1 TITLE

Algumas possibilidades

- 1) Strategies of Physics-Informed Neural Networks model optimization applied to Biochemical Reactions: A case study with *Lactobacillus casei*
- 2) The effects of nondimentionalization in Physics-Informed Neural Networks applied to the Kinetics of Biological Reactions: A case study with Lactobacillus casei
- 3) Physics-Informed Neural Networks applied to the Kinetics of Biological Reactions: Batch, Fed-Batch and CSTR
- 4) The simulation of Lactic Acid production in Batch, Fed-Batch and Continuous states using Physics-Informed Neural Networks

2 ABSTRACT

A FAZER

3 INTRODUCTION

Lactic acid (LA) is a molecule of great industrial and economical interest. LA and its derivatives are used in pharmaceutical, cosmetics and food industries (NANCIB et al., 2015). Since it is an organic molecule, occurs naturally is and perceived as "green" or environmentally friendly by many consumers (DE OLIVEIRA et al., 2021), its perceived value comes not only from the molecule chemical or technological features, but also from a marketing standing point. Another point that drove interest in LA growing since the beginning of the 21th century was its application as a raw material for the production of PLA (poly-lactic acid), an environmentally friendly alternative to plastics and that has found many applications (KOMESU; MACIEL; FILHO, 2017; LI et al., 2006). The industry of Lactic Acid will grow to 160 kt by 2025 (LÓPEZ-GÓMEZ et al., 2019), with a revenue forecast of almost USD 9 billion (DIN et al., 2021).

LA has actually two optical active forms, which can vary greatly in application. The L(+) form is prefered for some cosmetical, food and pharmaceutical applications since D(-)-Lactic Acid can be a harmful enantiomer ((POHANKA, 2020)). The chemical synthesis yields a racemic mixture (BÜYÜKKILECI; HARSA, 2004), while many bacteria can produce mainly one of the two enantiomers depending on the process conditions. The great economical and scientific interest in LA is demonstrated in Academia, with many studies conducted on these kind of microorganisms and their LA production using different carbon sources were evaluated (ALTAF; NAVEENA; REDDY, 2007, 2007; ALTIOK; TOKATLI; HARSA, 2006; DEY; PAL, 2013; REZVANI; ARDESTANI; NAJAFPOUR, 2017; THAKUR; PANESAR; SAINI, 2019; VERA-PEÑA; HERNÁNDEZ-GARCÍA; VALENCIA-GARCÍA, 2022). Thus, the large amount of scientific literature available and relevance made Lactic Acid production by *Lactobacillus casei* (ALTIOK; TOKATLI; HARSA, 2006) the ideal case study for the application of Physics-Informed Neural Networks in biological reactions and reactor simulation.

Physics-Informed Neural Networks (PINN) are an approach to employ deep neural networks capability of universal function approximators to solve complex numerical problems, such as stochastic and high order partial differential equations (PDEs) (RAISSI; PERDIKARIS; KARNIADAKIS, 2019; SANTANA et al., 2022). This technique can be applied to many cases where the numeric solution is highly complex, but also for simpler cases where it provides a robust framework for simulating system with well defined mathematical models. One of the greatest advantages of PINNs is the possibility to use the mathematical models themselves instead of raw experimental data, so it is possible to work with *small data*. Biological reactions kinetics pose a natural challenge for mathematical solution because, while many models may have a relatively simple mathematical description, this descriptions often involves multiple derivatives referencing each other, limiting the methods of solving it, or even requiring greater computer processing power and more sophisticated integration methods.

This work aims to determine the relevance of the deepness or wideness of the neural network chosen and the influence of the nondimensionalization of equations in performance and computational cost required by the system to achieve enough precision. This is done while evaluating the application of PINNs in biological reaction kinetics, using as a case study the production of Lactic Acid by *Lactobacillus casei* as described in the literature (ALTIOK; TOKATLI; HARSA, 2006) in CSTR (Countinuous stirred-tank), batch and fed-bacth reactors.

4 MATERIALS AND METHODS

4.1 Nondimensionalization

Nondimensionalization is a process in which variables are converted to nondimensional variables using nondimensional factors, in this work called nondimensional scalers. These scalers can substantially improve or worsen the error function, thus are an important for giving insides on how to improve the model (ALHAMA MANTECA; SOTO MECA; ALHAMA, 2012). The reduced dimension equation may need less computation resources, produce better results or be unable to attain a feasible solution. Since PINNs can be very sensitive to the value of the derivatives they are optimizing, the equations were nondimensionalized to evaluate the procedure impact on final results

We define each variable, represented by N, as an nondimensional variable. The variable is equals to the nondimensional variable multiplied by a coefficient:

$$N = N_A * N_S \tag{1}$$

where N is the variable itself, N_A is the nondimensional variable and N_S is the nondimensional scaler. The use of the subscript A represents the nondimensional variable version of N, and S represents the nondimensional scaler of N.

4.2 Reaction Kinetics Mathematical Model

The model of the production of Lactic Acid by *Lactobacillus casei* using *whey lactose* was proposed and validated by (ALTIOK; TOKATLI; HARSA, 2006) using experimental data of a batch reactor. The second experiment variant (with starting

lactose concentration 21.4g/L and duration of 9 hours) is used in this work. *L. casei* produces Lactic Acid through the classical Monod equation adjusted to represent product inhibition effect (represented by $(1 - P/P_M)$) and biomass inhibition effect (represented by $(1 - X/X_M)$), where X is the biomass concentration, P is the product concentration, P_M is the inhibitory product concentration, and X_M is the inhibitory biomass concentration. This step was necessary because LA production by *L. casei* is known to be controlled by inhibitory effects. Thus, the biomass reaction rate model includes product, substrate and production inhibitory effects. The biomass concentration derivative over time is given by:

$$r_X = \frac{\mu_{max}S}{K_S + S} X \left(1 - \frac{X}{X_M} \right)^f \left(1 - \frac{P}{P_M} \right)^h \tag{2}$$

where r_X is the reaction rate of biomass, μ_{max} is the maximum possible growth rate, S is the substrate (whey lactose) concentration, K_S is the Monod constant and f and h are factors that represent the toxicity or inhibitory potential adjustments.

The product formation rate is given by Luedeking-Piret kinetics, depending on the growth or decrease of biomass linearly (represented by dX/dt) and also on the biomass concentration itself:

$$r_{p} = \alpha r_{x} + \beta X \tag{3}$$

where α is the growth-associated product formation coefficient and β is the non-growth associated product formation coefficient.

The substrate consumption kinetics is given by a relationship including the substrate converted to product and the substrate used for biomass maintenance:

$$r_{S} = \frac{-1}{Y_{ps}} r_{p} - m_{s} X \tag{4}$$

where Y_{PS} is the product yield coefficient and m_{s} is the maintenance coefficient.

Equations 2 - 4 can be nondimensionalized as:

$$\frac{X_{S}}{t_{S}} r_{X_{A}} = \frac{\mu_{max} S_{A} S_{S}}{K_{S} + S_{A} S_{S}} X \left(1 - \frac{X_{A} X_{S}}{X_{M}} \right)^{f} \left(1 - \frac{P_{A} P_{S}}{P_{M}} \right)^{h}$$
(5)

$$\frac{P_S}{t_S} r_{P_A} = \alpha \frac{X_S}{t_S} r_{X_A} + \beta X_A X_S \tag{6}$$

$$\frac{S_{S}}{t_{S}}r_{S_{A}} = \frac{-1}{Y_{PS}} \frac{P_{S}}{t_{S}} r_{P_{A}} - m_{S} X_{A} X_{S} \tag{7}$$

4.3 Reactor Mathematical Model

The volume of liquid in the reactor is given by:

$$\frac{dV}{dt} = f_{\rm in} - f_{\rm out} \tag{8}$$

Which can be nondimensionalized as:

$$\frac{V_S}{t_S} \frac{dV_A}{dt_A} = f_{\rm in} - f_{\rm out} \tag{9}$$

The nondimensional concentration of a generic substance or biomass, N, can be given by:

$$V_{A}V_{S}\frac{N_{S}}{t_{S}}\frac{dN_{A}}{dt_{A}} = V_{A}V_{S}\frac{N_{S}}{t_{S}}r_{N_{A}} + f_{in}N_{in} - f_{out}N_{A}N_{S}$$
(10)

where N is the concentration of substance our biomass inside the reactor, V is the Volume of liquid inside the reactor, t is the time, f_{in} is the inlet flow rate (dm³/h) in the reactor, N_{in} is the concentration of N in the inlet flow, f_{out} is the outlet flow rate (dm³/h) of the reactor, and N_{out} is the concentration of N in the outlet flow. The subscripts S and A are defined on section 4.1.

This model can be used for representing both batch, fed-batch and CSTR models. For the batch case, both f_{in} and f_{out} are zero. For the fed-batch case, only f_{out} is zero. For CSTR, f_{in} and f_{out} are greater than zero and f_{in} = f_{out} at steady state.

4.4 Simulation using Physics Informed Neural networks

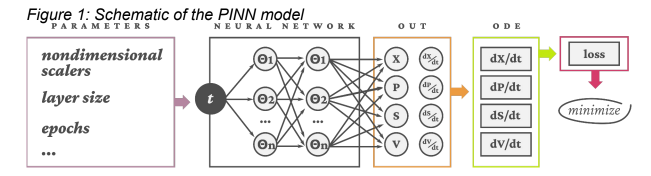
Physics-Informed Neural Network (PINN) was introduced to the scientific community in 2019 (RAISSI; PERDIKARIS; KARNIADAKIS, 2019). One of the most important points of the technique is to allow scientists to simulate physical, chemical or biological systems using differential equations as a font of data source for the optimization of Neural Networks (NN) created specifically to solve those problems. This is possible because NN are intrinsically universal approximators (SANTANA et al., 2022).

The overall loss function is defined as the weighted sum of the L^2 residuals (LU et al., 2021):

$$L = \sum w_N \left\| \left(\frac{dN}{dt} \Big|_{calculated} - \frac{dN}{dt} \Big|_{predicted} \right) \right\|^2$$
 (11)

where N represents X, S, P or V, and w_N is the weight of loss for N.

To test many different PINN configurations, a custom Grid Search and repetition loop was created. The PINN evaluation (without the repetition loop) scheme adopted in shown in 1. Both hyperparameters and equations parameters are fixed before each loop. Then, the Neural Network is fed with the data from the equation system. The outputs (labeled as OUT in the Figure) are the variables of interest (X, P, S and V) and their respective derivatives according to time. Then, the loss function is calculated comparing the given derivatives at step "n" compared with what their actual value should be. This can be achieved simply by making the difference (subtraction) of the derivative calculated (as show in equations 5-10) and the ones predicted by the NN.



In the original paper that introduced PINN (RAISSI; PERDIKARIS; KARNIADAKIS, 2019) questions suchs as how deep/wide the neural network must be to appropriately represent the mathematical models, the relevance of normalization and/or nondimensionalization and how the weights of loss functions can impact on the model loss and accuracy were raised. Because this subject still being debated and more studies need to be carried out (SANTANA et al., 2022), these points will be evaluated in the next topics. This work compares traditional (Euler), experimental (ALTIOK; TOKATLI; HARSA, 2006) and PINN solution for biological reactions, in specific the production of LA by *L. casei*, and strategies to improve performance and reduce loss in different reactors regiments (batch, fedbatch and CSTR). Therefore, is out of the scope of this work to discuss meticulously the implementation of PINNs.

4.5 Simulation configurations

The main questions to be answered are:

- Is it possible to approximate simple, ordinary differential equations (ODEs), with PINNs using lower numbers of neurons and layers? The simple Burgers equation required at least 20 neurons per layer and a few layers in another study (LU et al., 2021);
- Do the weights of the loss function of each variable impact significantly on the accuracy and number of steps required for acceptable results?
- What impacts does the independent variable (time) nondimensionalization cause to the loss?
- How does the dependent variables (X, P, S, V) nondimensionalization affect the system? Are some of them more relevant than the others?
- Since the same mathematical model of reactor is being used for the three cases (batch, fed-batch and CSTR), can the best configuration (group of parameters and hyperparameters, such as number of neurons per layer and

- nondimensional scalers) for the batch reactor also represent accurately the other two?
- Is it possible to find a pattern of relationship between hyperparameters, loss weights and nondimensionalization scalers in order to improve the traditional Random Grid Search, replacing it with a more rational and predictable approach?

The bacth reactor, by definiton, has closed inlets and outlets. The fed-batch reactor volume is allowed to increase indefinitelly and the CSTR volume is regulated by a function in order to open the outlet as the volume capacity (5 L) is approached. Batch starts with 5 L of solution and Fed-batch and CSTR starts with 1 L, all with initial concentrations of Biomass, LA and whey lactose as the same used in the experiment (case 2 of (ALTIOK; TOKATLI; HARSA, 2006)). CSTR is simulated for a greater time (96 h) while batch and fed-batch are both simulated for 10.6 h (slightly above the experimental time). The inlet flow of fed-batch is set to 2L/h, while the inlet flow of CSTR is set to 1 L/h. The outlet function for the CSTR is given:

$$f_{\rm in} * (V/V_{max})^7 \tag{12}$$

Were V_{max} is the maximum liquid volume in the reactor, set as the same of the maximum volume of the reactor used in the original experiment (5 L).

All simulations were run using all weights = 1, all nondimensionalization scalers (t_s , X_s , P_s , S_s , V_s) = 1, number of training points = 800, number of testing points = 1000, layer size = 22 neurons in 3 layers, epochs (iterations of Adam algorithm (KINGMA; BA, 2017)) = 30,000, learning rate = 1×10^{-3} , except when explicitly stated otherwise. The PINN model was solved using DeepXDE (LU et al., 2021), and the ner activated by a hyperbolic tangent function.

5 RESULTS

In order to try to find patterns between many of the hyperparameters and process parameters, tests were conducted in parallel instead of conducting a traditional Random Grid Search. Table 1 summaries the tests.

Table 1 - Conclusions for each test

| Test Name | Objectives | Conclusion |
|-----------------|---|---|
| t_S | Evaluate the effect of nondimensionalization scaler of time (t _s) | In general, the greater $t_{\mbox{\scriptsize S}}$ the greater the loss |
| nondim | Evaluate the effect of nondimensionalization scalers of X _S , P _S , S _S , and V _S | Batch, fed-batch and CSTR optimizations vary greatly, and there is no general rule |
| layer_size | 1) Evaluate the effect of layer size in performance 2) Check if it is possible to rationalize the ratio neuron/layers (N/L) in function of the loss | 1) 2) It was not possible to find any clear relationship between loss, N/L and total number of neurons |
| multiple_config | Evaluate if the same configuration can appropriately represent batch, fed-batch and CSTR | Almost all configurations tested can represent the batch reactor accordingly, and almost no one can represent fedbatch and CSTR |
| pinn_numeric_xp | Validate the proposed batch model, comparing with the solution using Euler Method and Experimental Data | The model was successfully validated |

Nondimensionalization effects can greatly improve CSTR and fed-batch performance, but easily made batch reactor simulation yield greater losses. The variation of $t_{\rm S}$ did not only not improve the simulation of batch reactor error, but also resulted in greater loss (2 - a). From all the values tested for the independent variable (t) nondimensionalization, the best $t_{\rm S}$ was found to be one. In other words, the best

model is found to be the one which does not nondimensionalize the time. The nondimensionalization of time, as a rule, increase the loss of the model and, in many cases, made the NN stagnate at points next to loss = 1. Thus, the nondimensionalization of time was ruled out as noneffective. It must be noted, however, that the 6 cases tested for time were greater or equal 1. Since the loss increased with the increase of $t_{\rm S}$, values of $t_{\rm S}$ <=1 may improve model performance in systems with multiple dependency (P and S depend directly on X, and X, P and S depend on V), such as the presented.

The nondimensionalization of X, P, S and V was conducted in sequence, with 6 different combinations. The layer size was tested as 5 layers with 32 neurons and 3 layers with 22 neurons. CSTR loss was reduced greatly using the smaller layer. The best cases for CSTR were using n1 and n6 (). The best cases for fed-batch was n4, with loss < 10⁻². Despite of that, none of these models were capable of representing adequately any o the reactors, even with more than 80,000 iterations.

Table 2 - Test Parameters

| Test Name | Parameters |
|-----------|--|
| t_S | t_1 : experiment time (9 h) t_2 : 1 / (μ * So / (K_s +So) t_3: |
| | |

Vale à pena fazer isso? Acho que só pra dissertação,

pro artigo não...

The layer size test was performed solely for the batch reactor case and is show in 2 - b. It was tested 12 different layer sizes. With a constant number of neurons of 22, experiments t lay7, t lay8 and t lay9 varied only the number of layers to 3, 4 and 5, respectively. While it is show a great improvement from 3 to 4 layers, the loss of 5 layers is about the same, for 22 more neurons to optimize. We can infer that, while increasing the number of layers may improve loss reduction, it can also need many more epochs to optimize, reducing the loss for a same given number of epochs. Experiments t lay4, t lay5 and t lay6 were done for 3, 4 and 8 layers respectively, and show that when using smaller number of neurons more deepness (number o layers) is necessary to reduce loss and get out of local minimums. From t lay 1 to 3, we can infer that ratios of neuron/layers (N/L) < 1 can yield better results than ratios > 1 for this range. From t lay 4 to 6, we can infer that an N/L > 4 can yield better results increasing the number of layers, with N/L varying from 5.33 to 2.67. From t lay 7 to 9, the loss reduces as N/L goes from 7.33 to 4.4. The loss is also smaller comparing t lay11 (N/L = 10.67) to t lay12 (N/L = 6.4). While there is a general trend in this data that N/L = 3 or less can generate better models, it was not possible to determine a general rule for the given system of equations. The best performance (smaller loss) was achieved in t lay12, the model with more neurons per layer (32 neurons in 5 layers, N/L = 6.4).

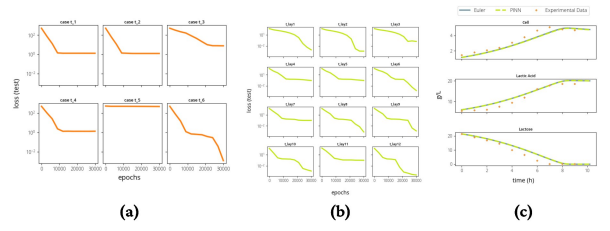


Figure 2: Nondimensional test, layer test and comparison between experimental data and PINN results

The plots shown are described on Table 1, and are: (a) $- t_S$ test; (b) $- layer_size$; (c) $- pinn_numeric_xp$.

Loss weights tests were performed in different layer sizes (8 neurons in 6 layers, 22 neurons in 3 layers, 32 neurons in 5 layers, and 90 neurons in 3 layers) (data not published). It is clear that the simulation for the batch reactor showed a good performance for any group of parameters, with the exception of W3, where $w_P =$ 3. Fed-batch is the model with the worst performance in almost every group of weights, but performs better at W2, with $w_X = 3$. Surprisingly, fed-batch loss was smaller in the deeper and narrower layer size (8 neurons in 6 layers), with an error of. 8.03×10⁻¹ after 120,000 epochs. Batch performed better in the most equilibrated layer size (32 neurons in 5 layers), with an error of 1.37×10⁻⁵ after 45,000 epochs. The only configuration were CSTR performed well (loss <10⁻²) was also the only one were it got smaller losses than the other two reactors, with layer size of 90 neurons in 3 layers and 120,000 epochs. This can't be determined as an optimal point, though, since both batch performed slightly worse than the other variations and fed-batch loss was greater than 10. This indicates that the CSTR model needs more effort to be solved, probably due to one more variant (fout) that the others don't have. The batch and fed-batch models, though, not only benefit from simpler models because of less computational cost, but also can achieve slower losses in fewer epochs in simpler Neural Networks. The best performance of batch reactor happened on test W3,

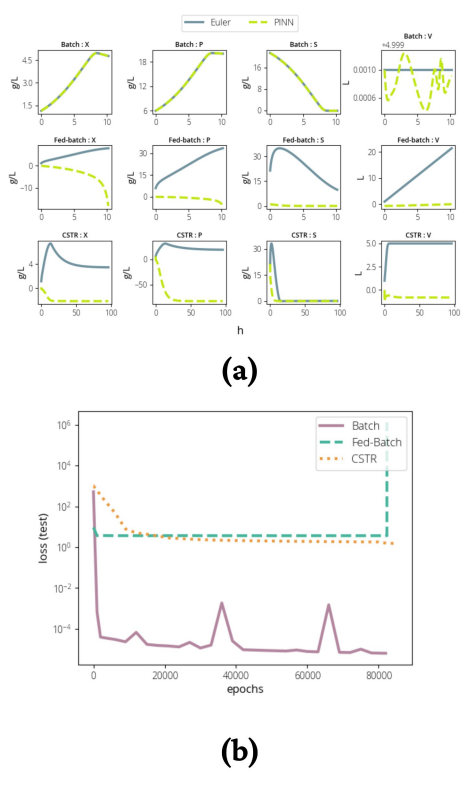
For visual validation of the model proposed, the system of equations was solved for the batch reactor case using the traditional Euler Method (with 240

discretization points in time) and the result is plotted, alongside PINN (60,000 epochs and layer size of 32 neurons and 5 layers) and experimental data (from (ALTIOK; TOKATLI; HARSA, 2006)) in 2 - c. We can conclude that the given PINN configuration can represent with great accuracy the biological reaction. The graphic doesn't answer, however, if the same model is capable of solve both the reaction and the volume variation, since in batch dV/dt = 0. The graphics and data (not published) show that the most probable culprit is the volume. Since the model wasn't being capable of evaluate the correct values of V and dV/dt, this impacted in X, P and S. Since P and S depend on both X and P, they were significantly affected. Thus, it was expected that a greater w_V would make the solver more sensitive to errors in the volume variable and, consequently, be able to produce more precise results. Since this could not be validated on the loss weight tests, it is possible that this specific problem requires a great amount of neurons (>90) and layers (>5) than the ones that were tested.

None of the combination of above configurations (layer size, number o neurons per layer, number of iterations/epochs, loss weights or nondimensional scalers) was capable of representing the fed-batch and CSTR reactors. The CSTR model employed in this work starts in transient state and is expected to converge to stationary state after some time. Then a final approach was tried, applying alongside the Adam algorithm, L-BFGS (BYRD et al., 1995) as a previous and post-processing step, combined with loss weights and nondimensionalization found to be the best fit on previous attempts. The result ins show in 3 - a. The name of each subplot is constructed as "Name of the reactor: Name of the variable". So "Batch: V" for instance, represents the volume in the batch reactor over time. While the batch reactor model presents a heavy fluctuation of volume, it is irrelevant and more than 3 orders of magnitude smaller than the total value (approximately 5 L). The CSTR model results are alike the ones for many other configurations and the same reactor: only one of the four variables values could be more or less acceptably predicted, while the others are clearly deviated. Fed-batch has the greatest loss of the three models in almost any configuration. Also, both fed-batch and CSTR errors are clearly stagnated in values of order of magnitude of about 10°, as show in 3 - b. Another very relevant topic is that, as shown in equation 5, there are terms raised to exponents. While ideally X and P would never be higher than X_m and P_m, respectively, this can't be assumed from a Neural Network that is learning and updating itself based on

errors. Because of this, fluctuation of errors on volume or biomass (which were determined to be the most sensible features, probably because they are directly involved in the equations of P and S) can rapidly destabilize the system, predict values of X and P that are greater than their respective maximum values and generate negative numbers raised to 0.5 for example. This was the cause of numerous errors, so many tests were not finished: they were prematurely interrupted, since the error was not a number (NaN) or infinity, and the system was not capable of recover from these fluctuations. While this may induce us to think that the losses values being too high were entirely related to the biomass and product inhibitory, the batch model shows clearly that they are not the main cause - at least not alone. The volume shows some fluctuations even though X, P and S are appropriately in accordance with the solution obtained using Euler Method. Therefore, we conclude that the main challenges faced by the system while trying to minimize the loss function were the volume balance, the cells balance, or both. The mere use of loss weights and nondimensionalization didn't prove effective for reducing these problems and enabling the same configuration to represent both batch, fed-batch and CSTR reactor models.

Figure 3: PINN comparison for Bacth, Fed-batch and CSTR reactors



(a) – The profile of biomass (X), product (P), substrate (S) and volume (V) for each reactor simulated. (b) – The total loss for each reactor over the number of epochs/iterations.

While PINNs are very interesting and open the doors of Machine Learning to many applications in fields of science that don't have a lot of data available, we believe a better approach would be to use PINNs to model only the reaction given a starting point, and to solve the physics-involved variables, which are much more straightforward, by a traditional numeric method. The system would be, then, a hybrid of PINN and traditional numeric integrator or solver system. We also could not find a clear relation between hyperparameters, nondimensionalization and loss weights that could serve as a guide for future works and substitute or rationalize the Grid Search.

6 CONCLUSIONS

Simpler Neural Networks, with less neurons and layers, were capable of producing acceptable losses for the reaction only, but performed poorly on balancing the reactions and the volume variation simultaneously. Almost all configurations produced high losses when dV/dt was not expected to be zero. Since the PINN performance for the batch reactor was exceptionally good and both CSTR and fedbatch models required a large amount of epochs to yield a still high loss, in cases where both the volume of the reactor is varying considerably and there are multiple reactions related to each other happening, a better approach may be combine the PINN for solving the biological or chemical set of equations (group of reactions), which is intrisically more complex, and use a regular numeric method, like Euler or Runge-Kutta, to solve the physics part of the system (volume and mass balance) which is usually more well-known and established.

Physics-Informed Neural Networks were shown to represent with adequate accuracy biological reactions with complex therms and multiple derivative dependency. However, all combinations tested failed to represent adequately both the concentrations and volume variation of the reactor. We also recommend future works to try employing nondimensionalization scalers for time $(t_{\rm S}) < 1$, even if the selected $t_{\rm s}$ is arbitrary and not based on actual constants from the equations, to explore new strategies to deal with these kind of physical and biochemical complex systems and to propose another ways to rationalize the process of creating a PINN model.

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