Optimal Insulin Dosing for Glucose Control in a Virtual Type-I Diabetes Patient through Reinforcement Learning

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Introduction

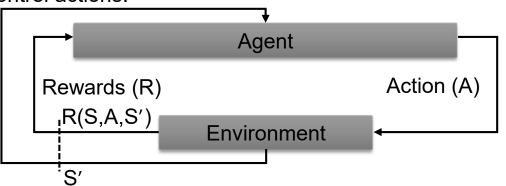
- Type 1 Diabetes Mellitus (T1DM): The body's immune system destroys β-cells, eliminating insulin production from the body
- T1DM patient depends on the exogenous insulin dosages
- Open loop control comprising multiple daily insulin injections generally leads to poor glycaemic control
- Closed loop control using an Artificial pancreas device system (APDS) is desired. APDS consists of:
 - Continuous glucose monitoring sensor (CGMS)
 - Controller that estimates insulin to be dosed based on glucose and other measurements
 - Insulin pump

Reinforcement learning and its application in APDS

- Uncertainty associated with external disturbances such as amount of meal, physical activity^{1,2}
- Inter and intra patient variability in glucose metabolism
- RL framework can account for such unexpected disturbances and individualize insulin dosing

State (S)

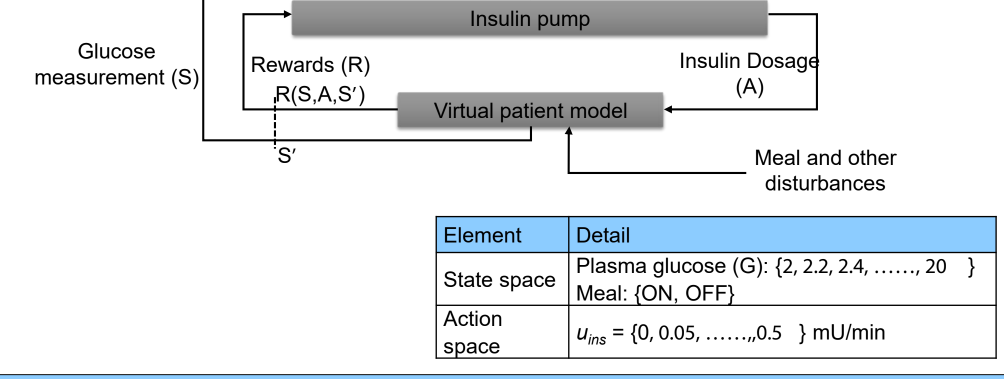
- RL algorithms that can help agent learn optimal control actions:
 - Model-based
 - Model-free algorithms
 - 1. Q-learning
 - 2. Dyna-Q
 - 3. $Q(\lambda)$



Mathematical model of a virtual diabetes patient³

- This model is used previously for in-silico testing of various control algorithms
- Glucose enters via:
 - Intestinal absorption through meals
 - Hepatic glucose production
- Glucose is removed via:
 - Utilization in RBC
 - Insulin-dependent glucose utilization in the liver
 - Glucose excretion takes place above the renal threshold
- In T1DM patient, only source of insulin is through APDS.
- Coupled model has 3 ordinary differential equations and 4 algebraic equations

Closed loop Type-1 glucose control problem



Online RL control algorithms



$$Q(S_{t}, A_{t}) = Q(S_{t}, A_{t}) + \alpha \left[R_{t+1} + \gamma \frac{arg \max}{a} Q(S_{t+1}, a) - Q(S_{t}, A_{t}) \right] \qquad \pi_{t}(S_{t}) = \begin{cases} P\left(\frac{arg \max}{a} Q(S_{t}, a)\right) = 1 - \epsilon \\ P\left(A \neq \frac{arg \max}{a} Q(S_{t}, a)\right) = \frac{\epsilon}{2} Q(S_{t}, a) \end{cases}$$
• **Dyna-Q:**

Repeat (for each episode)

Initialize S, A

Observe R, S

Update *Model*

Model(S, A) = R, S

Update Q

Until *S* is terminal

Repeat N number of times

Sample Model(S, A)

R,S = Model(S,A)

Q(s,a) = Q(s,a) +

Update Q

Initialize Q(s, a) arbitrarily, for all $s \in S$, $a \in A$ Repeat (for each episode) Z(s,a) = 0 for all $s \in S$, $a \in A$

Initialize S, A Repeat (for each step of episode)

Take action A, observe R, S' Chose A at S' using policy derived from Q $A = argmax_a Q(S, a)$ $\delta = R + \gamma Q(S, A) - Q(S, A)$ Z(S,A) = Z(S,A) + 1For all $s \in S$, $a \in A$ $Q(s,a) = Q(s,a) + \alpha \delta Z(s,a)$ If A = A, then $Z(s,a) = \gamma \lambda Z(s,a)$

else Z(s,a) = 0

Until S is terminal

Gaussian disturbances in:

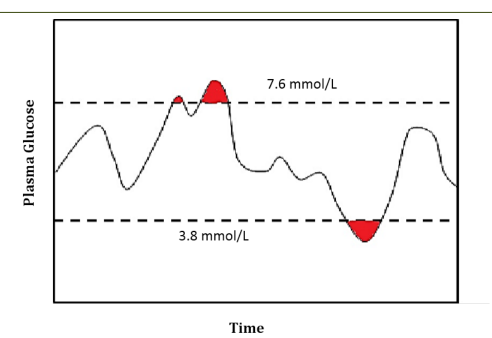
1. Amount of meal: 30% of the mean

S = S ; A = A

- 2. Time of meal: 60 min
- Control signal to the pump is updated every 10 minutes

Performance measure:

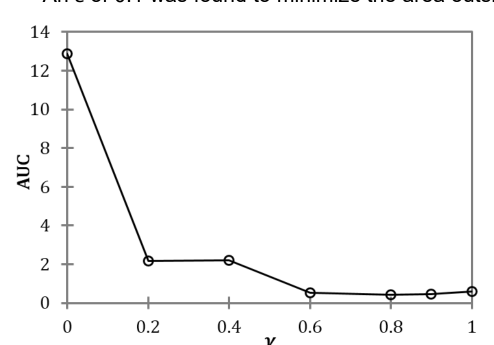
- Area under Curve (AUC) enclosed continuous glucose measurements with its upper and lower bounds of permissible limits during a 24 hour period is calculated.
- Lower AUC signifies better control.
- Average of AUC over all episodes (5000) is used to compare the performance of different algorithms

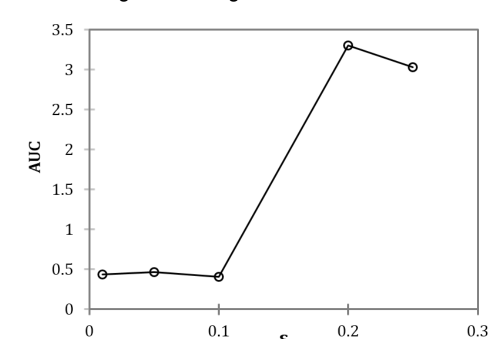


Results

Effect of tuning parameters of RL algorithm, namely, α , γ , and ϵ on the quality of control:

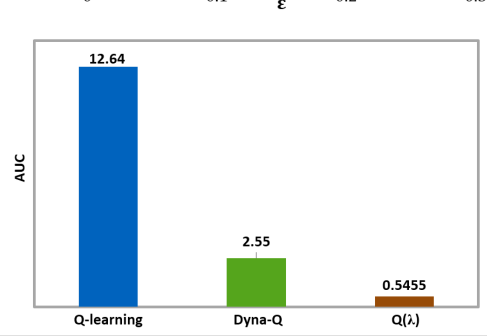
- Learning rate, α, above 0.2 yielded similar glucose control in the 5000 episodes simulated
- AUC reduced significantly on increasing discount factor γ and became steady at 0.6
- An ε of 0.1 was found to minimize the area outside the desired glucose range



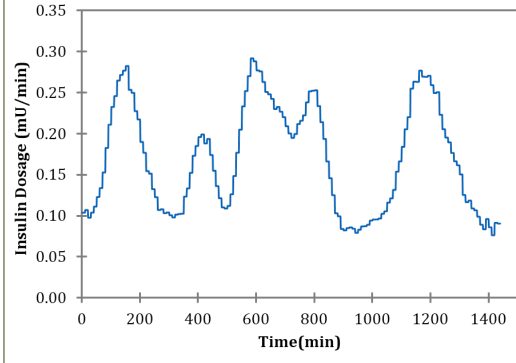


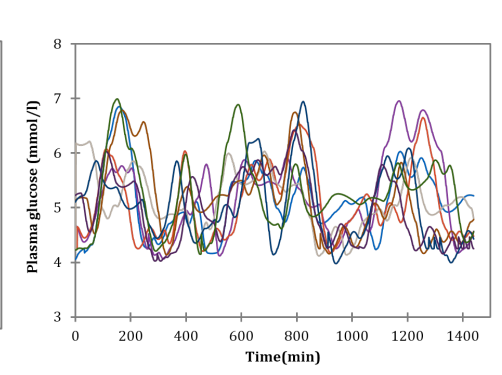
Performance comparison of RL algorithms:

- Dyna-Q which incorporates a planning agent performs better than Q-learning
- $Q(\lambda)$ outperforms both the algorithms
- Dyna-Q performance can be further improved by increasing the number of planning steps but this comes with an additional computational cost
- $Q(\lambda)$ is preferred over the other two as it performs better at relatively lower computational cost.



Application of Q(\(\lambda\)) algorithm:





- RL agent is able to control the glucose concentration during most of the days.
- Average glucose concentration is maintained between 4.5 to 5.5 mmol/L
- · Peaks in the insulin dosage plot coincides with the meal intake

Conclusion

- Reinforcement learning is a viable alternative to traditional controllers used in APDS.
- In the present study, application of RL based controllers was found to be effective in maintaining normoglycemia even in presence of disturbances in meal related inputs.
- More studies need to be performed to study the efficacy of RL based controllers when intra-day variability in a virtual-patient is present.
- Moreover, RL based controllers need to be supported with either estimation methods or clinical heuristics/rules to avoid any exploratory action that can be damaging to the patient

References

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- 3. Lehmann, E.D. and Deutsch, T., 1992. A physiological model of glucose-insulin interaction in type 1 diabetes mellitus. Journal of biomedical engineering, 14(3), pp.235-242.
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Initialize Q(s,a) and Model(s,a) arbitrarily, for all $s \in S$, $a \in A$

 $Q(s,a) = Q(s,a) + \alpha(R + \gamma argmax_a Q(S,a) - Q(S,A))$

S = select a state at random from the visited states

A = select a state at random from the visited states

 $\alpha(R + \gamma argmax_a Q(S, a) - Q(S, A))$

Repeat (for each step of episode)

Chose A at S using policy derived from Q