

DETAILED CHEMICAL CHARACTERIZATION OF PETROLEUM MIDDLE FRACTIONS BY CHEMOMETRICS ANALYSIS OF THEIR ULTRAVIOLET SPECTRUM

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Chemometric analysis has been applied in this work to correlate ultraviolet spectra of middle distillates and vacuum gas oils with hydrocarbon type analytical data obtained by high resolution mass spectrometry. The use of this methodology reduces the costs of chemical characterization of these types of petroleum fractions and may be applied to the characterization of diesel fuels and gas oils that are processed in fluid catalytic cracking or hydrotreating units.

Keywords: chemometrics, UVVIS Spectroscopy, hydrocarbon type analysis, chemical analysis of petroleum fractions.

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Se presentan resultados de la predicción de la composición detallada de fracciones medias y destilados de vacío del petróleo por tipo de hidrocarburo a partir del espectro ultravioleta visible de estas fracciones. Se aplican métodos quimiométricos para correlacionar los datos espectrales con la composición química por tipo de hidrocarburo determinada por espectrometría de masas de alta resolución. El uso de esta metodología permite reducir considerablemente el costo de caracterización detallada de fracciones medias y gasóleos de vacío; y puede ser aplicado como método de caracterización de diesel y gasóleos de vacío que se utilizan como carga a procesos de craqueo catalítico o de hidrotratamiento catalítico.

Palabras clave: quimiometría, espectroscopia UVVIS, análisis tipo de hidrocarburo, análisis químico de fracciones del petróleo.

INTRODUCTION

Detailed information about the individual components of a petroleum fraction is only possible for light fractions. Gas chromatography is the only analytical technique that allows separating and identifying all the components in a fraction that distills below 200°C (473,15 °F). In heavier petroleum fractions the occurrence of isomers is so great that no chromatographic column can separate all the possible components. Because of these limitations the appropriate way to express the chemical composition of these types of petroleum fractions is in terms of compound families and not of individual components.

Hydrocarbon type analysis by mass spectrometry is one of best methods used to define chemical composition of middle and heavy petroleum fractions. The technique as originally developed was applied to narrow petroleum fractions that could be volatilized in the source of the mass spectrometer.

Low resolution methods are usually applied to fractions of the sample to eliminate the interference between the compound types (ASTM D2786, ASTM D3239), but the Robinson method (Robinson, 1971) is the only low resolution method which can be applied to the complete fraction without any fractionation. High resolution methods allow performing the analysis without requiring previous sample separation (Fisher, 1974; Teeter, 1985; Bouquet, & Brument, 1990; Roussis, & Fitzgerald, 2001). These methods were developed under electron impact that produce a high fragmentation of the molecules but there are also methods which produce low fragmentation (Malhotra, Coggiola, Young, Tse, & Buttrill, 1984).

The results of these methods may be used for determining the potential that a feedstock has to generate selected products in Fluid Catalytic Cracking (FCC) and other petroleum processes. For example, the major reaction which occurs in the FCC unit is fission of paraffinic and cycloparaffinic bonds (this includes dealkylation of aromatic compounds). Aromatic rings are generally not cracked. Thus paraffins, cycloparaffins and monoaromatics are gasoline precursors in the FCC process (Fisher, 1990).

The methodology also could be used for following changes in the composition in of FCC feedstocks that

have been subjected to a hydrotreating process in order to improve their quality (Ng, S., Zhu, Y., Humphries, Zheng, Ding, Gentzis, Charland, & Yui, 2002) (Baldrich & Novoa, 1995).

The analysis by mass spectrometry is, however, very expensive and it is used only for research applications. On the other hand, routine analyses require fast and cheap methods that give the information required for the process engineer.

Ultraviolet spectroscopy is a low cost technique used for monitoring some properties of petroleum products such as the naphthalenes content of jet fuel (ASTM D1840), and the aromatics content in some petroleum fractions (Varotsis and Pasadakis, 1997). These types of applications rely on the strong absorption of ultraviolet radiation by aromatic compounds due to the occurrence of conjugated electrons their structure. (Scheinmann, 1973)

This paper deals with a chemometric approach for estimating, from the ultraviolet visible (UVVIS) spectrum of diesel fuel and gas oils samples, the detailed chemical composition equivalent to that obtained in the hydrocarbon type analysis by high resolution mass spectrometry.

It should be mentioned that in the literature there are very few applications of chemometrics to ultraviolet spectroscopy (UVVIS) of petroleum fractions. One of these applications has been reported by Wentzell, Andrews, Walls, Cooley, and Spencer (1999) who used a chemometric approach using ultraviolet spectroscopy data to estimate the hydrocarbon types in diesel fuel and light gas oil samples. In their research they established a good correlation between the UVVIS spectra and saturates and aromatic compounds contents of the samples determined by supercritical fluid chromatography (SFC) with FID detection. In their study they were able to quantitate three types of aromatic compounds (mono, di and polyaromatics).

This report attempts to correlate the ultraviolet spectra of middle distillate fractions and gas oils with more compound type families with the idea of reducing the time and the cost of a detailed chemical characterization for these types of petroleum fractions.

EXPERIMENTAL WORK

Samples to be used in this research were obtained by direct distillation of crude oils in laboratory distillation units according to ASTM D2892 (atmospheric samples) and ASTM D5233 (vacuum gas oils). Table 1 specifies the samples giving information about cut range in the distillation unit. The crude type from which the cuts were obtained was defined by the characteristics of naphtha and gas oil (Nelson, 1979).

Table 2 contains the chemical composition of all the samples used in this research. Hydrocarbon type analysis is expressed in terms of mono, di, tri, tetra, poly and aromatic sulphur compounds. The high resolution mass spectrometry analyses were performed in a double sector mass spectrometer Autospec VG Ultima. The samples were introduced to the source of the mass spectrometer throughout a Gas Chromatograph (GC) column connected. During the experiment the source was operated under high resolution conditions (dynamic resolution of 7000) at a temperature of 250 °C

Table 1. Behavior of the log-log slope with the dimensionless permeability modulus

SID	BOILING RANGE (°C)	CRUDE OIL	CRUDE OIL CLASSIFICATION
142056	315 - 371	Crude oil 1	Intermediate
157053	315 - 371	Crude oil 2	Intermediate
157054	371 - 427	Crude oil 2	Intermediate
172527	315 - 371	Crude oil 3	Intermediate
172528	371 - 427	Crude oil 3	Intermediate
172738	315 - 371	Crude oil 4	Intermediate
172739	371 - 427	Crude oil 4	Intermediate
173093	315 - 371	Crude oil 5	Intermediate
173094	371 - 427	Crude oil 5	Intermediate
173294	371 - 427	Crude oil 6	Intermediate
174010	371 - 427	Crude oil 7	Intermediate
174670	371 - 427	Crude oil 8	Intermediate
174715	371 - 427	Crude oil 9	Intermediate
181268	315 - 371	Crude oil 10	Intermediate
181316	249 - 315	Infantas L	Intermediate
181317	315 - 371	Crude oil 11	Intermediate
181318	371 - 427	Crude oil 11	Intermediate
181932	315 - 371	Crude oil 12	Intermediate
181933	371 - 427	Crude oil 12	Intermediate
182373	249 - 315	Crude oil 13	Naphthenic tendency
182374	315 - 371	Crude oil 13	Naphthenic tendency
182513	371 - 427	Crude oil 14	Intermediate
183253	371 - 427	Crude oil 15	Paraffinic-Intermediate
183846	315 - 371	Crude oil 16	Intermediate
184819	315 - 371	Crude oil 17	Intermediate
184820	371 - 427	Crude oil 17	Intermediate
174009r	315 - 371	Crude oil 18	Intermediate
182512r	315 - 371	Crude oil 19	Intermediate

Table 1, continue in next page

Table 1. Behavior of the log-log slope with the dimensionless permeability modulus

SID	BOILING RANGE (°C)	CRUDE OIL	CRUDE OIL CLASSIFICATION
173881	371 - 427	Crude oil 20	Intermediate
171520	315 - 371	Crude oil 21	Intermediate-Naphthenic
175913	371.1 - 426.6	Crude oil 22	Intermediate
176095	371.1 - 426.6	Crude oil 23	Intermediate
183251	249 - 315	Crude oil 24	Paraffinic-Intermediate
186006	315 - 371	Crude oil 25	Intermediate-Paraffinic
186510	315 - 371	Crude oil 26	Intermediate Naphthenic
186511	371 - 427	Crude oil 26	Intermediate Naphthenic
187527	249 - 315	Crude oil 27	Intermediate
187529	371 - 427	Crude oil 27	Intermediate
188127	249 - 315	Crude oil 28	Paraffinic-Intermediate
188128	315 - 371	Crude oil 28	Paraffinic-Intermediate
188129	371 - 427	Crude oil 28	Paraffinic-Intermediate
190404	315 - 371	Crude oil 29	Intermediate

(523,15K), the accelerating voltage was 8 KV and the ionization energy was 50 ev. The total chromatographic ion current was averaged for processing the spectra. The mass spectra data was converted to hydrocarbon type data using the Fisher modified matrix (Bouquet and Brument, 1990) included in proprietary software developed by Ecopetrol S.A.- ICP for running hydrocarbon type analysis by high resolution mass spectrometry.

The ultraviolet spectra were obtained in a Hewlett Packard 8453 instrument. The solvent used was spectroscopic grade cyclohexane. The thickness of the cells used was 1 mm. All the spectra were normalized taking into account the mass of the analyzed sample.

The chemometric data analysis was computed with the Unscrambler software V. 9.6 of Camo. To correlate the UVVIS spectral data and the hydrocarbon type information a PLS regression was performed on the data set. To validate the prediction models, full cross validation was employed, and the cross validation was also used to find the optimum number of latent variables to be used in the models.

Additionally, some samples not included in the calibration set were analyzed by this methodology in order to assess the performance of the method. These

samples include those from lab distillation units which were similar to those of the calibration set and some gas oils that had been hydrotreated in a bench scale unit. The last samples were used to test if the methodology could be applied to monitoring chemical changes in the type of processes used by refineries to improve the quality of some feedstocks.

RESULTS

According to the data presented on Table 2 and, as expected, the aromatic content of all the samples increases as these become heavier. Taking into account the crude oil classification, the most clearly observed trend is that naphthenic crude oils are characterized by low paraffin content. There is no additional clear trend with respect to the other hydrocarbon types and crude oil classification.

Figure 1 shows the ultraviolet spectrum of the samples in the region 190 to 400 nm. The spectra that are shown in this figure correspond to the normalized ones taking into account the mass of the analyzed sample and the dilution volume. The graph shows important differences between the different spectrums.

Table 2. High resolution mass spectrometry hydrocarbon type information of standard samples

SID	BOILING RANGE (°C)	PARAFFINS	MONOCYCLO PARAFFINS	DI+CYCLO PARAFFINS	MONO AROMATICS	DI AROMATICS	TRI AROMATICS	TETRA AROMATICS	PENTA AROMATICS	AROMATIC SULPHUR COMPOUNDS	POLY AROMATICS	SATURATED COMPOUNDS
142056	315 - 371	19,00	11,20	28,04	12,29	19,44	6,27	0,97	0,06	2,72	1,03	58,24
157053	315 - 371	16,85	24,26	29,51	11,14	8,25	2,83	0,71	0,09	6,37	0,80	70,62
157054	371 - 427	12,80	21,15	30,54	10,79	8,45	5,18	2,78	0,65	7,66	3,43	64,49
172527	315 - 371	9,15	24,34	29,56	14,59	12,87	3,00	0,57	0,05	5,87	0,62	63,05
172528	371 - 427	6,87	17,86	29,05	13,94	12,69	7,27	3,68	0,55	8,09	4,23	53,78
172738	315 - 371	18,32	21,62	20,24	16,99	14,12	2,63	0,69	0,10	5,27	0,79	60,18
172739	371 - 427	13,27	18,28	22,30	14,18	13,27	5,84	3,50	0,58	8,77	4,08	53,85
173093	315 - 371	18,48	20,34	19,62	11,78	18,34	5,96	0,80	0,07	4,52	0,87	58,44
173094	371 - 427	14,25	15,85	21,48	12,74	13,51	9,09	5,93	0,78	6,37	6,71	51,58
173294	371 - 427	5,14	11,41	25,29	16,12	17,45	9,37	5,34	0,80	9,07	6,14	41,84
174010	371 - 427	8,41	12,05	19,61	15,83	16,02	7,94	4,77	0,92	14,44	5,69	40,07
174670	371 - 427	7,71	18,15	30,90	12,74	10,72	6,14	3,84	0,81	9,00	4,65	56,76
174715	371 - 427	11,42	20,31	30,70	10,90	10,32	5,93	3,30	0,58	6,55	3,88	62,43
181268	315 - 371	9,88	25,21	26,86	15,27	13,21	3,15	0,58	0,03	5,80	0,61	61,95
181316	249 - 315	10,96	32,65	28,59	16,50	9,10	0,35	0,06	0,00	1,78	0,06	72,20
181317	315 - 371	9,53	26,13	27,61	14,14	12,17	3,17	0,72	0,10	6,43	0,82	63,27
181318	371 - 427	6,81	19,14	28,02	13,12	11,69	7,23	4,12	0,56	9,31	4,68	53,97
181932	315 - 371	11,36	12,79	34,22	14,23	16,53	4,86	0,90	0,07	5,03	0,97	58,37
181933	371 - 427	11,34	10,73	28,78	11,13	11,60	8,92	6,50	0,97	10,03	7,47	50,85
182373	249 - 315	2,87	15,97	52,63	19,70	5,37	0,28	0,05	0,00	3,13	0,05	71,47
182374	315 - 371	2,78	12,08	35,33	20,50	16,53	3,04	1,02	0,09	8,63	1,11	50,19
182513	371 - 427	14,77	14,28	27,78	7,95	9,01	9,07	7,57	0,87	8,70	8,44	56,83
183253	371 - 427	12,26	8,08	30,75	12,25	14,49	9,32	6,16	0,60	6,09	6,76	51,09
183846	315 - 371	14,38	10,51	31,92	9,18	19,83	8,14	1,38	0,07	4,58	1,45	56,81
184819	315 - 371	18,93	20,55	19,11	15,49	17,12	4,91	0,54	0,00	3,35	0,54	58,59
184820	371 - 427	13,52	15,69	21,56	13,87	13,17	8,37	5,63	0,77	7,41	6,40	50,77
174009r	315 - 371	14,09	18,72	17,53	18,38	15,01	4,00	1,01	0,11	11,15	1,12	50,34

Table 2, continue in next page

Table 2. High resolution mass spectrometry hydrocarbon type information of standard samples

SID	BOILING RANGE (°C)	PARAFFINS	MONOCYCLO PARAFFINS	DI+CYCLO PARAFFINS	MONO AROMATICS	DI AROMATICS	TRI AROMATICS	TETRA AROMATICS	PENTA AROMATICS	AROMATIC SULPHUR COMPOUNDS	POLY AROMATICS	SATURATED COMPOUNDS
182512r	315 - 371	13,58	11,06	33,18	9,55	19,54	8,26	1,34	0,05	3,44	1,39	57,82
173881	371 - 427	12,95	16,54	21,23	13,84	13,15	7,73	4,66	0,78	9,13	5,44	50,72
171520	315 - 371	7,20	10,80	33,95	16,73	19,13	4,42	1,09	0,11	6,57	1,20	51,95
175913	371,1 - 426,6	12,38	13,73	31,62	12,77	13,45	6,72	3,66	0,41	5,27	4,07	57,73
176095	371,1 - 426,6	10,70	10,41	33,73	13,97	16,23	6,91	3,14	0,39	4,51	3,53	54,84
183251	249 - 315	15,17	11,83	51,47	13,88	6,44	0,22	0,01	0,00	0,98	0,01	78,47
186006	315 - 371	32,92	20,22	13,63	6,58	14,42	6,03	0,78	0,02	5,40	0,80	66,77
186510	315 - 371	3,46	15,81	24,51	19,57	16,64	5,26	1,39	0,11	13,24	1,50	43,78
186511	371 - 427	4,13	12,00	23,48	14,27	14,26	8,85	5,15	0,79	17,06	5,94	39,61
187527	249 - 315	23,84	30,00	20,59	13,13	9,11	0,61	0,09	0,00	2,63	0,09	74,43
187529	371 - 427	16,26	22,31	22,87	11,29	8,37	5,93	3,81	0,69	8,46	4,50	61,44
188127	249 - 315	11,23	20,27	41,56	16,06	9,26	0,26	0,02	0,00	1,33	0,02	73,06
188128	315 - 371	8,95	16,42	34,88	14,79	15,24	4,21	0,67	0,09	4,75	0,76	60,25
188129	371 - 427	8,35	14,55	34,41	10,79	9,87	6,76	4,74	0,86	9,67	5,60	57,31
190404	315 - 371	20,92	23,02	21,98	8,12	15,81	5,58	1,12	0,05	3,42	1,17	65,92

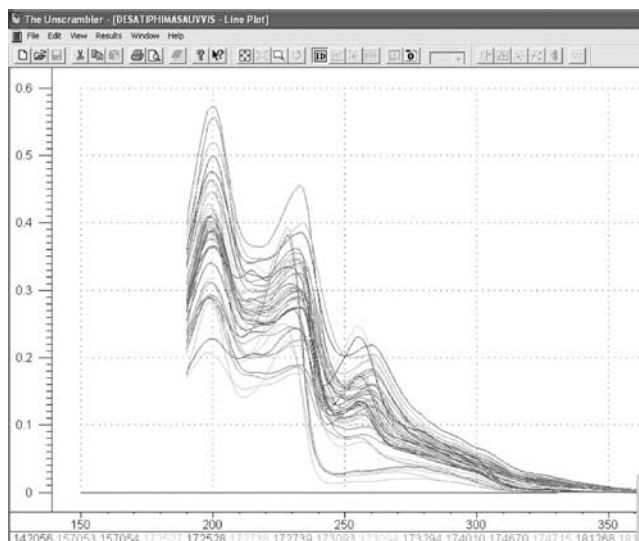


Figure 1. Normalized ultraviolet spectrum of standard samples

The performance of the individual models for each of the compound types is shown in Figures 2 to 8. The predictive models here developed to determine mono aromatics, di aromatics, tri aromatics, tetra aromatics,

poly aromatics (tetra⁺ aromatics), aromatic sulphur compounds and saturated compounds.

In all these graphs there appear the scores plot, the regression coefficient plot, the explained variance plot and measured versus predicted plot. The scores plot is a bi dimensional map of the scores of different cases in two defined principal components. This gives information about some trends in the samples. Inside the ellipse in the scores plot are similar samples in terms of their spectra. Those that lie outside are samples that show differences in terms of their spectra.

The regression coefficients plot summarizes the relationship between the spectrum and the properties. In PLS the regression coefficients could be calculated for any pair of component numbers. The total explained variance plot shows how much of the data variance is explained by each component. The last plot shows the predicted versus the experimental value.

Figure 2 shows the performance of the models for predicting the content of mono aromatics. The model

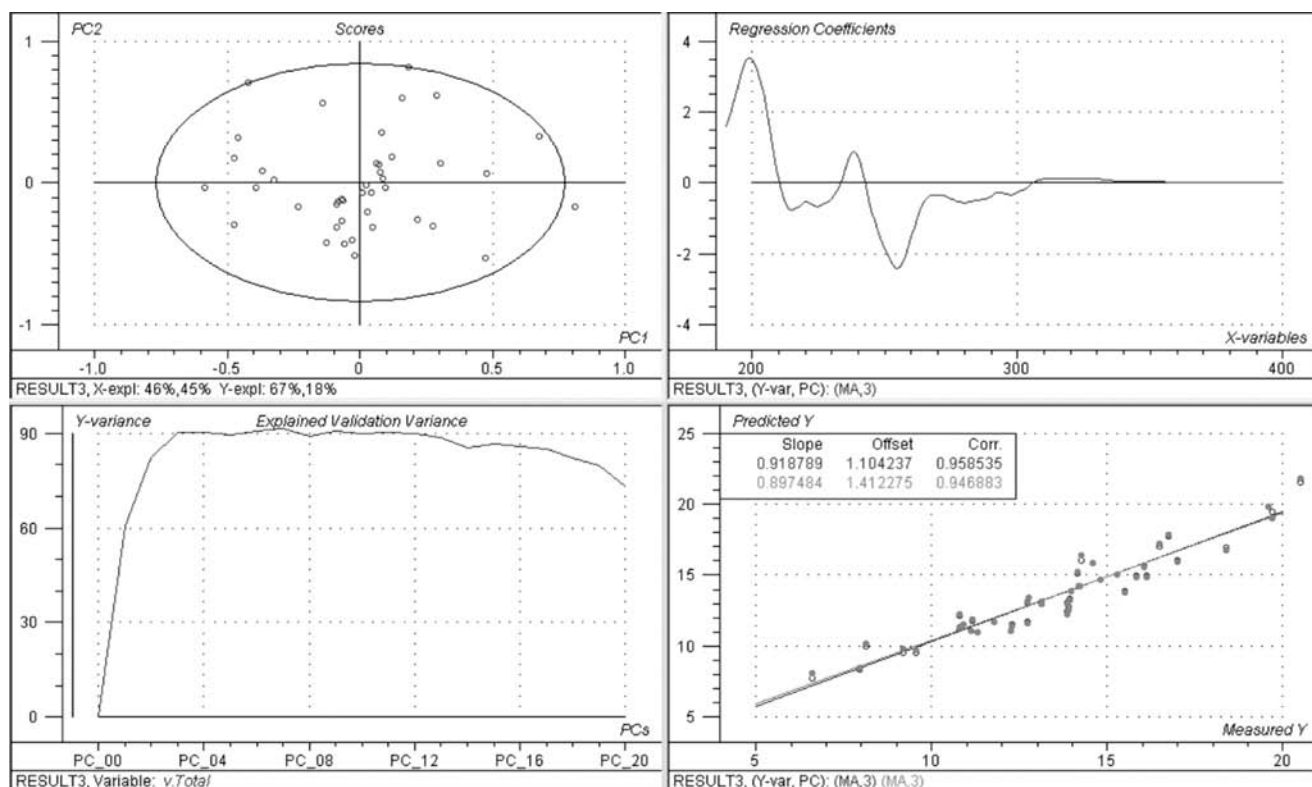


Figure 2. Graphic description of mono aromatics compounds prediction model

uses three components to explain the data variance. All the predicted values are close to the experimental values.

The information presented in Figure 2, shows a good relationship between predicted and experimental data both in calibration and validation. In the case of the third component there is a high positive correlation between the spectral region near 200 nm and 240 nm and a negative correlation between the spectra and the region near 255 nm.

Figure 3 shows the scores, regression coefficients, explained variance and regression line graphics for the content of di aromatic. The model requires 6 components to explain most of the variance of the data. Here, the regression coefficients plot indicates that in the sixth component these types of compounds are positively related to the spectral region around 210 nm, 236 nm, 256 nm and 299 nm. There is a negative relationship for di aromatic content in the spectral region near 200 nm, 228 nm, 244 nm and 272 nm.

Figure 4 shows the model predictions for tri aromatic compounds. In this case 2 components explain most of the variance and the regression coefficients plot indicates a positive correlation with the spectral regions around 261 nm, 240 nm and 214 nm. There is also a negative correlation with the 200 nm and 229 nm spectral regions.

Figure 5 shows the model prediction results for tetra aromatics. The regression coefficients plot indicates positive correlation between the spectral region around 266 nm, 294 nm and 309 nm and the tetra aromatics content. There is a negative correlation of this aromatic type with the spectral regions around 210 and 250 nm. A similar trend is observed with the poly aromatic compounds (see Figure 6).

These results indicate that when the aromatics become more complex they absorb energy at longer wavelengths of lower energy. This trend is to be expected. More complex aromatics are more highly conjugated than less complex ones and are thus more easily excited.

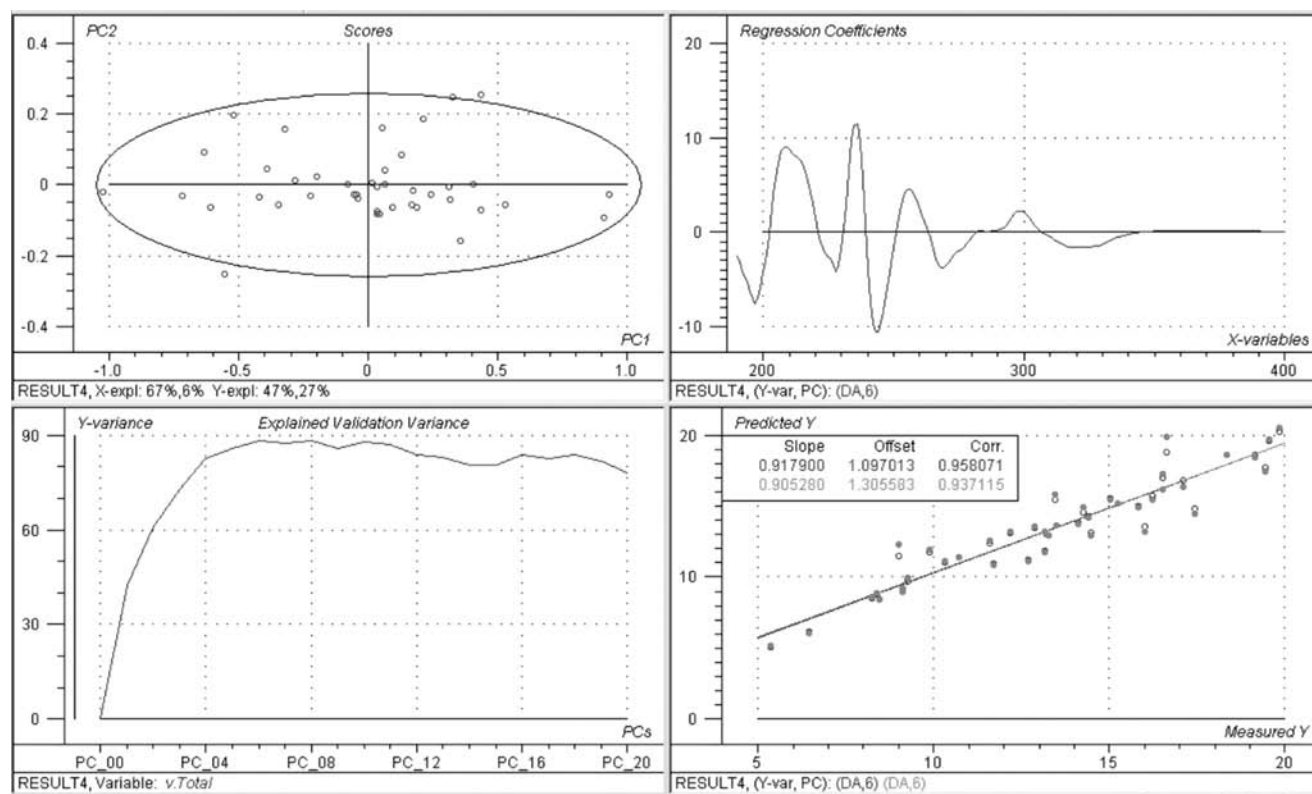


Figure 3. Graphic description of di aromatics compounds prediction model

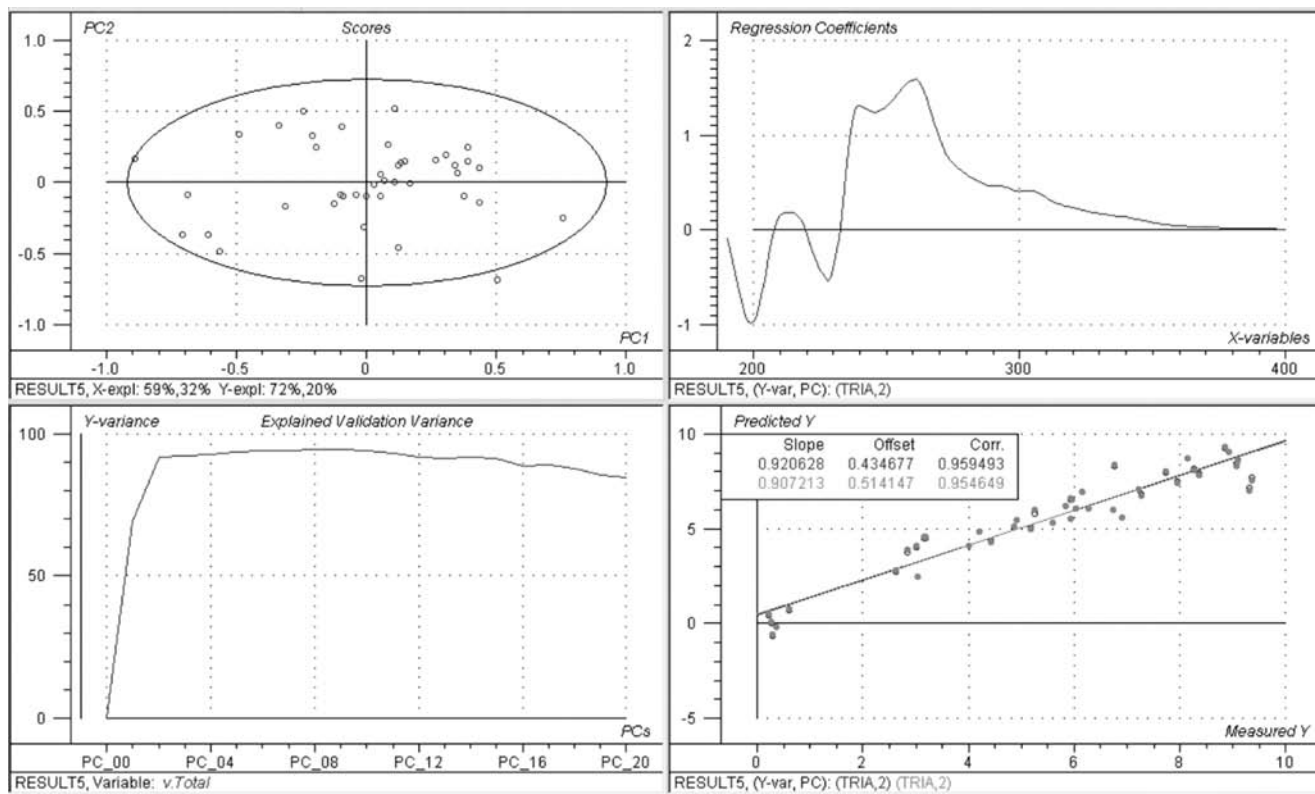


Figure 4. Graphic description of tri aromatics compounds prediction model

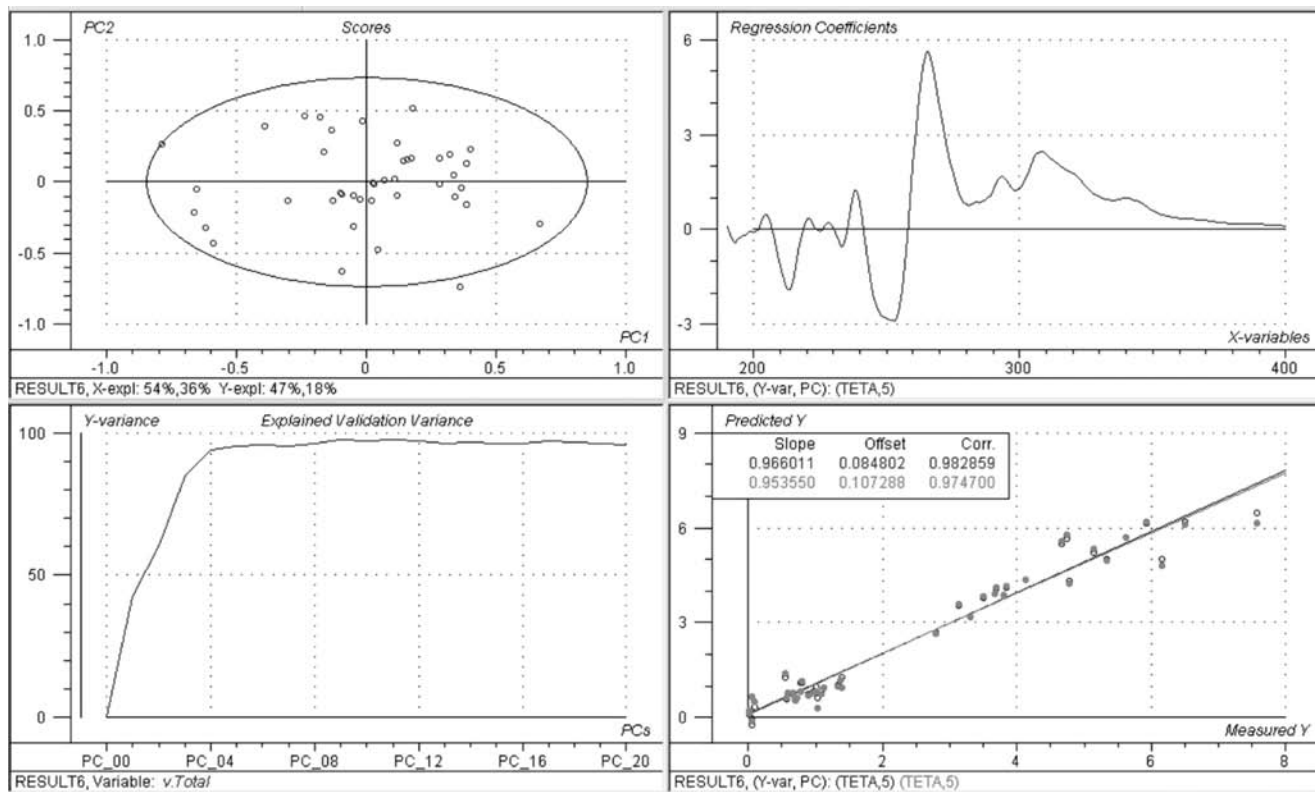


Figure 5. Graphic description of tetra aromatics compounds prediction model

Table 3. Statistical parameters of the predictive models

HYDROCARBON TYPE	PREDICTIVE MODEL			CALIBRATION				VALIDATION			
	REQUIRED COMPONENTS	EXPLAINED VARIANCE IN Y (%)	EXPLAINED VARIANCE IN X (%)	CORRELATION	RMSEC	SEC	BIAS	CORRELATION	RMSEP	SEP	BIAS
Monoaromatics	3	100	95,16	0,9585	0,878	0,888	1,13E-07	0,948	0,98	0,9921	0,0170
Diaromatics	6	88,6	99,64	0,9581	1,05	1,062	6,36E-07	0,9385	1,268	1,283	0,0340
Triaromatics	2	91,98	89,4	0,9594	0,756	0,765	-1,63E-08	0,9539	0,806	0,816	0,0190
Tetraaromatics	5	95,15	97,98	0,9828	0,397	0,402	-3,50E-08	0,9749	0,479	0,485	0,0130
Polyaromatics	4	85,58	92,6	0,985	0,427	0,432	-8,90E-08	0,9753	0,548	0,554	-0,0074
Aromatics sulphur compounds	11	93,7	99,94	0,9921	0,424	0,429	3,24E-06	0,9755	0,753	0,759	-0,0645
Paraffins	4	73,81	94,64	0,914	2,357	2,386	7,32E-07	0,8584	3,032	3,070	-0,0043
Monocycloparaffins	13	59,02	99,98	0,9431	1,869	1,892	-6,70E-06	0,7777	3,837	3,870	0,3160
Di+ cycloparaffins	11	66,72	99,95	0,9671	1,997	2,021	7,49E-07	0,8463	4,225	4,261	-0,3584
Saturated	3	87,33	87,64	0,9543	2,666	2,698	-1,18E-06	0,9445	2,930	2,965	-0,0770

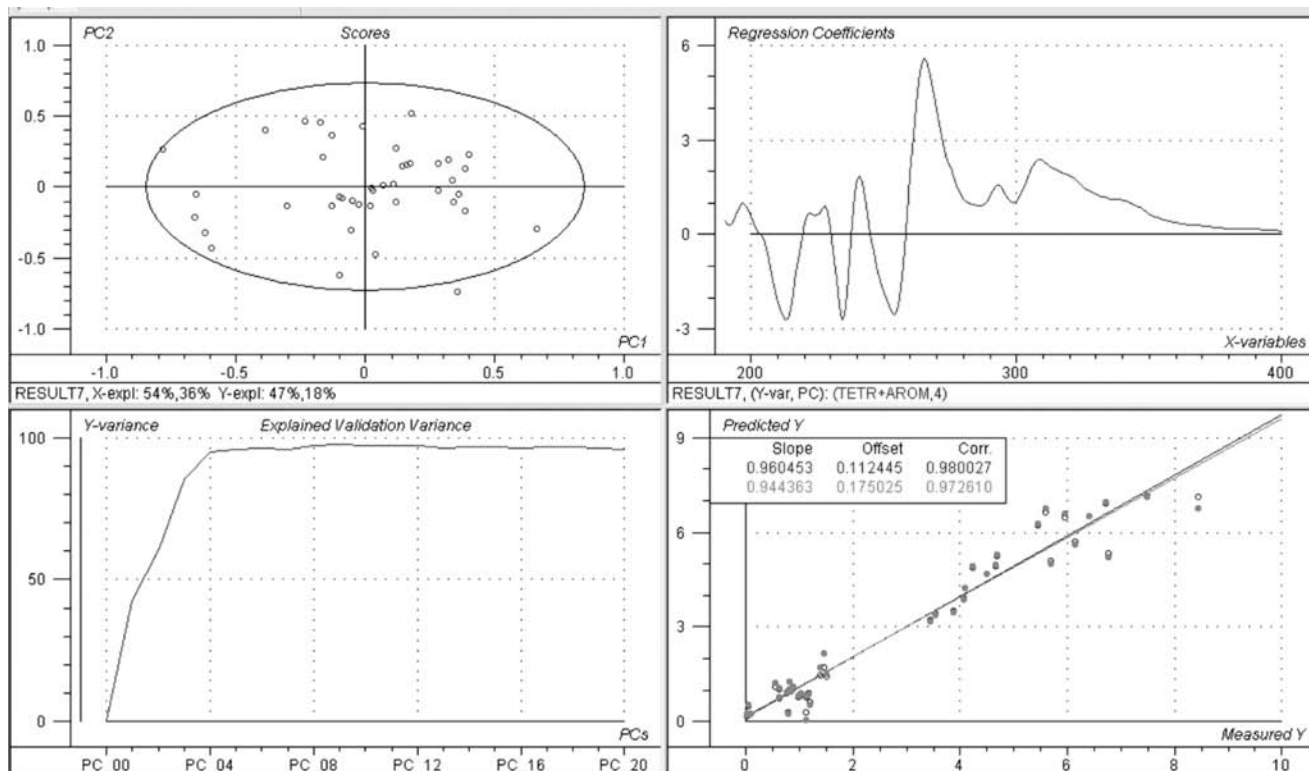


Figure 6. Graphic description of poly (tetra+) aromatics compounds prediction model

Figure 7 shows the model prediction results for the aromatic sulphur compounds. This parameter is positively related for the first component with the spectral region near 235 nm, 245 nm and 268 nm. There is a negative relationship with the spectral region near 200 nm, 238 nm, 254 nm, 262 nm and 279 nm.

Figure 8 shows how the saturates content is positively related for the third component with the spectral region near 223 nm, 216 nm and 255 nm and is negatively related with the spectral region near 200 nm, 239 nm and 266 nm.

All the above information indicates a high interference between the different types of compounds in the UVVIS spectral region. This information is important when developing analytical methods using this technique.

Table 3 summarizes the performance of the different predictive models obtained by correlating the UVVIS spectrum with the chemical composition of the calibration sample set. All the models were obtained without eliminating any of the calibration set samples despite

one or two samples which lie outside the enclosing ellipse. (See the scores plot in figs. 2 to 8).

For each model it is shown the number of components for explaining the variance in the data, the maximum variance explained in x and y variables, the correlation coefficient for calibration and prediction, the standard calibration error (SEC), the standard prediction error (SEP), the bias in calibration and prediction.

The prediction models improve when the component numbers required to explain the observed variance are lower and the explained variance in Y and X is higher. As a result, all the models for predicting aromatics are satisfactory while those which predict the cycloparaffins (monocycloparaffins and di and more complex cycloparaffins) are less so because of the high number of components (more than 10) and the low explained Y variance values (>70%). Also, the models for predicting cycloparaffins and paraffins show a high difference between the calibration and validation errors. These facts indicate that there are some problems in the models.

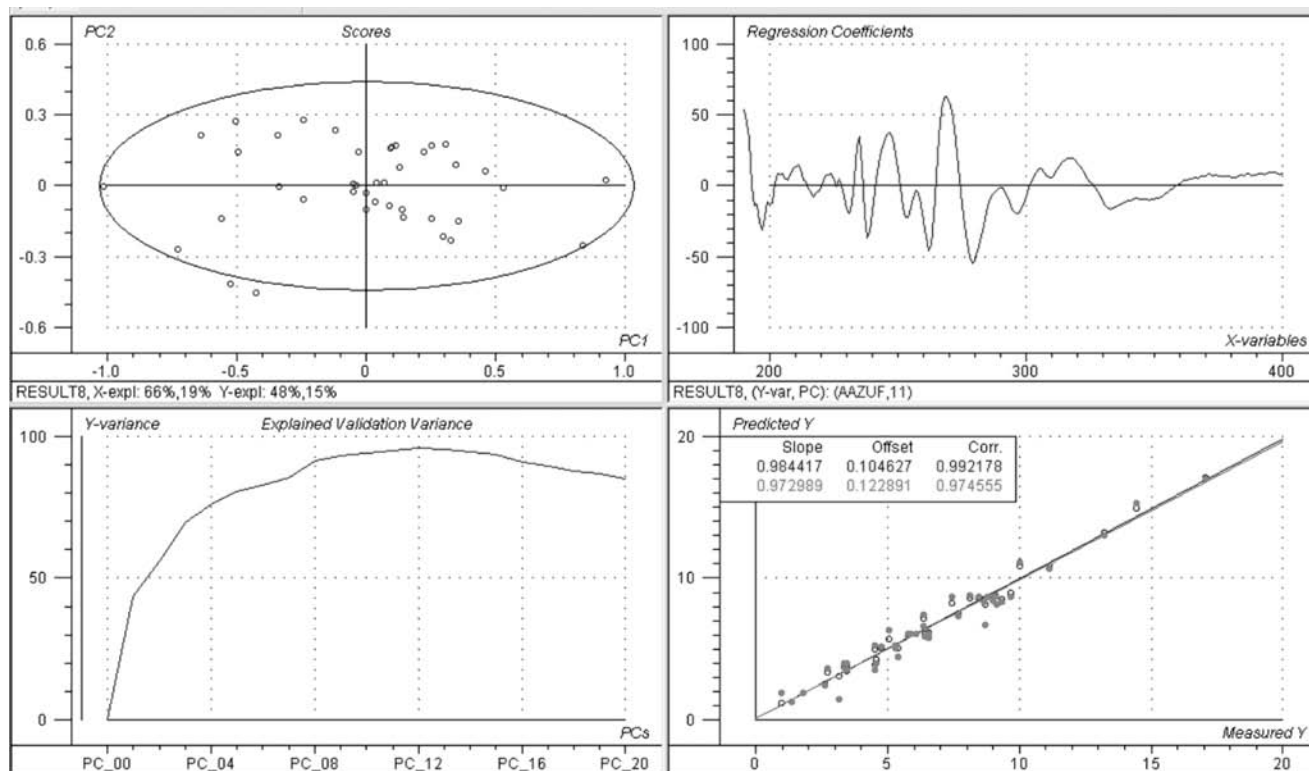


Figure 7. Graphic description of aromatic sulphur compounds prediction model

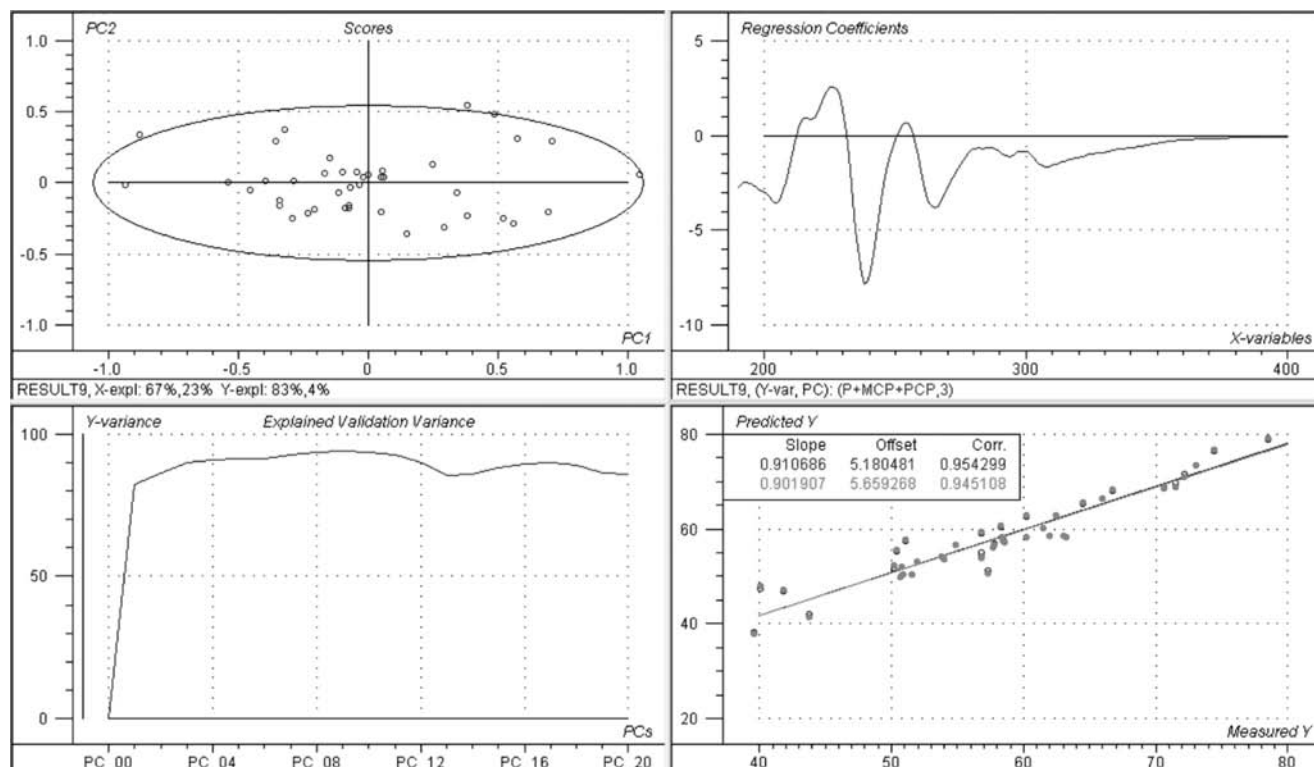


Figure 8. Graphic description of saturated compounds prediction model

Table 4. Performance of the predictive models in the characterization of different petroleum fractions

Sample id	Description	MONO AROMATICS		DI AROMATICS		TRI AROMATICS		TETRA + AROMATICS		SULPHUR AROMATIC COMPOUNDS		SATURATES		GLOBAL BALANCE
		Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	
176093	248-315 °C DIESEL FUEL ABO CRUDE OIL	15,17	0,83	9,08	0,80	-0,03	0,75	0,13	0,41	0,71	0,95	74,71	2,37	99,77
176094	315-371°C DIESEL FUEL ABO CRUDE OIL	14,00	0,90	15,82	1,15	3,33	0,62	0,17	0,58	2,38	0,76	64,38	2,60	100,07
176095	371-426.6°C LIGTH GAS OIL ABO CRUDE OIL	13,24	0,71	15,16	0,78	5,65	0,46	3,17	0,58	4,86	0,44	57,94	2,15	100,02
176096	426.6-482.2°C MIDDLE GAS OIL ABO CRUDE OIL	11,89	0,94	12,96	1,72	7,01	0,54	5,72	0,61	7,55	1,81	56,50	2,93	101,63
176097	482.2-530°C HEAVY GAS OIL ABO CRUDE OIL	12,52	3,14	12,18	7,42	11,10	2,07	13,19	1,52	9,71	6,74	42,00	8,90	100,69
150722	371-426.6°C LIGTH GAS OIL CHIGUIRO CRUDE OIL	14,19	0,74	16,76	0,96	6,58	0,53	4,80	0,50	4,41	0,71	53,13	2,22	99,87
150723	426.6-482.2 °C MIDDLE GAS OIL CHIGUIRO CRUDE OIL	14,58	1,73	15,49	2,84	8,54	1,25	8,97	0,75	8,35	3,38	46,11	4,80	102,04
150724	482.2-530°C HEAVY GAS OIL CHIGUIRO CRUDE OIL	13,88	3,24	13,80	6,97	10,41	2,09	13,17	1,53	6,37	6,70	41,70	9,17	99,33

Table 5. Comparison between predicted and experimental chemical composition data

Sample id	Description	MONO AROMATICS		DI AROMATICS		TRI AROMATICS		TETRA + AROMATICS		SULPHUR AROMATIC COMPOUNDS		SATURATES		Global balance
		Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	
176093	248 - 315 °C DIESEL FUEL	15,17	0,83	9,08	0,80	-0,03	0,75	0,13	0,41	0,71	0,95	74,71	2,37	99,77
	ABO CRUDE OIL													
176094	315 - 371 °C DIESEL FUEL	14,00	0,90	15,82	1,15	3,33	0,62	0,17	0,58	2,38	0,76	64,38	2,60	100,07
	ABO CRUDE OIL													
176095	371 - 426.6 °C LIGTH GAS OIL ABO	13,24	0,71	15,16	0,78	5,65	0,46	3,17	0,58	4,86	0,44	57,94	2,15	100,02
	CRUDE OIL													
150722	371 - 426.6 °C LIGTH GAS OIL	14,19	0,74	16,76	0,96	6,58	0,53	4,80	0,50	4,41	0,71	53,13	2,22	99,87
	CHIGUIRO CRUDE OIL													

Table 6. Following changes in chemical composition in hydrotreating process

Sample id	Description	MONO AROMATICS		DI AROMATICS		TRI AROMATICS		TETRA + AROMATICS		SULPHUR AROMATIC COMPOUNDS		SATURATES		Global Balance
		Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	Predicted	Deviation	
185145	VGO V Calipso Crude oil	10,15	0,98	9,20	1,79	7,14	0,74	6,05	0,55	7,34	2,26	59,56	2,22	99,43
	VGO V Calipso Crude oil	12,81	1,73	9,20	2,09	3,97	2,13	2,27	1,16	5,80	1,70	64,87	3,86	98,93
185146	HYDROTREATED AT 370°C													
	VGO V Calipso Crude oil	12,84	1,66	8,72	1,68	3,83	2,04	2,31	1,00	4,92	1,71	65,44	3,73	98,06
185147	HYDROTREATED AT 390°C													
	VGO VI Calipso Crude oil	9,39	1,31	8,63	2,74	7,67	0,81	7,00	0,76	6,38	3,12	59,33	2,87	98,40
	VGO VI Calipso Crude oil	12,71	1,71	8,89	2,09	4,20	2,07	2,44	1,15	6,44	1,91	64,29	3,81	98,97
185149	HYDROTREATED AT 370°C													
	VGO VI Calipso Crude oil	12,68	1,63	8,56	1,62	4,12	1,96	2,70	1,01	4,88	1,82	64,78	3,67	97,71
185150	HYDROTREATED AT 390°C													

The models for estimating total saturated compounds are better than those for estimating the individual saturated compounds models. Taking into account that saturated compounds do not absorb ultraviolet energy, the acceptable performance of the model for predicting saturated compounds content could be most related to the good relationship between the spectra and the aromatic compounds concentration. As saturated compounds content represents the difference between 100 and the total percentage of aromatics, it is reasonable to observe a good correlation between this difference and the experimental spectra. In general, all the correlation coefficients are high in calibration and validation indicating that the models are sound.

Tables 4 summarizes the chemical composition data predicted from the UVVIS spectra of different cuts obtained by distillation of some crude oils from the lab distillation units according to ASTM D2892 and D 5237. The tabulated data includes the predicted values and the expected deviation. The data presented in this table follow expected trends in the sense that for any given crude oil the heavier the cut the higher the aromatic and aromatic sulphur contents.

Table 5 that shows the predicted chemical composition of the samples that were not included in the calibration set are close to those obtained by mass spectrometry. The observed differences confirm that this method can be used to characterize crude oil fractions. This approach reduces the cost of chemical analysis and gives an alternative tool for obtaining this information.

Table 6 shows the results obtained by application of the method to monitoring the compositional changes in vacuum gas oils during a hydrotreating process. The data show that there is an increase in saturates and monoaromatics and a reduction in the other species as a result of the hydrogen addition. These changes follow the expected trends and the results allow the hydrogen consumption to be calculated from a material balance.

CONCLUSIONS

- All the correlations found in this work indicate that there is a strong relationship between the ultraviolet spectrum of the samples and the different types of compounds defined here.

- From the Ultraviolet spectrum of a middle petroleum fraction (diesel fuel and light vacuum gas oils) it is possible, using a chemometric approach, to predict the concentration of different compositional families such as mono, di, tri, tetra+ aromatics, aromatic sulphur compounds and total saturated compounds.
- The relative errors are low in species of high concentration like mono, di and tri aromatics but may be high for species present in low concentration such as tetra+ aromatics and aromatic sulphur compounds. In spite of this the method enables acquiring fast, cheap and accurate information about the concentration of the different compounds in middle petroleum fractions.
- The chemometric approach proposed here, using the ultraviolet spectrum to determine the chemical composition of petroleum fractions, has applications in the crude oil characterization of middle distillates and light and middle gas oils and could be used for following the trends in hydrotreating heavy petroleum fractions.

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