Prediction of Drug Loading in the Gelatin Matrix Using Computational Methods

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ABSTRACT: The delivery of drugs is a topic of intense research activity in both academia and industry with potential for positive economic, health, and societal impacts. The selection of the appropriate formulation (carrier and drug) with optimal delivery is a challenge investigated by researchers in academia and industry, in which millions of dollars are invested annually. Experiments involving different carriers and determination of their capacity for drug loading are very time-consuming and therefore expensive; consequently, approaches that employ computational/theoretical chemistry to speed have the potential to make hugely beneficial economic, environmental, and health impacts through savings in costs associated with chemicals (and their safe disposal) and time. Here, we report the use of computational tools (data mining of the available literature, principal component analysis, hierarchical clustering analysis, partial least squares regression, autocovariance calculations, molecular dynamics simulations, and molecular docking) to successfully predict drug loading into model drug delivery systems (gelatin nanospheres). We believe that this methodology has the potential to lead to significant change in drug formulation studies across the world.

1. INTRODUCTION

The global market for drug delivery systems is a multibillion-dollar industry, demand for which is growing in both developed and emerging economies (in part, driven by aging societies and rapid urbanization).1−9 Drug delivery systems deliver drugs at rates controlled by specific features of the systems, particularly their chemical composition (e.g., inorganic/organic components, molecular weights of their constituents, cross-linking density of polymers, etc.).10−12

The selection of the appropriate system (carrier and drug) to obtain optimal delivery is a challenge investigated by researchers in academia and industry, in which millions of dollars are invested annually.13 Experiments involving different carriers and determination of their capacity for drug loading are very time-consuming and therefore expensive. Consequently, approaches that exploit multivariate statistical methods, molecular simulations, docking methods, and mining the data in the literature14−19 have the potential to make hugely beneficial economic, environmental, and health impacts through savings in costs associated with chemicals (and their safe disposal) and time.

Computational/theoretical chemists/biochemists, biomedical/chemical engineers, and pharmacists have developed a variety of methodologies that can be applied to understand drug formulations. Principal component analysis (PCA) and hierarchical clustering analysis (HCA) are considered exploratory data analysis and unsupervised machine learning methods, where these techniques extract patterns from the independent factors (x-variables) only and irrelevant to the y-outcomes. Partial least squares (PLS) is a supervised pattern recognition method correlating the inputs with outputs and subsequently leads to the generation of a model.20 This data mining approach (through a retrospective analysis) combined with computer-aided analysis and simulation extracts knowledge from complex variables and responses obtained from historical records. The significant advantage of this approach is the possibility of uncovering interactions and linear relationships that might not be easily detectable with conventional experimental designs.21 Although not yet fully explored in drug formulation/delivery, multivariate statistical methods such as PCA and agglomerative HCA were previously used to develop drug delivery formulations. For example, PCA was utilized to generate a quantitative composition–permeability relationship for microemulsion formulations used to deliver testosterone transdermally, with a linear relationship between the lower-dimensionality data generated from the main principal components and the permeability coefficients of the different formulations.22 PCA and HCA were used to extract stable

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SMEDDS (self-microemulsifying drug delivery systems) and SNEDDS (self-nanoemulsifying drug delivery systems) formulations of lovastatin and glibenclamide, respectively,23,24 and PCA and PLS analysis were used to assess the qualitative and quantitative effects of different variables such as lipid/surfactant type and their concentrations on parameters related to storage stability.25 Furthermore, PLS was successfully employed to predict the sizes and polydispersity index (PDI) for lipid nanocapsules based on the quantitative mixture composition.26

Here, we extend these exciting studies by combining PCA, HCA, and PLS with molecular dynamics and docking analysis27 to give valuable insight into drug loading in a polymer matrix. As a model polymer matrix, we use protein-based nanoparticulate drug delivery systems (i.e., nanospheres composed of collagen-derived gelatin). Gelatin is an abundant and inexpensive protein,28 which is amphiphilic in nature due to its amino acid contents (ca. 12% anionic glutamic and aspartic acid, ca. 13% cationic lysine and arginine amino acids, and ca. 11% hydrophobic leucine, isoleucine, methionine, and valine),29 and gelatin-based matrices can in principle be used to deliver both small molecules and macromolecules.30−36 In this study, we focus on a selection of low-molecular-weight drugs used in the clinic, as depicted in Figure 1.

2. MATERIALS AND METHODS

2.1. Data Set. The data set contained four input variables (descriptors) and one output response (mass of drug loaded per 100 mg gelatin nanospheres determined experimentally) for different drugs. Data mining was performed through different databases such as PubMed and Web of Science to obtain the output response for 10 drugs: acyclovir,37 amphotericin B,38 cryptolepine,39 doxorubicin,40 5-fluorouracil (5FU),41 isoniazid,42 resveratrol,43 paclitaxel,44 and indomethacin.45

2.2. Calculation of Molecular Descriptors. The drugs were analyzed using Bioclipse version 2.6 (Bioclipse project, Uppsala University, Sweden).39 The four descriptors chosen were constitutional (molecular weight), electronic (number of hydrogen bond donors and number of hydrogen bond acceptors), and physicochemical (xLogP).

2.3. Hierarchical Clustering Analysis (HCA). The molecular descriptors generated using Bioclipse version 2.6 were subjected to hierarchical clustering analysis using JMP 7.0 (SAS, Cary, NC, USA). Ward’s minimum variance method was adopted to join the clusters and generate a dendrogram. Ward’s method is considered an agglomerative hierarchical technique where the merging in the dendrogram starts at the final clusters (leaves) and merging occurs stepwise until it reaches the trunk. Ward’s minimum variance criterion minimizes the total within-cluster variance. At each step, the pair of clusters possessing the minimum between-cluster distance is merged (i.e., the pair of clusters that leads to the minimum increase in the total within-cluster variance after merging is selected).45

2.4. Principal Component Analysis (PCA). PCA was used to extract patterns using an exploratory data analysis method that deals with the variances in sample observations. PCA was performed using JMP 7.0. Four principal components were calculated by taking a linear combination of an eigenvector of the correlation matrix built up from standardized original variables. The dimensionality of the data was reduced by extracting two main principal components possessing the two highest eigenvalues and plotting the data with respect to these two new orthogonal axes.

2.5. Partial Least Squares Analysis (PLS) for Model Generation and Validation of the Model. PLS was used to study correlations between the molecular descriptors and the output response. PLS was performed using JMP 7.0 using four latent vectors. The PLS generated model was validated by

Figure 1. Chemical structures of the substances studied herein: (A) acyclovir, (B) cryptolepine, (C) amphotericin B, (D) doxorubicin, (E) 5-fluorouracil (5FU), (F) isoniazid, (G) resveratrol, (H) paclitaxel, (I) indomethacin, and (J) curcumin.

2.5. Partial Least Squares Analysis (PLS) for Model Generation and Validation of the Model. PLS was used to study correlations between the molecular descriptors and the output response. PLS was performed using JMP 7.0 using four latent vectors. The PLS generated model was validated by
A Berendsen barostat, respectively.49 Out for 3 ns at 373 K and 1 bar using a v-rescale thermostat and constrained by the LINCS algorithm. The MDS were carried handle long-range electrostatic interactions. All bonds were force-

-probe molecules (with a calculated molecular weight of AGPRGQ(Hyp)GPAGPDGQ(Hyp)GP. Six hypothetical molecule. The primary sequence of the peptides was www.gromacs.org/). To prepare the gelatin system, 48 peptide

cgen means 15 × 15 Å as these dimensions were suitable to the size of the docked molecules and ensured a central position for them inside the gelatin matrix. Additionally, the genetic algorithm was used as the docking engine with 150 maximum poses. The type of calculation and ligand (as chosen using the software options) were Dock and Flexible, respectively, and the binding energies (ΔG, kcal/mol) reflecting the docking efficiencies were calculated.

### 3. RESULTS

Table 1 reports the molecular descriptors (number of hydrogen bond donors, number of hydrogen bond acceptors, xLogP, and molecular weight) for the investigated drugs. The dendrogram classifying these drugs according to HCA using Ward’s minimum variance method (an agglomerative type of analysis) is displayed in Figure 2. Isoniazid and 5FU were clustered together according to their four descriptors, Resveratrol and cryptolepine clustered together, whereas doxorubicin, acyclovir, and amphotericin B constituted separate clusters. Importantly, the loading pattern followed this classification (see Table 1) where SFU and isoniazid scored the highest loading masses followed by acyclovir, which is closest to the aforementioned drugs in the dendrogram. Cryptolepine and resveratrol were very close, with doxorubicin near to them. Amphotericin B had the lowest mass loaded into the nanospheres, which was clear from its separate branch (furthest distance) in the dendrogram.

Table 1. Descriptors of the Drugs, Amounts of Loaded Drug, and the Obtained Binding Energies from Docking of the Drugs on a Simulated Gelatin Matrix

| drug          | xLogP | no. H-bond donors | no. H-bond acceptors | molecular weight (g/mol) | actual amount of drug loaded (mg/100 mg gelatin) | Lamarckian genetic algorithm ΔG (kcal/mol) |
|---------------|-------|------------------|----------------------|--------------------------|-----------------------------------------------|---------------------------------------------|
| acyclovir     | −1.650| 3                | 8                    | 225.21                   | 8.74                                          | −3.94                                        |
| amphotericin B| 2.068 | 12               | 18                   | 923.49                   | 1.16                                          | 144.4                                       |
| cryptolepine  | 2.180 | 0                | 2                    | 233.30                   | 2.00                                          | −3.81                                        |
| doxorubicin   | −1.900| 6                | 9                    | 543.52                   | 2.10                                          | 58.29                                        |
| 5-fluorouracil| −0.760| 2                | 4                    | 130.00                   | 25.07                                         | −4.19                                        |
| isoniazid     | −0.683| 3                | 4                    | 137.14                   | 22.00                                         | −4.16                                        |
| resveratrol   | 2.050 | 3                | 3                    | 228.24                   | 1.96                                          | −3.74                                        |
| curcumin      | 1.95  | 2                | 6                    | 368.13                   | 3.50                                          | −2.59                                        |
| pacitaxel     | 6.15  | 4                | 14                   | 853.33                   | 0.52                                          | 173.5                                        |
| indomethacin  | 3.78  | 1                | 4                    | 338.14                   | 1.91                                          | −1.99                                        |

\[
Q^2 = \frac{\text{PRESS}}{\text{ISS}}
\]

where PRESS represents the predicted residual error sum of squares, while ISS stands for the total initial sum of squares. Moreover, a predicted versus actual correlation was obtained.

### 2.6. Molecular Dynamics Simulations (MDS) of the Gelatin Matrix.

Molecular dynamics simulations (MDS) were carried out using the GROMACS\textsuperscript{46} v. 4.6.5 freeware (http://www.gromacs.org/). To prepare the gelatin system, 48 peptide molecules were constructed, with 18 amino acids in each molecule. The primary sequence of the peptides was AGPRGQ(Hyp)GPAGPDGQ(Hyp)GP. Six hypothetical probe molecules (with a calculated molecular weight of 767.13) were added at random positions to the system. The force-field parameters were obtained from CgenFF\textsuperscript{47} (https://cgenff.paramchem.org/). The system was energy minimized by the steepest descent method. Molecular dynamics was subsequently carried out, with a time step of 2 fs, full periodic boundary conditions, and a cutoff distance of 1.2 nm for van der Waals and electrostatic interactions.\textsuperscript{48} PME was chosen to handle long-range electrostatic interactions. All bonds were constrained by the LINCS algorithm. The MDS were carried out for 3 ns at 373 K and 1 bar using a v-rescale thermostat and a Berendsen barostat, respectively.\textsuperscript{49}

### 2.7. Drug Docking in Simulated Gelatin Nanospheres.

The chemical structures of the studied drugs were drawn using ChemDraw Ultra version 10 (Cambridgesoft, Waltham, MA, USA). The corresponding “.mol2” files needed for docking experiments were obtained using Chem3D Ultra version 10 (Cambridgesoft, Waltham, MA, USA) after energy minimization using the MM2 force field of the same program. Docking analysis was generated by Argus Lab version 4.0.1 (Mark Thompson and Planaria Software LLC, Seattle, WA, USA). The hypothetical probe molecules were utilized to construct corresponding binding sites on the carrier (gelatin-probe), and the AScore was utilized for calculating the scoring function. The size of the display box in the x, y, and z dimensions were 15 × 15 × 15 Å as these dimensions were
A score plot of the drugs with respect to their descriptors after projecting the data into two main principal components is displayed in Figure 3, where principal component 1 and principal component 2 reflect 69.72 and 26.95% of the data variation, respectively (corresponding to 96.68% of total variance; Figure 3, top right panel), and 5FU and isoniazid are clustered together with acyclovir having the nearest score, and amphotericin B the furthest score. Figure 4 depicts the loading plots of the two main principal components. It is obvious that principal component 1 is mainly composed of the descriptors: the molecular weight, the number of the H-bond donors, and the number of the number of H-bond acceptors, while principal component 2 mainly depends on the remaining descriptor, xLogP. These results confirm the presentation of the four investigated variables in the two generated principal components.

The relationship between the obtained combined x-scores (combining the contribution from the four x-variables viz. descriptors) and y-scores is displayed in Figure 5, and the screen plot (Figure 5, bottom right) depicts the contribution of each individual latent factor to the combined x-scores with the first two factors accounting for 96.64% of the obtained scores. It is noteworthy that the generated x- and y-scores represent the distances of the points in space of all the dimensions to the main vector summarizing the final dimension (in the current case, there is a principal component or vector for the x-dimension comprising all the descriptors and another for the y-dimension representing the loaded mass). Therefore, the aforementioned scores can be negative numbers. Consequently, a generated model was developed, where

\[
Y = 13.175 + 0.115 \times \text{xLogP} + 0.001 \times \text{number of hydrogen bond donors} \\
+ 2.346 \times \text{number of hydrogen acceptors} \\
- 0.059 \times \text{molecular weight}
\] (1)

The values and the signs of the coefficients of the x-factors in the equation were indicative of the importance of increasing the number H-bond acceptors in the drug chemical structure in the presence of a balanced xLogP and low molecular weight to increase the loading of the drug. The model was validated by performing t-test statistical analysis between the actual experimental results for drug loading and the predicted drug loading using the model where no significant difference was obtained between the means at \( P < 0.05 \). The calculated \( Q^2 \) or the predicted R-squared after 5-fold cross-validation scored a value of 0.721 (a highly acceptable value).
drug molecule in a gelatin matrix after projecting its structure.

The obtained results can be explained by the fact that gelatin is a protein carrier with a relatively balanced hydrophilic/hydrophobic character displaying several hydrogen bond donor and acceptor groups with a repetitive sequence of amino acids -Ala-Gly-Pro-Arg-Gly-Glu-4Hyp-Gly-Pro- along its backbone. This structure can be transformed to some numerical values that are generated of each amino acid. Among which are the highly condensed variables “z-scale descriptors” that are derived from PCA analysis of several experimental and physicochemical properties of the 20 natural amino acids: z1, z2, and z3, which represent the amino acids hydrophobicity, steric properties, and polarity, respectively. Additionally, they are useful in QSAR analysis of peptides where they have proven effective in predicting different physiological activities.53–55 Herein, we used an extended scale (including 67 more artificial and derivatized amino acids)56 due to the presence of 4-hydroxyproline in the gelatin structure.

In this study, we expand the use of the first descriptor (z1) to predict the drug loading properties of nanoparticles. The first scale (z1) was chosen as it represents a lipophilicity scale that encompasses several variables (amino acid descriptors) such as the thin layer chromatography (TLC) variables, log P, nonpolar surface area (Snp), and polar surface area (Spol) in combination with the number of proton-accepting electrons in the side chain (HACCR).57 In this scale, a large negative value of z1 corresponds to a lipophilic amino acid, while a large positive z1 value corresponds to a polar, hydrophilic amino acid. Therefore, the gelatin typical structure amino acids (-Ala-Gly-Pro-Arg-Gly-Glu-4Hyp-Gly-Pro-) can be represented by their z1 values as follows: (0.24), (2.05), (−1.66), (3.52), (2.05), (3.11), (−0.24), (2.05), and (−1.66). Furthermore, an overall topological description of the repetitive sequence was accounted for by encoding the z1 descriptors of each amino acid into one covariance variable47 that was first introduced by Wold et al.58 The autocovariance value (AC) was calculated as follows

$$AC_{z, lag} = \frac{\sum_{i=1}^{N-lag} V_{z,i} V_{z,i+lag}}{N - lag}$$

where AC represents autocovariances of the same property (z-scale), i = 1, 2, 3, ..., N is the number of amino acids, lag = 1, 2, 3, ..., L (where L is the maximum lag, which is the longest sequence used) and V is the scale value.

Therefore, the AC value for the gelatin typical structure sequence was calculated with lag 1 scoring a value approaching zero (0.028), indicating a balanced hydrophobicity/hydrophilicity structure. In light of the above, the high loading of SFU and isoniazid can be ascribed to their amphiphilic nature with logP values approaching 0 and to the presence of several hydrogen bond donors and acceptors groups relative to their low molecular weight that is favorable in both diffusion through and entrapment in a protein matrix like that of gelatin nanospheres. Since there was a recorded deviation between the actual and the predicted values regarding isoniazid and SFU (may be attributed to their small molecular weight that helps their nonstoichiometric physical entrapment in the gelatin matrix), therefore, the results were further confirmed by molecular dynamics and docking experiments, where the drugs were docked on the gelatin matrix simulated structure. Figure 7 shows the molecular simulation of the gelatin nanosphere matrix. Interestingly, the best binding energy values \(\Delta G\) (−4.19 and −4.16 kcal/mol) corresponded to the highest loaded drugs SFU and isoniazid, respectively, followed by acyclovir (see Figure 8). In the same context, amphotericin B scored a highly positive \(\Delta G\) value, which explains its low loading values. The confirmation of the docking results with their experimental counterparts can be attributed to the
The binding energy due to hydrophobic forces, the binding energy due to van der Waals forces, and similar studies. 10 studies, which we recommend to increase in further number of the experimental studies that are involved in it simulated gelatin matrix. The only limitation of the model was loaded drugs through docking the investigated molecule on the relationship can highly estimate the molar masses of physically highly experimental values and the docking results. explaining the high correlation obtained between the real drug and its carrier that may lead to drug entrapment, which nearly all the possible interactions that can occur between the energy. As can be inferred, the equation terms encompass and involved in torsions (rotors) that were frozen due to binding, deformation is the energy due to rotational bonds and atoms (chg) is the binding energy due to H-bonding, $G_{\text{H-bond}}$ is the binding energy due to H-bonding to charged molecules, $G_{\text{deformation}}$ is the energy due to rotational bonds and atoms in Fig 8. Drug loading versus the obtained binding energy plot of the investigated drugs after docking on a simulated gelatin matrix built up using molecular dynamics simulation displaying an exponential relationship.

inclusive scoring function of the Arguslab software. This scoring function is based on the XScore calculated according to the following equation

$$\Delta G_{\text{bind}} = \Delta G_{\text{vdw}} + \Delta G_{\text{hydrophobic}} + \Delta G_{\text{H-bond}}$$

$+ \Delta G_{\text{H-bond (chg)}} + \Delta G_{\text{deformation}} + \Delta G_0$ (3)

where $\Delta G_{\text{bind}}$ is the total calculated binding energy, $\Delta G_{\text{vdw}}$ is the binding energy due to van der Waals forces, $\Delta G_{\text{hydrophobic}}$ is the binding energy due to hydrophobic forces, $\Delta G_{\text{H-bond}}$ is the binding energy due to H-bonding, $\Delta G_{\text{H-bond (chg)}}$ is the binding energy due to H-bonding to charged molecules, $\Delta G_{\text{deformation}}$ is the energy due to rotational bonds and atoms involved in torsions (rotors) that were frozen due to binding, and finally, $\Delta G_0$ represents the regression-obtained binding energy. As can be inferred, the equation terms encompass nearly all the possible interactions that can occur between the drug and its carrier that may lead to drug entrapment, which explains the high correlation obtained between the real experimental values and the docking results. An exponential model was generated correlating the actual experimental molar masses of the loaded drugs and their corresponding docking binding energies. This model was highly fitting with an obtained R-squared value of 0.95. This relationship can highly estimate the molar masses of physically loaded drugs through docking the investigated molecule on the simulated gelatin matrix. The only limitation of the model was the number of the experimental studies that are involved in it (10 studies), which we recommend to increase in further similar studies.

5. CONCLUSIONS

The current study introduces new approaches of interpreting and predicting drugs loading on protein carriers, such as gelatin nanospheres. These approaches comprise multivariate statistical methods such as hierarchical clustering analysis, principal component analysis, partial least squares regression, molecular dynamics, and docking. Moreover, the utilization of the amino acids z-scales descriptors represents a new and important asset in interpreting drug loading in protein-based carriers. We believe that this methodology has the potential to lead to significant change in drug formulation studies across the world.

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Notes
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