Experimental variability and data pre-processing as factors affecting the discrimination power of some chemometric approaches (PCA, CA and a new algorithm based on linear regression) applied to \((+/−)\)ESI/MS and RPLC/UV data: Application on green tea extracts

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**A B S T R A C T**

The influence of the experimental variability (instrumental repeatability, instrumental intermediate precision and sample preparation variability) and data pre-processing (normalization, peak alignment, background subtraction) on the discrimination power of multivariate data analysis methods (Principal Component Analysis -PCA- and Cluster Analysis -CA-) as well as a new algorithm based on linear regression was studied. Data used in the study were obtained through positive or negative ion monitoring electrospray mass spectrometry \((+/−\) ESI/MS\) and reversed phase liquid chromatography/UV spectrometric detection (RPLC/UV) applied to green tea extracts. Extractions in ethanol and heated water infusion were used as sample preparation procedures. The multivariate methods were directly applied to mass spectra and chromatograms, involving strictly a holistic comparison of shapes, without assignment of any structural identity to compounds. An alternative data interpretation based on linear regression analysis mutually applied to data series is also discussed. Slopes, intercepts and correlation coefficients produced by the linear regression analysis applied on pairs of very large experimental data series successfully retain information resulting from high frequency instrumental acquisition rates, obviously better defining the profiles being compared. Consequently, each type of sample or comparison between samples produces in the Cartesian space an ellipsoidal volume defined by the normal variation intervals of the slope, intercept and correlation coefficient. Distances between volumes graphically illustrates (dis) similarities between compared data. The instrumental intermediate precision had the major effect on the discrimination power of the multivariate data analysis methods. Mass spectra produced through ionization from liquid state in atmospheric pressure conditions of bulk complex mixtures resulting from extracted materials of natural origins provided an excellent data basis for multivariate analysis methods, equivalent to data resulting from chromatographic separations. The alternative evaluation of very large data series based on linear regression analysis produced information equivalent to results obtained through application of PCA an CA.

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1. Introduction

Extracts from tea leaves (Camellia sinensis) contain a large variety of compounds belonging to various chemical classes [1,2]. Different analytical investigation techniques were used for characterization of such complex mixtures. This includes gas and liquid chromatography, as single or multi-dimensional approaches [3–6], coupled to various detection techniques for unravelling organic compounds patterns as well as atomic absorption or optical emission/mass spectrometry inductively coupled plasma techniques for determination of the multi-elemental profiles [7–9]. Electrochemistry [10], anti-oxidant activity and chelating capacity [11], olfactory properties [12,13], texture properties (through image processing) [14], high resolution melting assay [15], UV [16], NIR [17] and DART-MS [18] spectrometric
the cumulative effects of the sample preparation, intra and interday instrumental variability made impossible the distinction between the compared data. Background subtraction of the (+)-ESI/MS spectra produced an increase of the measured true distances, while for the (−)-ESI/MS spectra the differences are minor. However, one can conclude that the data pre-processing modes do not add any additional discrimination power to the proposed evaluation algorithm.

4. Conclusions

Mass spectra obtained after ionization from the liquid state, under atmospheric pressure working conditions (either with positive or negative ion monitoring) of infused bulk solutions of complex mixtures (resulting from extraction of natural origin materials) are confirmed to represent an excellent data basis for making discrimination among samples, when applying chemometric algorithms, such as PCA or CA. Separation of compounds via a chromatographic method using a “classical” detection approach (such as UV spectrometry) are equivalent to mass spectra obtained from the bulk mixture, from the point of view of the chemometric algorithms. Comparison of chromatograms and mass spectra can be made with respect to the experimental shapes only, without any structural attribution of the compounds in the mixtures (holistic approaches). Grouping of the samples according to different chemometric approaches may be influenced by the sample preparation algorithm or the analytical investigational technique.

Instrumental repeatability and sample preparation induced variability are affecting to a less extent the chemometric algorithms used for discrimination between samples, compared to the instrumental intermediate precision. Mass spectrometric data are more influenced by the instrumental intermediate precision than chromatographic data. Variability issues should be considered and included in the experimental design of the analytical stages, to better assess and validate the dissimilarities between compared samples. It is recommended that data acquisition should be made in a single experimental session. MS experiments have the advantage to be performed faster compared to LC approaches.

An alternative data processing algorithm for discriminating against the samples was proposed. This algorithm is based on the linear regression model applied mutually to experimental data series. Each sample or comparison between two samples is thus represented as an ellipsoidal volume in the 3D Cartesian space, having the slope, the intercept and the correlation coefficient plotted on the three coordinates. These volumes are determined by the normal variation intervals of the slopes, intercepts and correlation coefficients. In fact, these normal variation intervals are equivalent to the experimental pro-

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Appendix A. Supplementary material

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