + 1), \cdots, \operatorname{cho}(t_0 - 1) \right], \\ \overrightarrow{\mathbf{ins}}(t_0, ph) &= \left[ \operatorname{ins}(t_0), \operatorname{ins}(t_0 + 1), \cdots, \operatorname{ins}(t_0 + ph - 1) \right], \\ \overrightarrow{\mathbf{cho}}(t_0, ph) &= \left[ \operatorname{cho}(t_0), \operatorname{cho}(t_0 + 1), \cdots, \operatorname{cho}(t_0 + ph - 1) \right], \\ \mathbf{\hat{ig}}(t_0, ph) &= \left[ \operatorname{ig}(t_0), \operatorname{ig}(t_0 + 1), \cdots, \operatorname{ig}(t_0 + ph - 1) \right] \end{aligned}$$
(1)

where  $\overleftarrow{cgm}(t_0, ph)$ ,  $\overleftarrow{ins}(t_0, ph)$ , and  $\overleftarrow{cho}(t_0, ph)$  are the CGM data, the delivered insulin and the ingested carbohydrates in the past ph minutes, respectively, while  $\overrightarrow{ins}(t_0, ph)$ ,  $\overrightarrow{cho}(t_0, ph)$  are the suggested amount of insulin and the meal information in the future ph minutes. For each ph a single model is identified as described in Section 3.1. The aim of each model is to depict the relation between ig in the future phminutes  $(\hat{ig}(t_0), \hat{ig}(t_0 + 1), \cdots, \hat{ig}(t_0 + ph - 1))$ , collected in the vector  $\hat{ig}$ , and the above mentioned signals. In particular, each single  $\hat{ig}$  value can be described as:

$$\hat{ig}(t_0+k,ph) = g\left(\overleftarrow{cgm}(t_0,ph),\overleftarrow{ins}(t_0,ph),\overleftarrow{cho}(t_0,ph),\overrightarrow{ins}(t_0,ph),\overrightarrow{cho}(t_0,ph)\right) \quad k = 0, 1, 2, \cdots, ph-1.$$
(2)

In the perspective of employing our solution in an MPC and in order to be able to accurately predict a glucose trend, an ensemble of these models can be trained, independently for each ph, and the predictions from these models can be combined to obtain a trend of future glucose concentration:

$$\hat{\mathbf{ig}}(t_0) = [\hat{\mathbf{ig}}(t_0, 5), \hat{\mathbf{ig}}(t_0 + 1, 5), \cdots, \hat{\mathbf{ig}}(t_0 + 4, 5), \\ \hat{\mathbf{ig}}(t_0 + 5, 10), \hat{\mathbf{ig}}(t_0 + 6, 10), \cdots, \hat{\mathbf{ig}}(t_0 + 9, 10), \\ \vdots \\ \hat{\mathbf{ig}}(t_0 + 54, 60), \hat{\mathbf{ig}}(t_0 + 55, 60), \cdots, \hat{\mathbf{ig}}(t_0 + 59, 60)].$$

$$(3)$$

#### 3.1. Proposed Architecture

We chose a simple model based on stacked Long Short-Term Memory (LSTM) cells [16, 51]. LSTMs are a special kind of RNNs, which are able to learn how to filter (e.g. forget) part of their hidden state during



Figure 1: (a) The basic structure of an LSTM cell. For each arrow pointing to a circle, an addition is performed. Dots represent vector/matrix multiplications. (b) Temporal unfolding and data flow on n stacked LSTM cells.

#### <sup>120</sup> the inference process in order to model long-term temporal dependencies.

Formally, a single LSTM cell with input  $\mathbf{x}(t)$ , output  $\mathbf{h}(t)$  and an internal cell state  $\mathbf{c}(t)$  is described by the following equations, also represented in graphical form in Figure 1(a):

$$\mathbf{c}_{in}(t) = \tanh(\mathbf{W}_{xc}\mathbf{x}(t) + \mathbf{W}_{hc}\mathbf{h}(t-1) + \mathbf{b}_{c})$$

$$\mathbf{i}(t) = \operatorname{sigmoid}(\mathbf{W}_{xi}\mathbf{x}(t) + \mathbf{W}_{hi}\mathbf{h}(t-1) + \mathbf{b}_{i})$$

$$\mathbf{f}(t) = \operatorname{sigmoid}(\mathbf{W}_{xf}\mathbf{x}(t) + \mathbf{W}_{hf}\mathbf{h}(t-1) + \mathbf{b}_{f})$$

$$\mathbf{o}(t) = \operatorname{sigmoid}(\mathbf{W}_{xo}\mathbf{x}(t) + \mathbf{W}_{ho}\mathbf{h}(t-1) + \mathbf{b}_{o})$$

$$\mathbf{c}(t) = \mathbf{f}(t)\mathbf{c}(t-1) + \mathbf{i}(t)\mathbf{c}_{in}(t)$$

$$\mathbf{h}(t) = \mathbf{o}(t)\tanh\mathbf{c}(t)$$
(4)

where each weight matrix  $\mathbf{W}_x$ ,  $\mathbf{W}_h \in \mathbb{R}^{d \times d}$  and  $\mathbf{b}$ ,  $\mathbf{x}(t)$ ,  $\mathbf{h}(t)$ ,  $\mathbf{c}_{in}(t)$ ,  $\mathbf{i}(t)$ ,  $\mathbf{o}(t)$ ,  $\mathbf{c}(t) \in \mathbb{R}^d$  while d represent the LSTM *dimension*, an hyperparameter defined upfront by design and constant among all cells. Respectively,  $\mathbf{i}(t)$ ,  $\mathbf{f}(t)$ ,  $\mathbf{o}(t)$  are called the *input*, *forget* and *output* gates, while  $\mathbf{c}_{in}(t)$  contains a vector of new candidate values for the cell state.

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During temporal unfolding, both  $\mathbf{h}(t)$  and  $\mathbf{c}(t)$  are passed to the temporal replica of the next cell in the fold. Models made of multiple, stacked LSTM cells can be easily conceived, by making the output of a given cell the input of the next one in the stack. The process of training through unfolding *n*-stacked LSTM cells is illustrated in Figure 1(b).

We trained multiple models, one for each  $ph \in PH$ . Depending on ph, the whole signal is sampled as <sup>130</sup> described in Eq. (1) and each sub-sample is split into two arrays  $\overleftarrow{\mathbf{X}}$  and  $\overrightarrow{\mathbf{X}}$ , the former representing past



Figure 2: Deep Glucose Forecasting (DGF) architecture. The input is split into two sets: past observations and estimated future inputs. Both branches are processed by n-stacked LSTM cells with dimension d. The output of the branches is concatenated into a final fully connected (FC) layer.

information given to the model, the latter representing the suggested therapy and meals for the future:

$$\begin{aligned} \overleftarrow{\mathbf{X}}(t_0, ph) &= \left[ \begin{array}{c} \overleftarrow{\mathbf{cgm}}(t_0, ph) \\ \overleftarrow{\mathbf{ins}}(t_0, ph) \\ \overleftarrow{\mathbf{cho}}(t_0, ph) \end{array} \right], \quad \overrightarrow{\mathbf{X}}(t_0, ph) = \left[ \begin{array}{c} \overrightarrow{\mathbf{ins}}(t_0, ph) \\ \overrightarrow{\mathbf{cho}}(t_0, ph) \end{array} \right] \end{aligned} \tag{5}$$

 $\overleftarrow{\mathbf{X}}$  and  $\overrightarrow{\mathbf{X}}$  are separately processed through two identical branches of the architecture, each being a stack of *n* LSTM cells. The output of both branches is then concatenated and processed through a final fully connected layer that produces the intended output. As the model aims to forecast the IG signals, the supervised architecture assumes to have access to the IG signal during training in order to use them as ground truth.

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More formally, leaving out the flowing of the internal cell states, the model is described by:

$$\begin{aligned} \mathbf{\widehat{h}}_{1}(t_{0}, ph) &= \mathrm{LSTM}_{1}(\mathbf{\widehat{X}}(t_{0}, ph)) & \mathbf{\widehat{h}}_{1}(t_{0}, ph) &= \mathrm{LSTM}_{1}(\mathbf{\widehat{X}}(t_{0}, ph)) \\ \mathbf{\widehat{h}}_{2}(t_{0}, ph) &= \mathrm{LSTM}_{2}(\mathbf{\widehat{h}}_{1}(t_{0}, ph)) & \mathbf{\widehat{h}}_{2}(t_{0}, ph) &= \mathrm{LSTM}_{2}(\mathbf{\widehat{h}}_{1}(t_{0}, ph)) \\ \vdots & \vdots & \vdots \\ \mathbf{\widehat{h}}_{n}(t_{0}, ph) &= \mathrm{LSTM}_{n}(\mathbf{\widehat{h}}_{n-1}(t_{0}, ph)) & \mathbf{\widehat{h}}_{n}(t_{0}, ph) &= \mathrm{LSTM}_{n}(\mathbf{\widehat{h}}_{n-1}(t_{0}, ph)) \\ \mathbf{\widehat{ig}}(t_{0}, ph) &= \mathbf{W}_{FC} \left[\mathbf{\widehat{h}}_{n}(t_{0}, ph) & \mathbf{\widehat{h}}_{n}(t_{0}, ph)\right] + \mathbf{b}_{FC} \end{aligned}$$
(6)

where  $\mathbf{W}_{FC} \in \mathbb{R}^{d \times ph}$ ,  $\mathbf{b}_{FC} \in \mathbb{R}^{ph}$  and  $\mathrm{LSTM}_n$  represents the *n*-th LSTM layer in the stack and it is described by Eq. (4).

The training process uses a Mean Squared Error loss function (MSE) with a default Adam optimizer (learning rate  $10^{-3}$ , 180 epochs), so that for each sample:

MSE := 
$$\frac{1}{ph} \sum_{t=t_0}^{t_0+ph} (\hat{\mathbf{ig}}(t, ph) - \mathbf{ig}(t, ph))^2$$
 (7)

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# Model hyperparameters were determined by a standard grid-search by using a separate set of validation data extracted by the same distribution of the training data, as detailed in Section 4.1. The complete solution is shown in Figure 2 and from now on we will refer to it as therapy-driven Deep Glucose Forecasting (DGF).

#### 4. Experiments

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In this section we report on a series of experiments in order to assess the performance of the proposed solution. We first describe the datasets used in our experiments, then we discuss the parameters settings and the measures used to evaluate our approach. In Section 4.3 we analyze how the performance of DGF varies by considering different configurations for the dimension d of each LSTM and the number n of stacked LSTMs, in order to identify the best combinations of these hyperparameters. Finally, we evaluate the prediction capabilities of the proposed solution on *in vivo* data collected during clinical trials [4].

#### 4.1. Dataset

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The *in silico* dataset has been generated using the UVA/Padova simulator [29, 30], which is equipped with a cohort of virtual patients and accepted by Food and Drug Administration (FDA) as a substitute to animals trials. This acceptance allowed the *in silico* synthesis of control algorithms directly testable on real patients. The UVA/Padova simulator includes a large nonlinear compartmental model able to simulate the glucose-insulin dynamics of the diabetic population [30]. The inter-subject variability of this population is modelled through different sets of metabolic parameters of this model.

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The simulator is equipped with three virtual populations (children, adolescents and adults), each composed of 100 subjects. In the most recent version of the simulator, the circadian variability of insulin sensitivity and meal absorption parameters have been added [52, 53, 54]. The UVA/Padova simulator allows also to simulate the so-called meal announcement, i.e. the patient can announce a meal intake to the controller in advance.

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In this paper, the population of 100 adults of the UVA/Padova simulator is used and two different scenarios are designed. Table [] shows Scenario 1, which is a four-day protocol simulated in closed-loop using the MPC-P to define the optimal insulin therapy. The first three days are used for model training, while the remaining day is used for validation purposes. The training scenario involves three meals per day with additional snacks in each day. Moreover, in order to define a real-life scenario, possible errors in the meal announcement are included, i.e. a limited events of unannounced meals or meals announced

|  |                   | Time   | CHO (g) | Insulin Bolus  |                            |   |  |                            |  |
|--|-------------------|--|---------|--|----------------------------|---|--|----------------------------|--|
|  |                   | 08:00  | 60      | Bolus on time  |                            |   |  |                            |  |
|  | y 1               | 13:00  | 60      | Bolus at 14:00   |                            | <i>T</i> T:   |  | Level's Delve              |  |
|  | Da                | 17:00  | 30      | No bolus   |                            | 1 ime   | CHO (g)  | Insuin Boius               |  |
|  |                   | 20:00  | 80      | Bolus on time  |                            | 08:00   | 50   | Bolus on time              |  |
|  |                   | 08.00  | 50      | Bolus on time  | y 1                        | 13:00   | 50   | Bolus on time              |  |
|  |                   | 08.00  |         | Bolus on time  | Da                         | Time         CHO (g)         Insuli           08:00         50         Bolus           13:00         50         Bolus           19:00         70         Bolus           23:00         20         Bolus           09:30         15         No           13:00         60         Bolus           19:00         70         Bolus           23:00         20         Bolus           10:00         60         Bolus at 12           17:00         25         Bolus           20:00         90         Bolus at 22           20:00         90         Bolus at 23           11:30         20         Bolus           11:30         20         Bolus           11:30         20         Bolus           11:30         20         No           20:30         100         Bolus           11:30         20         No           20:30         100         Bolus | Bolus on time  |                            |  |
|  | 5                 | Time           08:00           13:00           20:00           08:00           17:00           20:00           08:00           10:00           13:00           10:00           13:00           10:00           13:00           10:00           22:00           06:30           09:30           12:30           17:00           20:00           23:00           08:00           11:30           16:30           20:30 | 15      | No bolus   |                            | 23:00   | 20   | Bolus on time              |  |
|  | Day               | 13:00  | 35      | Bolus on time  |                            | 06:30   | 50   | Bolus on time              |  |
|  | _                 | 19:00  | 80      | Bolus on time  |                            | 00.30   | 50   | Dolus on time              |  |
|  |                   | 22:00  | 20      | Bolus on time  |                            | 09:30   | 15   | No bolus                   |  |
|  |                   | 06:30  | 40      | Bolus on time  | No.                        | 13:00   | 60   | Bolus at 12:00 for 50 g    |  |
|  |                   | 00.30  | 40      | Dorus on time  | D<br>D                     | 17:00   | 25   | Bolus on time              |  |
|  | ~                 | 09:30  | 20      | No bolus   |                            | 20:00   | 90   | Bolus at 20:15 for 70 g    |  |
|  | ay :              | 12:30  | 45      | Bolus at $12:00$ for $50$ g  |                            | 23.00   | 15   | No bolus                   |  |
|  | Ä                 | 17:00  | 20      | Insulin Bolus         Bolus on time         Bolus at 14:00         No bolus         Bolus on time         Bolus at 12:00 for 50 g         Bolus at 12:00 for 50 g         Bolus at 12:00 for 50 g         Bolus on time         Bolus at 12:00 for 30 g         No bolus         Bolus on time         Bolus on time         Bolus at 13:30 for 30 g         No bolus         Bolus on time         Bolus at 13:30 for 30 g         No bolus         Bolus on time         Bolus | 50                         | D l   |  |                            |  |
|  |                   | 20:00  | 70      | Bolus at 20:30   |                            | 08:30   | CHO (g)Insulin Bolu50Bolus on tim50Bolus on tim70Bolus on tim20Bolus on tim20Bolus on tim50Bolus on tim50Bolus on tim15No bolus60Bolus at 12:00 fo25Bolus at 12:01 fo15No bolus50Bolus at 20:15 fo15No bolus50Bolus on tim20Bolus on tim60Bolus at 13:00 fo20No bolus100Bolus on timCable 2: Testing Scenario. | Bolus on time              |  |
|  |                   | 23:00  | 20      | No bolus   | с<br>С                     | 11:30   | 20   | Bolus on time              |  |
|  |                   | 20.00  | 20      | Dl   | Day                        | 14:00   | 60   | Bolus at 13:00 for 30 g $$ |  |
|  | Day 4 Day 3 Day 2 | 08:00  | 35      | Bolus on time  |                            | 17:00   | 20   | No bolus                   |  |
|  | 4                 | 11:30  | 20      | Bolus on time  |                            | 20:30   | 100  | Bolus on time              |  |
|  | Jay               | 13:30  | 60      | Bolus at 13:30 for 30 g $$   |                            | 20100   | 100  | Donab on vinite            |  |
|  | Ι                 | 16:30  | 20      | No bolus   | Table 2: Testing Scenario. |   |  |                            |  |
|  |                   | 20:30  | 90      | Bolus on time  |                            |   |  |                            |  |

Table 1: Training Scenario.

with a wrong estimation of the amount. Scenario 2 lasts three days and it is reported in Table 2. The meal amounts and times of this protocol are designed to reproduce a real life scenario. Hypo treatments of 15 g are administrated to the patient in case glucose concentration falls below 65 mg/dl in both scenarios. Scenario

- 1 is used to perform model training, while Scenario 2 is exploited to assess the prediction capabilities of 170 the proposed model. The design of two different scenarios aims to simulate the randomness of the human behaviour regarding the variability of day to day eating habits. Specifically, Scenario 2, i.e. the testing scenario, is defined to reproduce eating patterns different from those present in Scenario 1, i.e. the training scenario. Moreover, an in vivo dataset is considered and it is composed of clinical data of a single T1D
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patient of the Padova clinical centre collected during experiments involved in the AP@Home project 5. The considered clinical trial lasted for a month, and it has been conducted through a fully automatic closedloop control 4. The closed-loop system was composed of an suitably modified android smartphone (the DiAs platform [38]), communicating wirelessly with the G4 Platinum CGM system, Dexcom Inc. and the AccuCheck Spirit Combo insulin pump, Roche Diagnostic. This dataset is challenging because the clinical

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of restriction.

trial has been conducted in free-living conditions, i.e. the patient could have a normal life without any type

#### 4.2. Parameter Settings and Evaluation Protocol

The accuracy of the model predictions is assessed by considering various prediction horizons, which are expressed in terms of minutes. In this paper, ph from 5 minutes to 60 minutes are considered. Since the proposed model is aimed to be included in the MPC-P controller, that is characterized by a sample time of  $T_s = 5$  minutes for the predictions, we consider PH =[5, 10,  $\cdots$ , 60] and the predicted signals are sampled every  $T_s$ . For a given patient p and a specific  $ph \in$  PH, we denote with  $\hat{ig}(\cdot, ph)$  the ph-steps ahead prediction on the entire testing scenario,  $ig(\cdot)$  the considered reference in Scenario 2, and ig its average value. The predictions of the model are evaluated in terms of Coefficient Of Determination (COD), FIT, and Root Mean Square Error (RMSE) defined as follows:

$$COD_{p}(ph) = 100 * \left(1 - \frac{||\widehat{\mathbf{ig}}(j,ph) - \mathbf{ig}(j)||^{2}}{||\mathbf{ig}(j) - \mathbf{ig}||^{2}}\right)$$
$$FIT_{p}(ph) = 100 * \left(1 - \frac{||\widehat{\mathbf{ig}}(j,ph) - \mathbf{ig}(j)||}{||\mathbf{ig}(j) - \mathbf{ig}||}\right)$$
$$RMSE_{p}(ph) = \frac{1}{N_{\text{sample}}} ||\mathbf{ig}(j) - \mathbf{ig}(j,ph)||$$
(8)

where  $j = ph, ph + T_s, \dots, N_{\text{sample}} \cdot T_s$  is used to index the considered samples, and  $N_{\text{sample}}$  is the number of samples of the signal when sampled every  $T_s$  minutes for Scenario 2. COD and FIT are equal to 100% for perfect predictions and can reach negative values in case of bad performances. The average value of each metric ( $\overline{COD}, \overline{FIT}, \overline{RMSE}$ ) for all PH is used as main outcome to evaluate the overall performance as follows:

$$\overline{COD} = \frac{1}{N_{ph}} \sum_{i=1}^{N_{ph}} \left( \frac{1}{N_p} \sum_{p=1}^{N_p} COD_p(ph(i)) \right)$$
$$\overline{FIT} = \frac{1}{N_{ph}} \sum_{i=1}^{N_{ph}} \left( \frac{1}{N_p} \sum_{p=1}^{N_p} FIT_p(ph(i)) \right)$$
$$\overline{RMSE} = \frac{1}{N_{ph}} \sum_{i=1}^{N_{ph}} \left( \frac{1}{N_p} \sum_{p=1}^{N_p} RMSE_p(ph(i)) \right)$$
(9)

where  $N_{ph}$  is the total number of considered  $ph \in PH$ , i.e.  $N_{ph} = 12$ , and  $N_p$  is the total number of patients involved in each testing experiment.

#### 185 4.3. Ablation Study

In order to have a comprehensive idea of the model behaviour, an ablation study is performed to assess the model's structure and the function of its different components. Specifically, the goal of this study is to evaluate the contribution of the model components and to observe how these affected the predictive capabilities.

|              |        | $\overline{COD}$ (%) |       |       | $\overline{FIT}$ (%) |       |       | RMSE  |       |       |
|--------------|--------|----------------------|-------|-------|----------------------|-------|-------|-------|-------|-------|
|              | n<br>d | 1                    | 2     | 3     | 1                    | 2     | 3     | 1     | 2     | 3     |
| 0            | 16     | 79.48                | 79.27 | 79.12 | 55.93                | 55.59 | 55.40 | 12.57 | 12.65 | 12.72 |
| ILIC         | 32     | 81.31                | 81.14 | 79.60 | 57.94                | 57.12 | 56.27 | 11.93 | 11.79 | 12.41 |
| $\mathbf{S}$ | 64     | 82.10                | 81.93 | 81.32 | 58.84                | 58.66 | 57.98 | 11.68 | 11.72 | 11.93 |
|              | 16     | 67.67                | 71.92 | 71.61 | 45.03                | 48.17 | 47.92 | 29.09 | 27.43 | 27.56 |
| /IVC         | 32     | 67.20                | 72.57 | 72.05 | 44.96                | 48.93 | 48.65 | 29.13 | 27.03 | 27.18 |
| -            | 64     | 62.06                | 71.70 | 72.50 | 41.41                | 48.44 | 49.17 | 31.01 | 27.29 | 26.91 |

Table 3: Performance metrics on each combination of d - n.

In order to maximize the generalization capability of the proposed algorithm, we train the model on different patients with eating habits drastically different with respect to those observed in the testing scenario. To do so, the population of 100 adults is split in two parts  $(N_p = 50)$ : the model is trained on the first  $N_p$  patients in Scenario 1, and the tests are conducted on the second half of the patients in Scenario 2. The same is performed but considering the other half of patients in each scenario. The final results are obtained by averaging the estimates from the two different train and test groups. This data separation has 195 been chosen in order to test the capability of the model to represent subjects not belonging to the training set but also to check the model robustness against a variation in meal sizes and correlation of meal sizes between a day.

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Firstly, the study focuses on the choice of both the size of the hidden units in each layer  $(d \in \{16, 32, 64\})$ and the number of LSTM layers  $(n \in \{2,3\})$ . From Table 3 it is possible to observe that increasing the number of hidden units for each LSTM entails a slight improvement of the performance indices while the number of stacked LSTM does not affect the final performance. This may be due to the fact that the amount of data and its variability are not sufficient to train a model with a larger number of parameters. For this reason all the subsequent experiments have been performed considering two stacked LSTM with d = 64. Higher values for d did not produce any significant improvement.

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Secondly, we analyze how the different features considered as input for the network influence the final performance. For this reason, we removed past insulin  $(\overleftarrow{ins})$  and carbohydrates  $(\overleftarrow{cho})$  from the input stream. The model performance in this case is evaluated both on *in silico* and *in vivo* testing data.

Denoting with  $\mathbf{X}^*$  the modified input array, the vector representing past information given to the model is defined as follows:

$$\overleftarrow{\mathbf{X}^{*}}(t_{0}, ph) = \left[\overleftarrow{\mathbf{cgm}}(t_{0}, ph)\right].$$
(10)



Figure 3: Comparison of the prediction performance on both in silico and in vivo testing data in term of COD (a), FIT (b) and RMSE (c) on Scenario 2 and real-life data.

Figure 3 shows the comparison of the prediction performance of the models with  $\left[\overleftarrow{\mathbf{X}^{*}}(t_{0}, ph) \ \overrightarrow{\mathbf{X}}(t_{0}, ph)\right]$ and  $\left| \overleftarrow{\mathbf{X}}(t_0, ph) \ \overrightarrow{\mathbf{X}}(t_0, ph) \right|$  as inputs, respectively. It denotes that the introduction of the information regarding the past insulin therapy and ingested meals guarantees an improvement in the prediction performance of the model. In particular, considering the results obtained by using *in silico* data, there is a slight improvement in performance. Indeed, the virtual patients belonging to the training and testing groups are subsets of the same population.

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Since the LSTM is able to learn the behaviour of the population, the additional information about past history does not provide a significant improvement. On the other hand, Figure  $\overline{\beta}$  shows a significant gap in performance computed on real life data. Indeed, if the LSTM is trained on in silico data, the lack of past therapy information lowers the performance on *in vivo* testing dataset. Hence, the past evidence ins and the help mitigating the differences in the data distribution. The model obtained considering these additional information is able to generalize to new unseen data and improve the overall glucose control. 220

5. Discussion

The proposed solution is a population average model identified on the 100 adults of the UVA/Padova simulator. An average model could ideally limit the performance since it describes the average dynamics of the population, so we aim to test both its prediction and generalization capabilities on a dataset different from the one used in training. Hence, the proposed algorithm is tested on a 1-month dataset containing all the data collected during the clinical trial [4] for a single patient. This dataset is challenging because it includes eating patterns not present in the training dataset, but also it includes all the problems experienced during the clinical trial.

The performance obtained by the DFG model and the linearized average model (AVG) of the UVA/Padova

| Predictor                       | COD (%) | FIT (%) | RMSE  |
|---------------------------------|---------|---------|-------|
| AVG                             | 15.23   | 11.48   | 46.82 |
| DGF                             | 69.85   | 48.44   | 27.29 |
| DMP <b>13</b>                   | 80.79   | 61.44   | _     |
| DGF Fine Tuned                  | 73.23   | 49.69   | 26.63 |
| DGF Fine-tuned + Exp. Filtering | 84.05   | 60.14   | 21.09 |

Table 4: Prediction matrices.

simulator are reported in Table 4. By considering the first two rows of Table 4. the DFG model shows su-230 perior prediction performance against the AVG. Moreover, DGF approach proves to be able to generalize over different datasets by achieving interesting improvements with respect to AVG, despite being an average model.

#### 5.1. Fine tuning

The drawback of an average model is that it can not fully describe the variety of individual glucose 235 response of the entire population. The definition of an individualized insulin therapy by exploiting a patienttailored model can substantially improve the effectiveness of the glucose control. Hence, in order to improve the DGF model performance, the LSTM trained on *in silico* data is fine-tuned on data of the specific patient. In this context, Fine-Tuning (FT) slightly modifies the weights of the LSTM in order better fit the individual behaviour of the real patient. For the purpose of this paper, we have decided to extend the 240 partial retraining entailed by any FT to the entire architecture, by using a suitably small learning rate  $(10^{-5})$  for 100 additional epochs on the FT dataset. A filtered version of the collected CGM data  $(ig_R)$  is used as reference for the signal ig. The signal  $ig_R$  is obtained by using a retrofitting algorithm, which is able to reconstruct an accurate continuous-time BG profile by exploiting BG samples from the fingerstick and CGM data from the sensor 56. In order to provide a good amount of information to describe the dynamic 245 of a specific patient, the FT dataset contains two days picked up randomly among the available ones.

The FT technique improves the accuracy by 6% on the 2-layer stacked LSTM, as reported in the row 4 of Table 4. Figure 4 reports the performance metrics as a function of ph for the 2-layer stacked LSTM with in vivo testing dataset. The higher the ph, the lower the performance. This is motivated by the fact that the further you want to predict the more difficult it becomes. The improvements of the fine-tuned models 250 are evident for large ph values where the performance increases with respect to the original model without FT. These improvements are less obvious if we compare their performance against the performance of the individualized model (MMP) presented in 13 and reported in Table 4. In 13 the individualized model is identified from real-data on the base of the a-priori knowledge acquired through the analysis of the patient

real-data, while here real-data are used to adjust the model pre-trained on in silico data. 255



Figure 4: A comparison of the prediction performance of 2-layer stacked LSM compared to its fine-tuned version in term of COD (a), FIT (b) and RMSE (c) on the clinical dataset.

#### 5.2. Output filtering

The main limitation of the proposed approach is that the network is trained on noisy input measurements but a noiseless signal is required as output. Since the measurement noise that affects the CGM data is not negligible, the network would try to reduce the noise on the output in order to improve the overall quality of the prediction. However, this effect can be limited and in order to further improve the prediction smoothness an exponential filtering is applied to the predictions.

The predicted  $ig(\cdot, ph)$  is filtered by the following exponential filter:

$$\hat{ig}_{exp}(t_0 + k, ph) = \alpha \cdot \hat{ig}(t_0 + k, ph) + (1 - \alpha) \cdot \hat{ig}_{exp}(t_0 + k - 1, ph) \qquad k = 0, 1, \cdots, ph - 1$$
(11)

where  $\alpha \in (0, 1)$  is the smoothing factor and is defined as follows:

$$\alpha = \frac{2}{w_{\exp} + 1} \tag{12}$$

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and  $w_{exp}$  is the length of the window used by the filter. It is set to 5, i.e. the minimum window observed by the model and it is kept fixed for all models. Figure 5 shows the noisy CGM data, the output of the retrofitting algorithm  $ig_R$ , which represents the ground truth, and the output of the exponential filter  $ig_{exp}$ . It may be noted in Figure 5 that there is a scaling problem in the signal  $ig_{exp}$ . This effect can be explained by considering that  $ig_{exp}$  is rescaled by using the minimum and maximum training values of **ig**. These values are computed with the 100 adults of the UVA/Padova simulator and they are different from the minimum and maximum values computed on the entire clinical trial of the selected patient. A compromise solution could be to use the minimum and maximum values of the FT dataset. Therefore it is not a suitable solution

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because this means to neglect the well-known inter-day variability. In order to cope with this problem, a larger dataset of the patient is required to capture the individual variability without compromising the testing dataset. The last row of Table 4 shows the results obtained by applying the exponential filter to



Figure 5: Glucose profile of a 24h study-case: the yellow line is the noisy CGM data, the light blue line is the output of the retrofitting algorithm  $ig_R$  presented as reference, and the red line is the output of the exponential filter  $ig_{exp}$ .

the output of the 2-layer stacked LSTM model. It is evident that the filtering technique is able to highly improve the performance by partially removing the noise that affects the CGM data.

#### 275 6. Conclusions

A solution based on deep learning to predict a trend of future glucose concentration is proposed. The models are trained by using the 100 *in silico* adult patients of the UVA/Padova simulator. Each model is used to predict a glucose profile for a fixed prediction horizon and the individual predictions are then aggregated to obtain a profile of future glucose levels.

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In order to assess the generalization ability, the models have been tested on real data collected during a 1-month clinical trial in free-living conditions. The achieved results show that the proposed approach can significantly improve predictive performance despite being an average model. In order to individualize the trained models, a fine-tuning is applied to each model separately considering a really small portion of the specific patient. Satisfactory results have been obtained in terms of prediction capabilities. We plan to investigate more in details how the data affect the model training and to improve the individualized models by exploiting a larger amount of patient data.

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