Modeling and Simulation of Cell Signaling Networks for Subsequent Analytics Processes Using Big Data and Machine Learning

Máximo Eduardo Sánchez-Gutiérrez1 and Pedro Pablo González-Pérez2*

1Colegio de Ciencia y Tecnología, Universidad Autónoma de la Ciudad de México, Ciudad de México, México. 2Departamento de Matemáticas Aplicadas y Sistemas, Universidad Autónoma Metropolitana, Unidad Cuajimalpa, Ciudad de México, México.

ABSTRACT: This work explores how much the traditional approach to modeling and simulation of biological systems, specifically cell signaling networks, can be increased and improved by integrating big data, data mining, and machine learning techniques. Specifically, we first model, simulate, validate, and calibrate the behavior of the RIK/AKT/mTOR cancer-related signaling pathway. Subsequently, once the behavior of the simulated signaling network matches the expected behavior, the capacity of the computational simulation is increased to grow data (data farming). First, we use big data techniques to extract, collect, filter, and store large volumes of data describing all the interactions among the simulated cell signaling system components over time. Afterward, we apply data mining and machine learning techniques—specifically, exploratory data analysis, feature selection techniques, and supervised neural network models—to the resulting biological dataset to obtain new inferences and knowledge about this biological system. The results showed how the traditional approach to the simulation of biological systems could be enhanced and improved by incorporating big data, data mining, and machine learning techniques, which significantly contributed to increasing the predictive power of the simulation.

KEYWORDS: Big data techniques, modeling and simulation approach, cell signaling networks, machine learning techniques, predictive analytics

Introduction

Big data techniques are commonly applied whenever raw data is too large to be processed by a computer system. Big data also refers to when the management systems or database servers cannot provide the required data in a reasonable time due to problems with loading, searching, selecting, and saving.\(^1\)\(^3\)

It is difficult to find a widely accepted definition of the term big data because it comes from very different origins or domains. In real-time computer systems, response time becomes an essential variable.

- **Velocity.** It means the speed with which data is generated, collected, and processed. Let us think about the remarkable speed with which data is generated in applications such as search engines, the stock market, e-commerce platforms, and social networks, to name a few examples. In real-time computer systems, response time becomes an essential variable.

- **Variety.** Indicates the non-homogeneity or diversity of the data because it comes from very different origins or sources, implying that the data are of very different types, such as numeric, Boolean, categorical, nominal, ordinal, structured text, unstructured text, images, and videos among others.

However, in recent years, other “Vs” have been proposed to contribute to the definition of big data, such as Veracity and Value. The term Veracity refers to the low quality that sometimes characterizes large-scale data. In other words, the need to contend with the uncertainty in the data, mainly due to the

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wide variety that characterizes them derived from the different sources that generate them. The veracity problem in large-scale data commonly occurs in unstructured text data, commonly generated in sources such as social networks, emails, and chats, due to the freedom that characterizes its creation. On the contrary, the term Value refers to the importance and significance that big data can provide in making decisions that lead to companies, businesses, and institutions being much more profitable and successful.

It is undeniable that the computational simulation approach could be enhanced and improved by integrating big data techniques, which would be a valuable support for the acquisition, processing, and analytics of the large volume of data that computer simulations produce continuously. Specifically, big data techniques provide a means to obtain and evaluate large-scale data produced for computational simulations, as well as to extract causal and temporal relationships between input and output patterns, which will allow us to carry out further predictions and inferences about the behavior of the simulated system.

Biology has been one of the disciplines strongly favored with the support of computational simulation. In the last 2 decades, a wide range of computational simulations of biological systems and processes, such as protein folding, artificial faldamer design, molecular docking, and cell signaling networks, among others, have been developed based on a wide range of mathematical and computational models (see a survey of these models). In particular, the modeling and simulation of cell signaling networks have ranged from continuous and discrete mathematical models such as systems of differential equations or numerical methods, respectively; to computational models, such as cellular automata, Petri nets, Boolean models, rule-based systems, multiagent systems, and artificial neural networks (ANNs).

During the execution of the simulation of the biological system, large volumes of data are generated, eg, proteomics, genomics, interactomics, and metabolomics, among others, which, once acquired, structured, and stored, will undoubtedly constitute a valuable input for the predictive analytics application, commonly based on data mining techniques.

Therefore, in this piece of work, we are exploring how much the traditional approach to modeling and simulation of biological systems, specifically cell signaling networks, can be increased and improved by integrating big data and machine learning techniques.

Essentially, in this work we (1) model the PI3K/AKT/mTOR signaling network, identifying cellular compartments, signaling elements, the types of interactions between them, and the kinetic parameters and initial parameters concentrations that characterize the interactions, and the signaling elements, respectively; (2) simulate, verify, and calibrate the expected behavior of the PI3K/AKT/mTOR signaling network on the Big-Data Cellulat bioinformatics platform; (3) generate large volumes of data describing the behavior of the simulated biological system over time; and (4) apply exploratory analysis, feature selection, and analytics processes to the resulting biological dataset, to obtain new inferences and knowledge about this biological system.

We state that when the computational simulation of a biological system has been finely tuned and verified then, beyond the observed simulated behavior and the subsequent in silico experiments carried out, one of its strengths lies in the large volume of reliable biological data that it can produce. As a result, these data, through exploratory analysis and analytics process, can produce new inferences and knowledge about the simulated biological system.

**Material and Methods**

**The PI3K/AKT/mTOR signaling pathway**

Intracellular signaling is one of the essential molecular mechanisms for controlling cell activity, and it is involved in almost all cell functions, including cell division, growth, differentiation, and death. Cancer progression, malignancy, and treatment resistance are all influenced by signaling pathways. The intracellular signaling pathways linked with NF-kB (nuclear factor kappa B), TGF-β (transforming growth factor beta), Notch, and PI3K/AKT/mTOR are the most commonly altered in cancer.

The PI3K/AKT/mTOR signaling pathway is engaged in many tasks relevant to cancer biology, including cellular proliferation, survival, migration, angiogenesis, and apoptosis, making it one of the most critical processes in cancer growth. The research of anticancer targets working through the PI3K/AKT/mTOR signaling pathway requires a comprehensive understanding of this signaling pathway: the characteristics of its signaling elements, the complex interactions that occur between them—signal amplification, activation, deactivation, phosphorylation, dephosphorylation, complex formation, and several others—and the global behaviors that ensue; which requires the use of approaches such as systems biology, big data, and data mining. It is critical for cancer treatment to understand the activity of PI3K/AKT/mTOR and how it interacts with other pathways that are regulated by the presence of specific molecules.

In several types of cancers, including brain, breast, ovarian, and renal carcinomas, unregulated activation of the PI3K/AKT/mTOR signaling pathway contributes to cellular change and tumor growth. The PI3K/AKT/mTOR pathway is important for a variety of cellular processes because it contains a complicated signaling mechanism including 3 key modulators proteins: PI3K (Phosphoinositide-3-kinase), AKT (Serine/threonine kinase, also known as protein kinase B), and mTOR (Serine/threonine kinase, mammalian Target of Rapamycin).

**Big-Data Cellulat: the cell signaling network simulator**

The Big-Data Cellulat platform was conceived as a computer simulator for cellular signal transduction systems.
The simulator is based on a model that integrates (1) the concept of tuple space\textsuperscript{12,32,33} for the representation and interaction of chemical reactions and reactants and (2) an action selection mechanism based on the Gillespie algorithm\textsuperscript{34,35} for the selection and execution of chemical reactions. The joint use of these 2 approaches allows Big-Data Cellulat to exhibit a series of key characteristics required to simulate cell signaling systems and the consequent in silico experimentation. On one hand, the representation based on tuple spaces provides the simulation with characteristics such as multi-compartmentalization, localization, and topology; on the other hand, the selection and execution of chemical reactions based on the Gillespie algorithm provide the simulation with synchronization, timing, and a selection based both on the rate/affinity of the chemical reaction and on the availability of the reactants.

**Representation of the chemical reactions and reactants.** A tuple is an ordered collection of information or knowledge, and as a knowledge representation, tuples aid in representing the chemical reactions and reactants. In a tuple space, the interaction and synchronization between functions, procedures, objects, programs, and even intelligent agents, occur through reading, modifying, writing, and destroying tuples in the shared tuple space.\textsuperscript{32,33} Based on these considerations, the translation of the structures and elements involved in cell signaling to abstractions of the tuple spaces is shown in Figure 1 and described in Table 1.

**Figure 1.** Use of tuple spaces to represent cell compartments, reactants, and chemical reactions involved in cell signaling. Note that the selection and execution of chemical reactions are coordinated by an action selection mechanism based on the Gillespie algorithm.
where $Moli$, $1 \leq i \leq k$, is the number of molecules required of reactant $i$, $n_i$ is the number of available molecules of reactant $i$, and $reqMol_i$ is the number of molecules required of reactant $i$, $1 \leq i \leq k$.

2. Selection of the next chemical reaction to run.

$$\psi \leq \sum_{i=1}^{n} \frac{Rate_i}{RTot}$$

where $\psi$ is a random number, $0 \leq \psi \leq 1$ and $RTot$ is the summation of the rates ($Rate_i$) of all reactions.

3. Determination of the delay (suspenion) between the last reaction executed and the next reaction to be executed.

$$Stop_{time} = \frac{-\ln(\tau)}{RTot}$$

where $\tau$ is a random number, $0 \leq \tau \leq 1$.

The main characteristics and functionality of Big-Data Cellulat simulator. As a computational simulator of cell signaling systems, Big-Data Cellulat exhibits characteristics that are crucial when trying to emulate the structure and behavior of this type of complex biological systems such as compartmentalization, localization, topology, interaction, coordination, synchronization, timing, and selection and execution of chemical reactions considering their rate/affinity. As previously mentioned, these characteristics emerge from the joint use of (1) a tuple space model for the representation and interaction of the structures, elements, and components involved in cell signaling and (2) coordination and action selection mechanism based on the Gillespie algorithm, for the selection and execution of chemical reactions. At this point, it is necessary to note that the functionality exhibited by the Big-Data Cellulat simulator can be described in terms of the characteristics mentioned above. That is, each of these characteristics constitutes in itself a feature that the simulation tool provides to the user during the phases: (1) creation of the simulation; (2) execution, calibration, and validation/verification of the simulation; (3) execution of in silico experiments; and (4) production and recording of massive data for intelligent data analysis tasks.

Farming big cell signaling data

As pointed out by Tolk, among the big-data methods closely related to the modeling and simulation approach are data farming and crowdsourcing. Both methods are beneficial when applied to steps of traditional modeling and simulation studies. In particular, data farming uses computational simulations (in silico experiments) to grow data. Once the data are produced and stored—in this case by the Big-Data Cellulat simulator—it can be analyzed using various techniques and models, such as data mining and machine learning, to discover causal relationships between them. When the Big-Data Cellulat simulator is used, data farming takes place once the simulation is launched, encompassing the following actions:

1. Selection/filtering of the features (signaling elements) required for the dataset integration.
2. Selection of the sampling factor ($K$ milliseconds, seconds or minutes, with $K$ integer, $K > 0$) for data recording.
3. Identification of the path, file name, and extension where the data will be stored.

Exploratory data analysis and feature selection techniques

The purpose of the exploratory data analysis is to generate as many insights and information about the data as possible and find any problems in the dataset. One of the most common issues found in datasets is missing values. Two frequently used techniques to handle missing values in a dataset are dropping rows or columns and replacing missing values with central tendency values such as mean, median, and mode. Deleting rows or columns with missing values may produce a model that works poorly if the percentage of missing values is excessive compared to the complete dataset. On the contrary, inputting missing values prevent data loss but do not factor the covariance between features.

Another common issue concerning datasets is the unequal distribution of classes within a dataset, known as data imbalance. Some techniques can be used to solve the class imbalance...
problem; resampling by oversampling or undersampling\textsuperscript{36} and ensemble methods.\textsuperscript{37} Oversampling can be carried out by generating as many synthetic samples as needed, selecting the most common value in the class and repeating it, or repeating a randomly selected value from the smallest class. In contrast, undersampling is a technique that decreases the number of samples of the most significant class down to the smallest class size. These 2 techniques can be combined to oversample the minority class and undersample the majority class. On the contrary, ensemble methods typically use boosting or bagging to build several estimators on a different randomly selected subset of data.

One advantage of the Big-Data Cellulat simulator is that it can farm data by computational simulations (in silico experimentation). In the context of data imbalance, artificial generation of data points is unnecessary because the simulation can be run multiple times and, consequently, join the resulting datasets. In this case, the class imbalance can be dealt with undersample techniques to reduce the majority classes. In general, raw datasets contain various data types, including numerical and categorical information. Feature engineering deals with these heterogeneous datatypes using various techniques that convert different data types to numerical vectors.\textsuperscript{38} For example, to encode categorical or numerical features, one can use Dummy Encoding, Count Encoder, One-Hot Encoder, and idmax, among others. Similarly, feature binning converts continuous to categorical variables.

**Predictive process**

Once the raw dataset is preprocessed, and the class information is encoded, the data are ready to be fed to a predictive model. In this work, we choose a multilayer perceptron (MLP)\textsuperscript{39} to predict the cellular state or states that should characterize the cell, given a particular activation/deactivation configuration of the signaling elements that make up the network. An MLP can be implemented as a classifier because it finds the most appropriate boundary between 2 or more classes. Hence, it may discern the structural differences between 2 or more given classes, identify the space that separates each one, and determine the likelihood of a given data point belonging to a particular class. An MLP is a neural network connecting multiple neurons or perceptrons, partitioned into the input layer, the hidden layer, and the output layer. The neurons compose a directed acyclic graph, meaning that the paths connect nodes in layers from one layer to the next, as shown in Figure 2. Each neuron, excluding the input ones, has a nonlinear activation function, a bias, and connecting weights which the MLP train by backpropagation in a supervised learning fashion\textsuperscript{40} so that the error value can be updated in a much more successful way.

When developing a neural network model, 3 stages are needed before its deployment: (1) dataset preprocessing, (2) performing feature engineering, and (3) dividing the dataset into training and testing sets using a cross-validation strategy. The input dataset to the machine learning model usually requires partitioning the data into training and test sets. Data belonging to the training set contains a known output or label, from which the model learns to generalize to other data. On the contrary, the test set is used to test our model’s prediction capabilities. In this work, we perform a cross-validation schema to split the dataset by partitioning the available data into 3 sets (see Figure 3).

**Methodological approach**

The methodological approach followed in this work integrates the key aspects of traditional modeling and simulation with current big data, data mining, and machine learning techniques, involving the following activities:
1. **Modeling.** Modeling the PI3K/AKT/mTOR signaling network, identifying cellular compartments, signaling elements, the types of interactions between them, and the kinetic parameters and initial concentrations that characterize the interactions and the signaling elements, respectively.

2. **Creation of the computational model (simulation).** Assembly in Big-Data Cellulat of cell structures (cell compartments, cells, tissues), chemical reactions with their kinetic parameters, and reactants with their initial molar concentration value.

3. **Simulation, validation, and calibration.** Simulation, verification, and validation of the expected behavior of the PI3K/AKT/mTOR signaling network in the Big-Data Cellulat bioinformatics platform.

4. **Data farming.** Generation of big data describing the behavior of the simulated biological system over time.

5. **Intelligent data analysis.** Application of data mining and machine learning techniques to the resulting biological dataset to obtain new inferences and knowledge about this biological system.

**Results and Discussion**

*The modeling of the PI3K/AKT/mTOR signaling network*

The main results obtained in the modeling phase are illustrated in Figure 4 and described in Table 2. Figure 4 shows the resulting model of the PI3K/AKT/mTOR signaling network-integrated from segments, cascades, particular types of interactions, as well as other theoretical-experimental aspects reported in the
specialized literature on this antiapoptotic signaling pathway and its role in cancer. In the representation of the signaling network illustrated in Figure 4, the nodes represent signaling elements such as membrane receptors, proteins, and transcription factors; the arcs establish the different types of interaction that occur between the signaling elements such as activation, inhibition, compounding, among others, while the rounded edge rectangles correspond to the final cell states achievable from specific activation/inhibition combinations of signaling elements. Note that the primary cellular compartments involved in intracellular signaling have also been identified. On the contrary, Table 2 provides examples of the reactions that formalize the interactions between the signaling elements illustrated in Figure 4. As shown in Table 2, each reaction is characterized by its kinetic parameters and the concentration micromolar initial of the reactants involved; all these parameters are required in the subsequent simulation creation phase. It should be noted that the global signaling network illustrated in Figure 4 involves more than 60 reactions.

The simulation of the PI3K/AKT/mTOR signaling network

The behavior of the simulation of the PI3K/AKT/mTOR signaling pathway over time (on a millisecond scale) is captured in the snapshots illustrated in Figure 5. All in silico experiments carried out during this phase aimed to validate, verify, and calibrate the simulated biological system, which was facilitated by running the simulation in phases of incremental complexity. That is, the signaling pathway was split into the following signaling segments identified from a biological approach: (1) from the activation of transmembrane receptors to the activation of proteins and the formation of compounds in the juxtamembrane region, (2) from the activation of transmembrane receptors to the activation of key proteins and the formation of compounds in the cytoplasm, (3) from the activation of transmembrane receptors to the activation of transcription factors in the nucleus, and (4) from the activation of transmembrane receptors to the triggering of final cellular processes.

The simulation verification, validation, and calibration processes aim to reduce the error between the simulated micromolar concentration values of the target signaling elements and their concentration values of these signaling elements observed in the real biological system. The simulation validation was based on the analysis of differences between simulated concentration values and measured concentration values, using statistical indices such as the mean bias error (MBE), the mean absolute error (MAE), the mean square error (MSE), and the root mean square error (RMSE).

### Table 2. Examples of chemical reactions defined as part of the modeling of the PI3K/AKT/mTOR signaling pathway.

| REACTION | REACTANTS | INITIAL CONC. (µMOL) | K_M (µMOL) | V_MAX (µMOL/µL/SEG) | V_S |
|----------|-----------|----------------------|------------|---------------------|-----|
| Cyt + RK → CytRK | Cyt RK | 0.1 0.25 | 34.2 | 7.6 | 2.22 × 10⁻⁵ |
| CytRK + JAK → CytRKJAK⁺ | JAK Cyt RK | 0.012 0.0001 0.25 | 34.2 | 7.6 | 2.22 × 10⁻⁵ |
| CytRKJAK⁺ + STAT → STAT⁺ | STAT Cyt RK | 0.4 0.0001 0.25 | 74.1 | 49 | 6.61 × 10⁻⁵ |
| RAS⁺ + PI3K → PI3K⁺ | PI3K RAS | 0.9 0.8 | 53.4 | 49 | 0.0915 |
| PIP3⁺ + Akt → Akt⁺ | PIP3 Akt | 0.27 0.2 | 1.1 | 22.1 | 4.3554 |
| PDK1⁺ + Akt → Akt⁺ | PDK1 Akt | 1.0 0.2 | 36 | 22.3 | 0.6027 |
| Akt⁺ + p27⁺ → p27 | Akt p27 | 0.2 0.27 | 7.8 | 8.4 | 0.2810 |
| Akt⁺ + FKHR⁺ + FOXO⁺ → FKHR/FOXO | Akt FKHR FOXO | 0.2 0.4 0.4 | 74.1 | 49 | 0.2630 |
| FKHR/FOXO → Apoptosis inhibition | FKHR/ FOXO | 0.4 | 74.1 | 49 | 0.2630 |
| STAT⁺ → Proliferation/Angiogenesis | STAT⁺ | 0.4 | 74.1 | 49 | 0.2630 |
| p27 → Cell cycle activation | p27 | 0.27 | 7.8 | 8.4 | 0.2810 |
The big cell signaling dataset produced by the computer simulator

As previously described, the main product resulting from the data farming stage is a large volume of input-output patterns produced by in silico experiments carried out by Cellulat bioinformatics framework. In this case, the product is the simulation of the behavior of the PI3K/AKT/mTOR signaling network from different concentration settings of its signaling elements.

Table 3 shows the characteristics of the resulting dataset in terms of the number of features, number of instances, and number of classes. The dataset comprises 35,574 instances characterized by 77 attributes and 6 classes in which instances are classified as Autophagy, Proliferation, Inhibition Apoptosis, Cell Growth, Proliferation Angiogenesis, and Cell Cycle Activation.

Table 3. Characteristics of the PI3K/AKT/mTOR signaling dataset resulting from the data farming process.

| BIOLOGICAL DATASET            | NUMBER OF FEATURES (SIGNALING ELEMENTS) | NUMBER OF INSTANCES | NUMBER OF CLASSES (CELLULAR PROCESSES) |
|-------------------------------|------------------------------------------|---------------------|----------------------------------------|
| PI3K/AKT signaling network    | 77                                       | 35,574              | 6                                      |

In more than one class. In other words, the classes are not mutually exclusive. Table 4 shows the number of instances in each class for each sampling period.

As mentioned earlier, machine learning requires a dataset with which the learning process can be carried out. First, this dataset needs to go through a preprocessing stage, including data cleansing, to transform it into a format that a machine learning algorithm can understand. In our dataset, as shown in Table 4, the signaling pathway identifies 6 classes in which the instances are grouped as Autophagy, Proliferation, Inhibition Apoptosis, Cell Growth, Proliferation Angiogenesis, and Cell Cycle Activation. The discrepancy between instances reported in Tables 3 and 4 is due to the classes not being mutually exclusive. To prevent that one instance can be assigned to more than one class, the initial representation of the signaling database was transformed to a suitable scheme that would allow its processing by machine learning algorithms, as shown in Table 5. With this new assignment of patterns to classes, the number of instances grouped in each of them underwent the update shown in Table 6 and Figure 6.

Observe in Figure 6 that the number of instances is imbalanced; the majority class is about 20.7 times the minority class. This class imbalance can lead to models biased toward the majority class, causing the wrong classification of the minority class.
To alleviate this problem, we are presented with 2 options: oversample the minority classes or undersample the majority classes. As the dataset comes from a data farming stage of the patterns produced by in silico experiments carried out by the Cellulat bioinformatics framework, the number of samples that we can obtain is only restricted by the framework processing time; this presents us with the opportunity to use the undersampling technique. As stated in Table 6, the dataset is composed of 77 descriptors, and one attribute representing the class, the whole relation of features (signaling elements) and interactions are shown in Figure 4.

Results of the exploratory data analysis and feature selection

As previously stated, because the number of instances is imbalanced, in this work, we handle the class imbalance by undersampling the majority classes avoiding the need to generate samples from the same dataset artificially. This technique produces a random subsample of a dataset by removing random observations of the majority classes, and the redistribution of samples is shown in Figure 7.

Once the dataset was balanced, we explored a set of feature selection techniques to rank the features according to their saliency to get an idea of the importance of the features. These techniques consider data variance, chi-square stats, feature ranking with recursive cross-validated feature elimination, a linear model with iterative fitting, a meta estimator that fits randomized decision trees, and Pearson correlation. An excerpt of the 77 ranked features is available in Table 7. As each technique ranks the features differently, Table 7 shows the relative importance of the features for each technique. It is important to note that if we group the features in the top 10, 20, . . ., some features are found to belong in the same tier across selection techniques, eg, PI3K*, AKT, BAX*, AMPK*, among others.
After ranking the features, we can select the ones that meet specific criteria, eg, the ones that explain 80% of the total variance, and in general, the features that are the most likely to be class-dependent and therefore more relevant for classification. Below we present the prediction results produced by the MLP model after applying several selection techniques using (1) the entire dataset (77 features/descriptors), (2) the domain expertsuggested feature selection (12 features/descriptors), (3) the complete set of primary descriptors identified by each technique, and finally, (4) the complete set of primary descriptors identified by each technique enriched with a subset of features according to domain experts.

Results of the predictive process on the big cell signaling dataset

The results of the 4 experiments that explored the effects of the different selection techniques are shown in Table 8. To have a baseline to compare against, we use the entire dataset without any feature selection, ie, the 77 features. This first experiment achieved an accuracy of 96.15%. In contrast, the second experiment used the 12 features suggested by the domain expert, obtaining only 93.64% accuracy. The 12 suggested features (signaling elements) in this case were CycD, BCL2, E1F4E, P27, BCLXL1, STAT*, XIAP, PI3K, AKT, P21, RAS, and FKHR-FOXO (see the whole PI3K/AKT/mTOR signaling pathway in Figure 4). Regarding the number of features, only 15.6% of the features were used in the second experiment; this means that there is room for improvement in the number of features selected and accuracy. This also means that the features suggested by the domain expert may not be the best ones to describe the data.

We ran a third group of experiments to investigate the effect of selecting different subsets of features by different techniques. The results of such a group of experiments are listed in the Selected features column of Table 8; in this column, we can see that the Recursive feature elimination technique yields a better accuracy rate using only 23 of the 77 original features.

Finally, the results of the Selected and domain knowledge features experiment in Table 8 show that accuracy wise, not only it is not beneficial to add the suggested domain features, but if we compare the cardinality (number of features) of both experiments, we can also see an increment. In this regard, note that the Pearson correlation technique worsens the accuracy by adding only one more expert-suggested feature set. Contrarily, the accuracy of the low variance technique significantly improves when adding 10 features proposed by the expert knowledge; this unusual behavior may be explained by considering that it is the smallest selected feature set, and it may not be enough discriminative information. This behavior is schematized in Figure 8.

Concerning the selected subsets, we can note that from the 12 domain expert-suggested features, BCL2 appears in the top 10 of 3 feature selection techniques, BCLXL1 also appears in the top 10 of 3 columns in Table 7, the same occurs for XIAP and AKT. In addition, inside Table 7’s top 10, BAX* and PI3K* appear in 4 columns. On the contrary, the domain expertsuggested features do not appear to impact the second half of the top 20; nevertheless, mTOR-RICTOR* and SHP2 appear in all the selected subsets, while 14-3-3*, GRB2, SOS, CytRKJAK*, and SHC are selected by 4 techniques.

Strengths and weaknesses

The data generated by the Cellulat bioinformatics framework includes inputs and their discrete labels; this presents the opportunity to tackle the supervised learning task as a classification problem. Random forest (RF), support vector machines (SVM), and ANNs are some of the most prevalent classification algorithms. A common idea in big data and machine learning (ML) is that the more data you have, the more accurate your results will be; nevertheless, massive datasets come with their own set of problems. Unstructured data formats,
Table 7. First 20 features were obtained after applying several feature selection techniques.

| PREDICTOR'S RELATIVE IMPORTANCE | LOW VARIANCE | CHI² | LASSO | EXTRA TREES | PEARSON CORRELATION |
|---------------------------------|--------------|------|-------|-------------|---------------------|
| 1                               | BAX*         | BCLXL1 | XIAP  | E1F4E       | TSC2*               |
| 2                               | AMPK*        | PI3K* | BCLXL1 | BCLXL1      | XIAP               |
| 3                               | RAS*         | 4EBP1 | BAX*  | 14-3-3*     | FOXO*              |
| 4                               | PI3K*        | BCL2  | mTOR-RICTOR* | AKT         | FKHR*              |
| 5                               | BCL2         | PDK1* | mTOR1* | STAT*       | CRAF               |
| 6                               | ULK1*        | ULK1* | PI3K*  | FOXO*       | 14-3-3*            |
| 7                               | 14-3-3-BAD   | SHC   | RHEB  | BAX*        | mTOR-RICTOR*       |
| 8                               | PDK1*        | GRB2  | BCL2  | XIAP        | AKT                |
| 9                               | 4EBP1        | SOS   | CytrKJAK* | mTOR-RAPTOR | mTOR-RAPTOR       |
| 10                              | AKT          | CytrKJAK* | BCLXL1 | RHEB*       | BAX*               |
| 11                              | mTOR-RICTOR* | SHP2  | AMPK*  | mTOR-RICTOR* | AMPK*             |
| 12                              | 14-3-3*      | AMPK* | GRB2  | FKHR*       | RAS*               |
| 13                              | GRB2         | RAS*  | LKB1* | P21         | SHC                |
| 14                              | SHP2         | AKT*  | 14-3-3-BAD | CRAF      | CytrKJAK*         |
| 15                              | SOS          | BAX*  | SHC   | SHP2        | SOS                |
| 16                              | CytrKJAK*    | 14-3-3 | ULK1* | CASP2*      | GRB2               |
| 17                              | SHC          | mTOR-RAPTOR | SHP2  | LKB1*       | SHP2               |
| 18                              | mTOR1*       | 14-3-3 | SOS   | AKT*        | RHEB*              |
| 19                              | 4EBP1*       | AKT   | TSC2* | PIP3        |                   |
| 20                              | RHEB*        | mTOR-RICTOR* | PIP3  | STAT*       |                   |

These features were ranked by their discriminative power and, to ease its understanding, partitioned in sets of 10. This partitioning lets us analyze the shared features among techniques.

Table 8. Accuracy results of MLP machine learning algorithm.

| SELECTION TECHNIQUE | CARDINALITY | ACCURACY OF SELECTED FEATURES | ACCURACY OF SELECTED AND DOMAIN KNOWLEDGE FEATURES | CARDINALITY |
|---------------------|-------------|------------------------------|-----------------------------------------------|-------------|
| Feature selection techniques | Low variance | 12 | 82.71 | 92.45 | 22 |
|                       | Lasso       | 18 | 94.85 | 94.76 | 27 |
|                       | Recursive feature elimination | 23 | 96.62 | 96.41 | 31 |
|                       | Pearson correlation | 48 | 96.54 | 95.65 | 49 |
|                       | CHI²        | 54 | 95.84 | 95.84 | 54 |
|                       | Extra trees | 54 | 95.84 | 95.84 | 54 |
| Base line (all features) | 77 | 96.15 | | |
| Domain knowledge features | 12 | 93.64 | | |

Abbreviation: MLP, multilayer perceptron.

The bold-evidenced accuracies compare the best results of the algorithmically selected features with the algorithmically selected features plus the domain-knowledge suggested features. The italicized rows show the baseline accuracy results after using the complete set of features and the domain-knowledge suggested features, i.e., the results without feature selection and without automatic feature selection, respectively.
noisy and poor-quality data, unbalanced input data distribution, unlabeled data, and high dimensionality are common problems. Another important consideration is that some machine learning algorithms were created assuming that the complete dataset could fit in memory. Big data ignores these assumptions, rendering standard algorithms useless or severely slowing them down.

Support vector machines, which try to find the optimal hyperplane with the highest margin between classes, a random forest, which can be described as an ensemble of classification trees, where each tree votes on the class assigned to a given sample, and ANNs, which can be described as parallel computing units that can separate nonlinear data, are some of the most common algorithms for supervised classification. As the number of samples and classes grows, so does the complexity of these algorithms. Building a random forest, e.g., becomes more expensive as the number of trees increases. In SVMs, the worst-case scenario is if the training set contains as many support vectors as samples. Although multiclass SVMs exist, their canonical implementation requires the training of a separate SVM for each class. Selecting the appropriate architecture for a specific problem in ANNs, such as the MLP used in this work, is still an open research issue.

The data mining approach to big data, empowered by machine learning techniques presented in this work, ameliorates the concerns mentioned early by acquiring, processing, and analyzing large data volumes to reduce its complexity.

Conclusions

With this study we try to improve the traditional approach to modeling and simulation of biological systems—specifically, cell signaling networks—by integrating big data, data mining, and machine learning techniques. As a result, new inferences and knowledge were obtained from the dataset generated from the simulated system, which allowed increasing the predictive capacity of the latter.

First, the behavior of the PI3K/AKT/mTOR signaling network was modeled, simulated, verified, and calibrated; subsequently, large volumes of data describing the behavior of the simulated biological system over time were produced by running the simulation (data farming); and finally, exploratory data analysis, feature selection techniques, and analytics processes were applied to the resulting biological dataset, obtaining new inferences and knowledge about this biological system.

The resulting dataset was obtained by farming a large volume of input-output patterns produced by in silico experiments carried out by the Cellulat bioinformatics framework. These input-output patterns represent the activation/deactivation state of the 77 elements that make up the PI3K/AKT/mTOR signaling network (input pattern) and the cellular processes associated with the configuration (output pattern). The cell signaling dataset was used as input to the machine learning process using an MLP algorithm. However, for it to be helpful, we went through a cleaning and a preprocessing stage; this process involved the statistical feature selection techniques to evaluate the saliency of the features (signaling elements in the PI3K/AKT/mTOR signaling network). The predictive model resulting from applying the MLP automated learning algorithm yielded new knowledge about the simulated system by allowing the prediction of the cellular state or states associated with a specific input pattern made up of a reduced number of signaling elements.

Finally, the results of the evaluation of the machine learning model for the different selected subsets show that the use of
feature selection techniques not only improves the accuracy rate of the MLP but also improves its performance, because only 30% of the original 77 characteristics are necessary to improve the baseline.

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Author Contributions
Both authors contributed equally to conceptualization, methodology, validation, formal analysis, investigation, data curation, original draft preparation, review and editing, visualization, supervision, and project administration. Both authors have read and agreed to the published version of the manuscript.

ORCID iDs
Maximiliano Eduardo Sánchez-Gutiérrez https://orcid.org/0000-0003-1101-5956
Pedro Pablo González-Pérez https://orcid.org/0000-0001-7223-9035

Data Availability
The Cell signaling dataset that support the findings of this study are available at https://raw.githubusercontent.com/elMaxPain/files/master/CRISP/PI3KAKT_50_123_500ms.arff.

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