

EVALUATION OF WAVELENGTH SELECTION METHODS FOR 2D FLUORESCENCE SPECTRA APPLIED TO BIOPROCESSES CHARACTERIZATION

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Abstract - In biotechnological processes, the productivity and costs depend strongly on the control of the operating conditions. For this reason, sensors that allow the monitoring of variables of interest become quite important. 2D fluorescence spectroscopy is one promising option among those that are being applied for this purpose. In the present work, three methods were evaluated to select the best excitation/emission wavelength pairs of 2D fluorescence spectra to infer product, substrate and cellular concentrations throughout a fermentation using a multiple linear chemometric model: Exhaustive Search (ES), *Stepwise Regression* and Genetic Algorithm (GA). The *Stepwise Regression* presented unsatisfying results, while GA always led to good R^2 values in short computational times. However, for the proposed problem, the ES showed the best performance, finding the global optimum in a few minutes.

Keywords: Fermentation; 2D fluorescence spectroscopy; Wavelength; Variable selection.

INTRODUCTION

Due to the high dependence of biological systems (microorganisms and produced metabolites) on the physical and chemical characteristics of the cultivation medium, bioprocesses can become quite complex (Scheper *et al.*, 1996, Scheper *et al.*, 1999). Therefore, the maintenance of product quality and reduction in production costs of these processes are highly associated with monitoring and controlling the operating conditions. For this purpose, there is still a need for sensor development in order to achieve reliable online monitoring and, consequently, to decrease the time delay between data analysis and

appropriate control actions (Hulhoven, 2006, Surribas *et al.*, 2006).

For classical properties such as temperature, pressure, pH and dissolved oxygen, measurement methods are already well established, enabling their online monitoring. On the other hand, the measurement of biological variables such as metabolites, substrate and cellular concentrations demands further improvement (Haack *et al.*, 2004). Several authors have studied different techniques that enable the inference of the variable of interest using chemometric models (Schügerl, 2001; Ferreira *et al.*, 2003; Benoudjita *et al.*, 2004; Becker *et al.*, 2007). Among these, the advantages of 2D fluorescence

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spectroscopy have been highlighted in many recent studies (Boehl *et al.*, 2003; Hantelmann *et al.*, 2006; Ganzlin *et al.*, 2007; Rhee and Kang, 2007; Ödman *et al.*, 2009; Jain *et al.*, 2011; Bogomolov *et al.*, 2011; Rossi *et al.*, 2012). In this technique, the fluorescence emission of metabolites present in the cultivation medium is monitored when they are excited with UV-Visible radiation in a specific wavelength range. The molecules that emit fluorescence are called fluorophores. This non-invasive technique can be considered an evolution of the fluorescence sensor that could monitor only the fluorophore NAD(P)H and then would obtain information on cellular metabolism. Once the 2D fluorescence spectroscopy is capable of monitoring all fluorophores, it can yield information not only about cellular metabolism, but also about the cultivation medium (Scheper *et al.* 1999, Ganzlin *et al.* 2007).

Since biomass contains a number of natural fluorophores like NAD(P)H, flavins and aromatic amino acids, its concentration can be directly correlated with the trajectories of the fluorescence of these analytes during the cultivation. On the other hand, substrates and products like glucose, ethanol and glycerol do not exhibit any fluorescence. In this case, their concentrations can still be indirectly predicted using the signals of available fluorophores (Rossi *et al.*, 2012). In the literature, there are several examples of successful indirect prediction models based on 2D fluorescence spectroscopy (Skibsted *et al.*, 2001; Solle *et al.*, 2003; Lee *et al.*, 2005; Surribas *et al.*, 2006; Rossi *et al.*, 2012). However, the metabolism of a microorganism can shift during the cultivation as a result of environmental changes, such as sequential uptake of different substrates, availability of oxygen or accumulation of inhibitors. As a consequence, the relation between the predicted variables and the respective analytes can also change. For processes in which these shifts are relevant, a segmented modeling approach applying multiple calibration has been studied (Ödman *et al.*, 2009).

Therefore, the 2D fluorescence spectra, which contain a great amount of information, need to be correlated with the variables of interest through a chemometric model. This task is not trivial because the matrix data has a high degree of collinearity. This linear dependency is a consequence of the fact that the same fluorophore can absorb radiation in a range of excitation wavelengths as well as emit fluorescence in a range of reemission wavelengths with different intensities. Usually, PCA and PLS techniques

are used in order to convert the spectrum data into prediction models for the desired variables (Solle *et al.* 2003). A drawback of these techniques is that the whole spectrum is needed to obtain the principal components or latent variables, since each predictor is a linear combination of the spectrum variables. This inconvenient can be overcome when methods for direct selection of variables are employed. Thus, a limited number of excitation/ emission wavelength pairs that can satisfactorily represent the system can be truly selected, enabling the construction of simpler sensors to monitor only those selected wavelength pairs. It makes the large-scale application of 2D fluorescence more feasible.

Regarding the discussion above, the aim of this work is to evaluate three different methods (i.e., Exhaustive Search, *Stepwise Regression*, and Genetic Algorithm) to select directly those wavelength pairs in order to infer product (ethanol), substrate (glucose) and cellular (biomass) concentrations throughout a fermentation using a multiple linear chemometric model. These results are further compared to those obtained using full-data PLS, which is the conventional technique employed to solve this kind of problem.

MATERIALS AND METHODS

Experimental Data

In this work, batch fermentation data, provided by Professor B. Hitzmann of the *Institut für Technische Chemie, University of Hannover*, and first presented in the work of Solle *et al.* (2003), were used for evaluating the selection methods. In that experiment, a fermentation with *S. cerevisiae* H620 using supplemental glucose Schatzmann medium was carried out in a 1.5 L bioreactor at a constant temperature of 30 °C and controlled pH of 5.5. Throughout the batch, 2D fluorescence spectra were recorded every 6 minutes with a BioView®-spectrometer (Delta Light & Optics, Denmark) as described by Stärk *et al.* (2002). Each spectrum contains 150 excitation/emission wavelength pairs that are the parameters to be selected by the studied methods.

In addition, 10 samples were withdrawn along the fermentation for off-line analysis of the variable of interest (i.e., ethanol, glucose and biomass concentrations) by high-performance liquid chromatography (HPLC). The measured values of these variables are given in Table 1.

Table 1: Experimental data of the single batch fermentation presented in the work of Solle *et al.* (2003).

Time [h]	Biomass [g/L]	Ethanol [g/L]	Glucose [g/L]
0	0.28	0.41	30
2.5	0.47	0.65	28.80
5	1.34	2.14	25.20
6.5	2.31	4.32	17.90
8	4.30	9.38	3.54
9.5	5.14	9.42	0.01
14.1	6.25	6.22	0
16	7.28	4.05	0
18.1	8.92	1.48	0
19.4	9.81	0.01	0

Reference Model

When *S. cerevisiae* is grown in glucose, even under aerobiosis, biomass and ethanol are produced and a diauxic pattern can be observed (Zhang *et al.*, 1997). The system can be modeled by the following three differential equations:

$$\frac{dC_g}{dt} = \frac{C_b \cdot \mu_G}{Y_{GB}} \quad (1)$$

$$\frac{dC_{et}}{dt} = \frac{C_b \cdot \mu_G}{Y_{GE}} - \frac{C_b \cdot \mu_E}{Y_{EB}} \quad (2)$$

$$\frac{dC_b}{dt} = C_b \cdot \mu_E + C_b \cdot \mu_G \quad (3)$$

C_g , C_{et} and C_b are the glucose, ethanol and biomass concentrations. The parameters μ_G and μ_E are, respectively, the specific growth rates in glucose and ethanol, while Y_{GB} , Y_{GE} and Y_{EB} are, respectively, the yield coefficients with respect to the conversion from glucose to biomass, glucose to ethanol and ethanol to biomass. The diauxic growth can be considered in the model as described by Solle *et al.* (2003): μ_G is only greater than zero if glucose is present and μ_E is only greater than zero if glucose is absent, and then there is growth on ethanol.

Due to the small amount of off-line data, concentration values were calculated for each point for which the 2D-fluorescence spectrum was available by numerical integration applying the Euler method. For this purpose, the dynamical model discussed above was fitted to the experimental data by minimizing the sum of squared errors using as initial values for the parameters (specific growth rates and yield coefficients) the ones reported by Solle *et al.* (2003). Thus, the adjusted values for the

model parameters used in this work were: $\mu_G = 0.469 \text{ h}^{-1}$, $\mu_E = 0.058 \text{ h}^{-1}$, $Y_{GB} = 0.162$, $Y_{GE} = 0.460$ and $Y_{EB} = 0.421$. Alternatively, the offline data could have been interpolated using other techniques, such as splines, as evaluated in another work of our group (Escobar *et al.* 2011). The generated concentration data were pre-treated by discounting the mean value and dividing by the standard deviation and used as reference in the evaluation of the selection methods.

Problem Definition

In our previous work (Ranzan *et al.*, 2011), it was shown by PCA analyses that 3 excitation/emission wavelength pairs could satisfactorily describe the variables of interest. A similar result was found by Haack *et al.* (2004) applying 2D fluorescence spectroscopy in an analogous system. Therefore, the selection methods were applied to find the best 3 excitation/emission wavelength pairs to predict the variable of interest (ethanol, glucose and biomass concentrations) at the same time using a multiple linear chemometric model.

For Exhaustive Search (ES) and the Genetic Algorithm (GA), the problem was structured as an optimization problem, which should choose 3 excitation/emission wavelength pairs while minimize a given objective function (FO). This function was the unexplained variance (S), defined as the quotient of the residual squares (SS_{err}) sum and the total squares sum (SS_t) as presented in Equations (4) to (6).

$$SS_{err} = \sum (y_i - f_i)^2 \quad (4)$$

$$SS_t = \sum (y_i - y_m)^2 \quad (5)$$

$$S = \frac{SS_{err}}{SS_t} \quad (6)$$

where y_i is the concentration according to the experimental data interpolated using the reference model (i.e., Equations (1) to (3)), f_i is the concentration predicted by the adjusted multiple linear chemometric model and y_m is the average concentration for the experimental data interpolated by the reference model. The minimization of S is equivalent to maximizing the coefficient of determination (R^2), given by

$$R^2 = 1 - \frac{SS_{err}}{SS_t} \quad (7)$$

For the *Stepwise* method, according to the interactive tool for *Stepwise Regression* of the Statistical toolbox of Matlab used in this work, the criterion was the root-mean-square error (RMSE), which is presented in Equation (8).

$$\text{RMSE} = \sqrt{\frac{\sum (y_i - f_i)^2}{n}} \quad (8)$$

n is the number of data points used of the reference model, which coincides with the number of 2D fluorescence spectra recorded.

However, in order to better compare the results, all results are presented in the form of the determination coefficient (R^2).

Selection Methods

In this work, three selection methods were evaluated: (i) Exhaustive Search (ES), (ii) *Stepwise Regression* and (iii) Genetic Algorithm (GA). All methods were run in Matlab® for Windows with a Core2Quad computer.

(i) Exhaustive Search (ES)

The ES was evaluated by a simple algorithm implemented in Matlab®, where all possible combinations of three excitation/emission wavelength pairs were tested. Thus, the achievement of the best three pair combination, meaning the lowest S , is ensured. Because the three models, for ethanol, glucose and biomass concentrations should be evaluated at the same time, the individual S 's were summed in order to have only one function to be minimized.

(ii) Stepwise Regression

Stepwise Regression is considered to be an efficient method for variable selection (Jouan-

Rimbaud *et al.*, 1995b). It proceeds as an automatic sequence of linear regressions, where in each step the best variable according to a specified criterion can be included in the model or the worst can be eliminated (Benoudjita *et al.*, 2004). In the function *Stepwise* from the Statistical toolbox of Matlab, which was used in this study, the criterion that decides which variable should be included or eliminated from model is the F-Test. Unlike with ES, the three models could not be evaluated at the same time and the number of excitation/emission wavelengths in the model could not be fixed due to the way the method is implemented. Thus, for each variable of interest, the final number of pairs and which pairs were chosen by the method could differ from the others.

(iii) Genetic Algorithm (GA)

The GA is a stochastic optimization method in which a new set of possible solutions is evaluated at each step, called a generation. Its randomness remains in the fact that the first set of solutions, called the initial population, is produced randomly and the further generations are produced from the combinations of that initial population (Neumann and Witt 2010).

In this work, GA was implemented in Matlab®. Each possible solution, called an individual of the population, was a three element vector containing integers from 1 to 150 that coincide with the 150 pairs available to be selected. The operators used to create the new generations were crossover (combination of two solutions of the last generation), mutation and elitism; while the stop criterion was the number of generations. As can be seen in Table 2 four sets of operator values were evaluated, and the generation number was fixed at 50. The convergence and stop criterion will be further discussed. In order to consider the randomness of the achieved solution, each set was executed ten times and the results presented are always the mean value of them.

Table 2: The four sets of GA parameters tested.

	Generations	Individuals	Reproduction rate (%)	Mutation rate (%)	Elitism rate (%)
Rep80Ind10	50	10	80	10	10
Rep80Ind100	50	100	80	10	10
Rep60Ind10	50	10	60	30	10
Rep60Ind100	50	100	60	30	10

RESULTS AND DISCUSSION

Table 3 presents the summary of the results for the evaluated methods of variable selection. The same table also includes the results of the PLS technique that will be further discussed.

Due to the great number of objective function (FO) evaluations, the algorithm ES needed the longest CPU time to achieve the solution. For this problem it expended ca. 9 minutes, ca. 650 times more than GA, which consumed ca. 0.7 seconds to converge to a set of three excitation/emission wavelength pairs. On the other hand, the optimality of the solution was guaranteed: pairs 11 (Emission 510 nm/ Excitation 270 nm), 46 (Emission 370 nm/ Excitation 330 nm) and 104 (Emission 530 nm/ Excitation 430 nm). The predictions obtained by the adjusted chemometric models for C_{et} , C_b and C_g are presented in Figure 1(a) and the fluorescence intensity signals of the chosen pairs can be seen in Figure 1(b).

Figure 1(b) shows that each fluorescence signal pattern of the selected pairs is different from the others. This fact indicates a coherent selection, because repeated signal patterns do not add information to the model. In addition, in Figure 1(a), the good agreement of the chemometric models with the reference models, which was already perceived in the R^2 values, is highlighted.

In Table 3, it can be seen that *Stepwise Regression* failed to give good R^2 values; all of them are smaller than 0.81. Analyzing R^2 as a function of the number of pairs added to the chemometric model (Figure 2), it can be observed that the chemometric models should use at least seven pairs in order to obtain R^2 values similar to the others methods. It clearly shows the inefficiency of *Stepwise Regression* for this application. This result can be ascribed to the high degree of collinearity of the matrix data, a problem that was also pointed out by Jouan-Rimbaud *et al.* (1995a).

Table 3: Result summary for the evaluated methods.

	Pairs in the model	R^2			FO evaluations	CPU Time (s)
		<i>Ethanol</i>	<i>Biomass</i>	<i>Glucose</i>		
PLS	3 ^b	0.9876	0.9984	0.9969	-	0.0006
ES	3	0.9922	0.9961	0.9958	3,307,800	518
Stepwise	3	0.6363	0.7975	0.8072	-	-
Rep60Ind100	3	0.9875 ^a	0.9970 ^a	0.9955 ^a	4510 ^a	0.7922 ^a

a- Mean value of 10 executions.

b- Model with 3 latent factors.

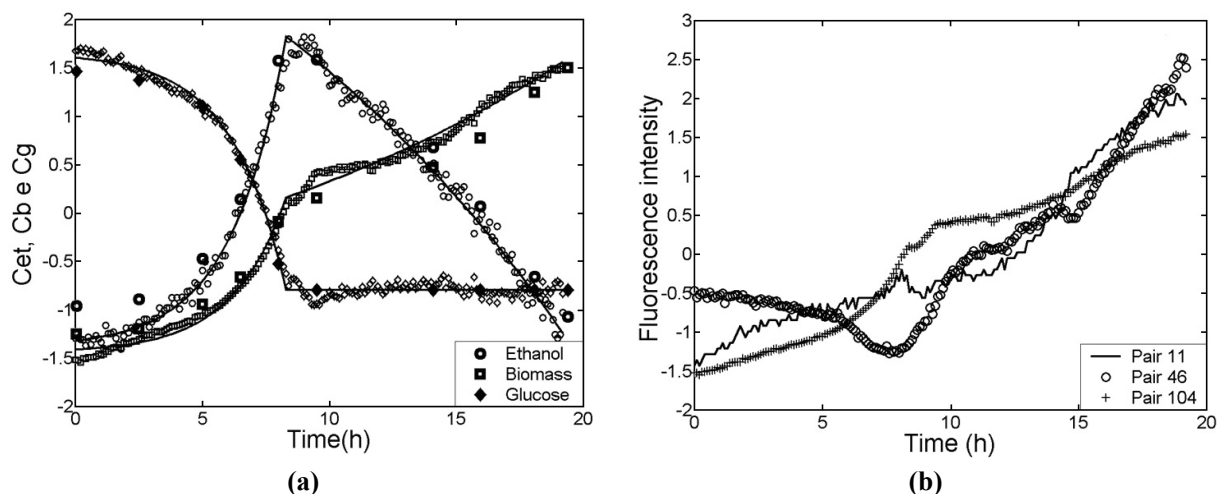


Figure 1: (a) Comparison of chemometric models (lines with open symbols) with reference models (solid lines) and off-line data (symbols); (b) Signal output patterns of the selected excitation/emission wavelength pairs.

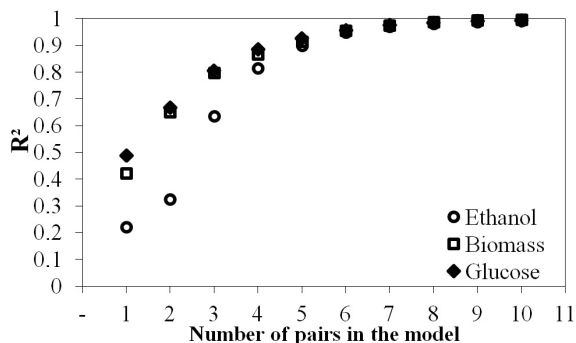


Figure 2: R^2 as a function of excitation/emission wavelength pairs in the models for the *Stepwise Regression* selection method.

The GA set presented in Table 3 (Rep60Ind100) showed the best performance among the four evaluated ones. In Figure 3, the results for the ten executions of all sets can be compared. All sets needed very low CPU times, less than one second, and presented satisfactory R^2 values, always higher than 0.9.

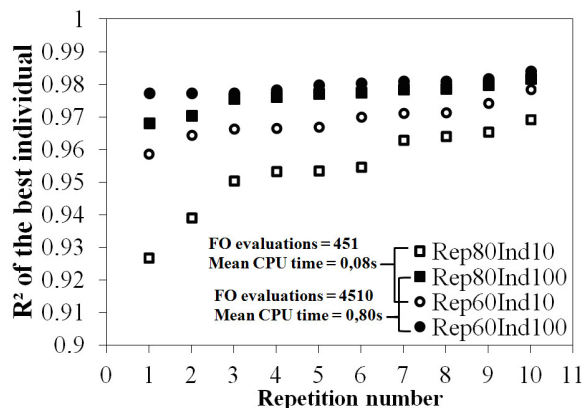


Figure 3: R^2 for the ten executions for the four sets of GA parameters.

Regarding the similarity of the R^2 values, the sets with higher mutation rate demonstrated better performance, while Rep80Ind10 presented the widest range of R^2 values for the final solution. This result may be related to the fact that, while the crossover operation leads the population to converge by making the population alike, the mutation operation reintroduces diversity back into the population helping the algorithm to escape from local optima (Konak *et al.*, 2006). Nonetheless, it is important to mention that the chosen excitation/emission

wavelength pairs were usually different for each Rep60Ind100 repetition, as can be seen in Table 4.

Table 4: Chosen pairs for the ten repetitions of Rep60Ind100.

Repetition	1	2	3	4	5	6	7	8	9	10
FirstPair	10	10	46	11	19	46	47	46	47	33
SecondPair	46	59	52	46	60	76	73	61	73	59
ThirdPair	103	102	102	104	101	95	102	94	102	108

In Figure 4, the evolution of the mean R^2 values of the population can be seen throughout the generations. It is possible to observe that the four sets converged to a solution, meaning that a greater number of generations would probably not enhance the R^2 values of the final solution. This fact indicates that the number of generations was appropriate and could even be decreased, if a reduction in CPU demand is desirable.

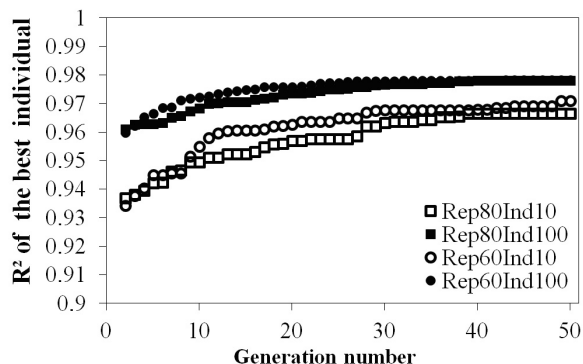


Figure 4: Mean R^2 of the ten executions as a function of the generation number for the four sets of GA parameters.

The better CPU performance of GA may not be relevant in the specific case under study, but this result become quite important when problems with more variables or with more complex prediction models need to be solved. Another example, in which low CPU times are necessary, is the calibration method for bioprocess analyzers based on 2D fluorescence spectroscopy proposed by Oliveira *et al.* (2008). They used only one off-line sample and employed the data of the theoretical model simulation in order to calibrate the chemometric model. At the same time, the results of the chemometric and theoretical model were periodically compared to calibrate the parameters of the theoretical model. This procedure continued in a cycle throughout the cultivation. According to their results, the new method presented better prediction than the usual method using a great amount of off-line analyses for calibration.

In Figure 5, the chosen excitation/emission wavelengths pairs of Rep60Ind100 and ES can be compared by their location in the spectrum. The typical regions of some fluorophores according to Marose *et al.* (1998) are also indicated. In this figure, it can be observed that pair 104, which was selected by ES and whose signal is very similar to the biomass concentration trajectory, lay in the region of riboflavin fluorescence. The good correlation of this

fluorophore, NAD(P)H and tryptophan with biomass growth has already been reported in the literature and indicates coherence in the selection (Bogomolov *et al.* 2011). Pair 46 is in a transition region of fluorescence peaks, probably being tryptophan and pyridoxine. However, pair 11 lies outside of any fluorescence peak in the spectrum. In the same figure, it can be also observed that GA tended to select wavelength pairs in spectral zones near to pair 46, 104 and the NAD(P)H fluorescence region.

The fluorescence signals of all pairs chosen by the ten repetitions of GA are plotted in Figure 6 together with the ones chosen by ES. In this figure, it can be verified that GA, except for repetition 5 and 10, selected three different patterns of fluorescence signals and that these patterns are analogous to the fluorescence signals selected by ES. This behavior can be ascribed again to the high degree of collinearity of the data matrix. In the case of repetitions 5 and 10, GA selected two wavelength pairs with the same pattern of fluorescence signal, but located in different spectral regions (proteins and riboflavin). Even though GA could not always find the optimal solution, in contrast to *Stepwise Regression*, it was able to deal more adequately with the linear dependency of the pairs, presenting satisfactory and coherent results in all the tested cases.

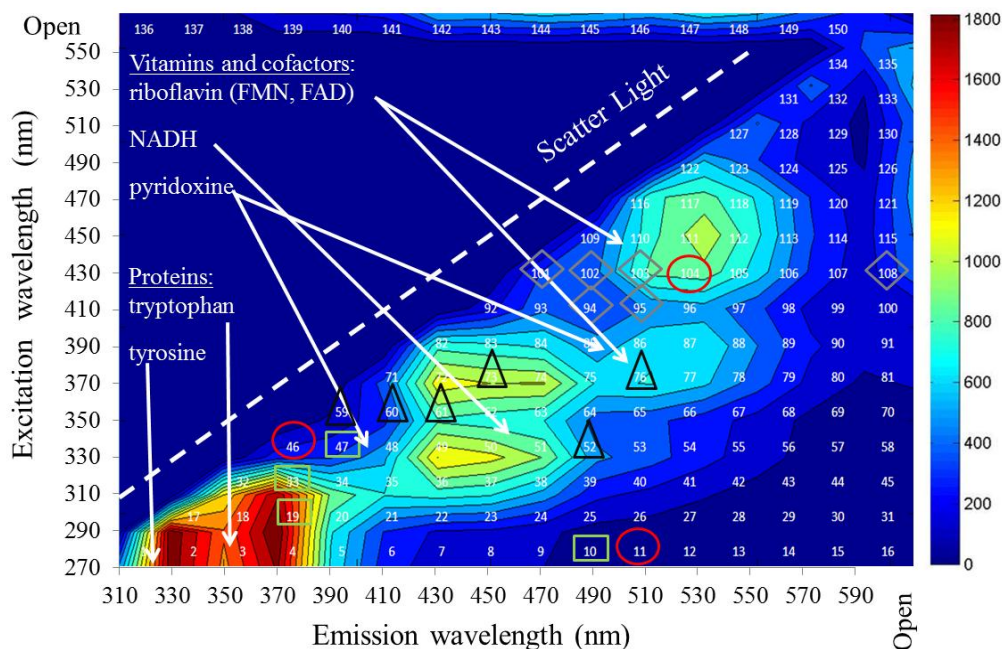


Figure 5: 2-D fluorescence spectrum with excitation/emission wavelength pairs selected by ES (circles). The first, second and third pairs selected by Rep60Ind100 repetitions are indicated with squares, triangles and lozenges, respectively.

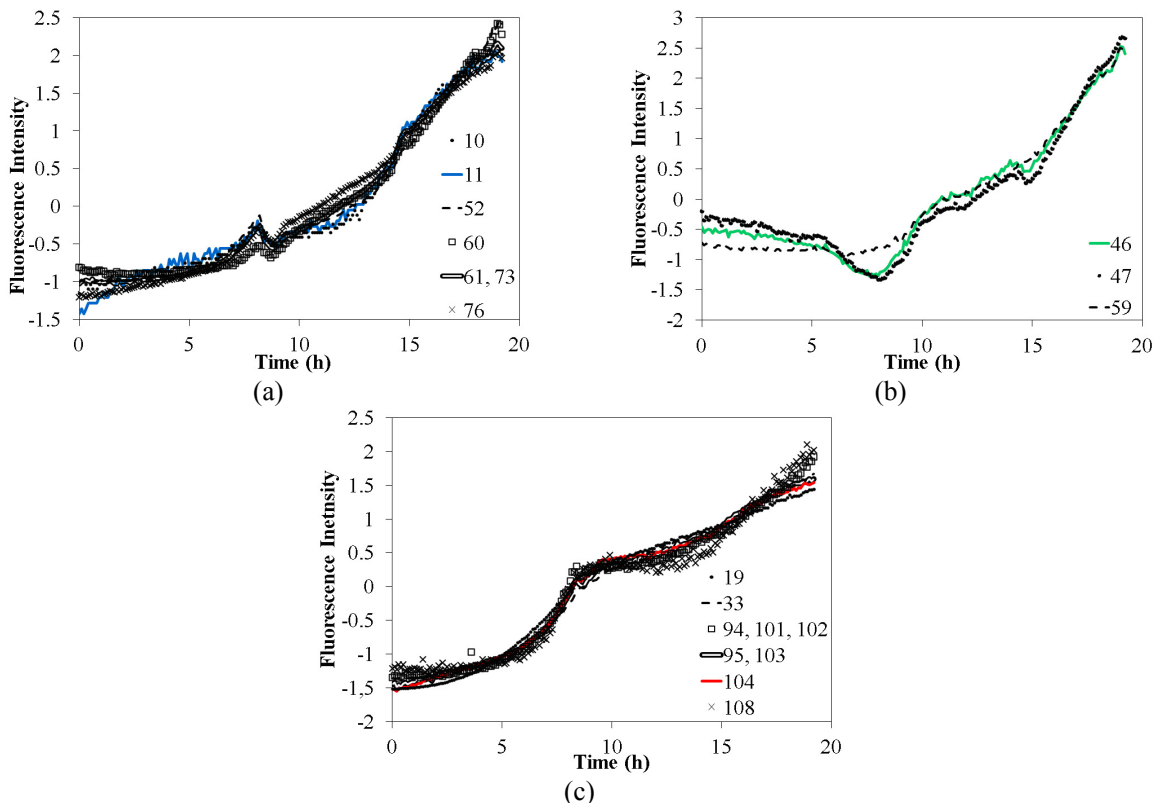


Figure 6: Fluorescence signal of the selected excitation/emission wavelength pairs.

At this point, we can understand the reason why ES selected pair 11 that is outside of any fluorescence peak. For some reason, excitation/emission wavelength pair 11 presented a fluorescence signal analogous to the ones located in the NAD(P)H region. This observation alerts to the fact that the solution of the optimization problem should be confronted with the fluorescence spectrum behavior. For the problem studied in this work, the GA solutions, that selected excitation/emission wavelength pairs near to the fluorescence peak regions, may lead to a more reliable prediction model than the optimal solution of ES.

Comparison to the PLS Technique

In order to compare the performance of the selection methods to the PLS approach, which is a traditional approach to deal with spectroscopic data, the function *plsregress* from the Statistical toolbox of Matlab was used. Since those methods aimed to find the best 3 excitation/emission wavelength pairs, the function was set to use 3 latent factors as well. The results can be seen in Table 3. The PLS technique achieved slightly better models for biomass and

glucose but worse for ethanol in impressively low CPU times.

Considering the results of ES and GA, it can be said that the two methods gave R^2 values comparable to the traditional PLS technique with one big advantage: they do not require the complete spectrum, enabling the construction of small on-line sensors. In addition, because the wavelength pairs selection of these methods is directly based on the process variables, signals related to noise or influenced by physicochemical changes in the reaction medium (e.g., temperature, pH, viscosity, concentration of secondary substances, etc) may be avoided.

CONCLUSIONS

For the problem stated in this work, *Stepwise Regression* presented unsatisfactory performance. It was necessary to add more than six excitation/emission wavelength pairs to the multiple linear chemometric model in order to obtain R^2 values close to the ones found by the ES and GA methods. This fact may be ascribed to the high collinearity of the matrix data. GA could not always find the global

optimum and each call returned a different solution due to the random character of the method. Nevertheless, it was able to achieve solutions with satisfactory R^2 values in low CPU times and few evaluations of the objective function, dealing well with the linear dependency of the variables. The ES method showed the best performance, finding the global optimum in a few minutes. However, the coherence of the optimal solution should be verified by confronting it with the 2D fluorescence spectrum behavior. It also ought to be mentioned that this method could take excessive computational effort for a problem with more variables, for a more complex prediction model or when the optimization is performed in cycles.

In conclusion, it can be said that ES and GA should be considered as good alternatives for solving or at least for pre-screening possible candidates in analogous problems of variable selection. When compared to the traditional PLS technique, these two methods present an advantage. They can provide prediction models based on only a few wavelengths pairs and do not need the complete spectrum, thus enabling the construction of small on-line sensors.

NOMENCLATURE

C_b	Biomass concentration	g.L^{-1}
C_{et}	Ethanol concentration	g.L^{-1}
C_g	Glucose concentration	g.L^{-1}
ES	Exhaustive search	
GA	Genetic algorithm	
NAD(P)H	Nicotinamide adenine dinucleotide phosphate	
Pair 11	Emission 510 nm/ Excitation 270 nm	
Pair 46	Emission 370 nm/ Excitation 330 nm	
Pair 104	Emission 530 nm/ Excitation 430 nm	
R^2	Determination coefficient	
RMSE	Root mean square error	
S	Unexplained variance	
SS_{err}	Residual squares	
SS_t	Total squares sum	
Y_{EB}	Ethanol to biomass yield	g.g^{-1}
Y_{GB}	Glucose to biomass yield	g.g^{-1}
Y_{GE}	Glucose to ethanol yield	g.g^{-1}
μ_E	Specific growth rates in ethanol	s^{-1}
μ_G	Specific growth rates in glucose	s^{-1}

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