Alternative Metodology for Gold Nanoparticles Diameter Characterization Using PCA Technique and UV-VIS Spectrophotometry

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Abstract  The rapid developments in nanostructured materials and nanotechnology is starting to have a profound impact in many applications of the biomedical areas, including biomolecules, tissue engineering, and detection of biomarkers, cancer diagnosis, cancer therapy, and bioimaging. Nanotechnology applications hinge frequently on the availability in well-characterized size distributions for gold nanoparticles. In this work we implemented a methodology in order to determine gold nanoparticles size by using the UV-Vis spectra and the multivariate analysis with the principal components' analysis method. The sample size of gold nanoparticles used for correlation of this methodology were previously known (5, 10, 15, 20, 30, 40, 50, 60, and 80 nm), they were supplied by Sigma Aldrich and Ted Pella Inc. All GNPs were measurement with a spectrometer Shimadzu UV-VIS 1800 Rayleigh model over the range from 400 to 700 nm. The UV-VIS spectra was compared with the implemented methodology in order to show excellent behavior and similitude in results, which one suggests this is an outstanding usefully tool to determine the gold nanoparticle diameter.

Keywords  Gold Nanoparticles, UV-Visible Spectra, Principal Component Analysis

1. Introduction

Nanoparticles form a class of materials with diameters in the range from 1 to 100 nm. These systems are produced commonly in nature, and they are called colloids, aerosols and submicron solids; these are dust particles and suspended solids in aqueous systems, smoke, clouds and mist. Other nanoparticles are deliberately made for unique applications in areas of medicine and electronics[1-3]. Fabricated metal colloids have been used for hundreds of years in manufacture of stained glass[4-6]. Gold is a highly functional metal, which has a lot of applications in chemistry and material's science[7] and it is known as one of the best metals to form stable nanoparticles. Recently, nanotechnology has attracted considerable interest in the field of scientific measurement, with a particular focus on gold nanoparticles (GNPs) and their usefulness in biomedical analysis[8-12]. GNPs have shown a particular feasibility for Surface Enhanced Raman Spectroscopy (SERS) where they serve to greatly increase the sensitivity and specificity of Raman classification[13-16].

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It can apply a variety of techniques to characterize the metal nanoparticles intended for a particular use, including spectroscopic methods (UV-visible absorption, Raman scattering) or microscopy (transmission electron microscopy, scanning atomic force microscopy). The precise synthesis of nanoparticles with narrow size dispersion presents particular challenges, the most common diagnostic of size distribution is electron microscopy, which is expensive and time consuming.

UV-visible spectroscopy is widely used in the study of nanomaterials as a diagnostic of nanoparticle formation. Used in conjunction with affinity labeling, UV-visible spectroscopy often provides the means of choice to gauge response in an analysis using nanoparticles. It has been further suggested that the spectroscopic properties of nanoparticles can provide an indicator of their size distribution by fitting the position of the surface plasmon resonance (SPR) to a simple wavelength function[17-19].

In this article, it is suggested a methodology by which principal component's analysis (PCA) can be used to determine gold nanoparticle size from a simple measurement of the UV-visible spectrum. Unlike, previous methods, which rely solely on calibrating the positions of peaks in the SPR spectrum, PCA analysis of variance in the full dimensionality of the absorbance measurement.

The PCA function is an unsupervised procedure to analysis of the inherent multivariate structure of the data. It
reduces the dimensionality of a data set by finding an alternative set of coordinates: principal components (PCs). PCs are linear combinations of the original variables, which are orthogonal to each other and designed in such a way that each one successively accounts for the maximum remaining variance of the data set. Plots of principal component scores against one another can reveal relationships, such as natural data clustering, differentiation and outliers. In addition, plotting the principal component loadings as a function of a given variable can determine whether that variable accounts for a substantial variance.

Important applications relating to imaging processing, data compression, pattern recognition, clustering, classification and time series prediction have developed over last years based upon PCA[20-22]. PCA methods applied with various spectroscopic techniques have grown to become key tools in the biomedical analysis, including techniques capable of assisting in the diagnosis of multiple diseases such as breast and cervical cancer [23-24], and leukemia[25].

2. Experimental Section

It was analysed nine different GNPs samples of known sizes in colloid solution. Table 1 shows these reference products along with technical information.

| GNP Diameter (nm) | Concentration (Particles/ml) | Supplier          |
|-------------------|------------------------------|-------------------|
| 5                 | 5×10^13                      | Ted Pella Inc.    |
| 10                | 5.7×10^12                    | Sigma-Aldrich     |
| 15                | 1.4×10^12                    | Ted Pella Inc.    |
| 20                | 7×10^11                      | Sigma-Aldrich     |
| 30                | 2×10^11                      | Ted Pella Inc.    |
| 40                | 9×10^10                      | Ted Pella Inc.    |
| 50                | 4.5×10^10                    | Ted Pella Inc.    |
| 60                | 2.6×10^10                    | Ted Pella Inc.    |
| 80                | 1.1×10^10                    | Ted Pella Inc.    |

It was taken five spectra of each gold colloid suspension in the range from 400 to 700 nm. These spectrums were recorded with an 1800 Rayleigh spectrophotometer, of 1 nm acquisition resolution. Samples were illuminated by halogen and deuterium lamps. The absorbance measurements were made using 0.5 cm path length quartz cuvettes. Quartz cell containing distilled water served as a reference. Figure 1 shows visible spectra of gold colloids for 5, 10, 15, 20, 30, 40, 50, 60 and 80 nm diameters from bottom to top, respectively. The observed shifts are congruent with the increased nanoparticle diameter[18]. The SPR is clearly visible as a peak in the range between 514 and 550 nm. For small particles, this peak is damped due to the reduced mean free path of the electrons.

These measurements encompassed a total of 45 visible spectrums. In order to have a spectrum's comparator, it was normalized each spectrum to the SPR feature, which in each case represented the maximum. The spectrums were pre-processed by baseline correction with asymmetric least squares smoothing[26-28].

Unlike other algorithms, this one is faster and simpler, even for large data sets, and asymmetric weighting apply everywhere. This algorithm served well to remove noise background and additional external sources of noise.

Spectroscopy techniques for biological samples evaluations are strengthened from this approach to remove fluorescence, shot noise, dark current and spikes from cosmic rays. In Figure 2 is compared a raw spectrum with one normalization and baseline smoothing. All the algorithms for data analysis were implemented in Matlab R2009a commercial software.

PCA offers a way of identifying patterns in sets of correlated data and expressing these characteristics in such a way that highlight their similarities and differences[29]. Main advantage of PCA is that the patterns in data have been identified; they can be compressed by reducing the number of dimensions, without much loss of information[30-31]. PCs are linear combinations of the original variables, which are orthogonal to each other and
constructed so that each one successively accounts for the 
maximum remaining variance of the data set. Plots of 
principal component scores, against one another, reveal 
relationship's existence among the samples, including data 
clustering and segmentation, as well as outliers. In addition, 
the principal components' loadings can be plotted to 
determine which variables cause the greatest variance.

It is often useful to have a comparative measure of the 
width of the variance from the mean along different 
dimensions in a variable space. The covariance offers such 
a measure. The covariance is always measured between two 
component dimensions. If we calculate the covariance 
between one dimension and itself, it is gotten the variance 
in that component. A useful way to assemble all the 
possible covariance values between all the different 
dimensions the variable space for an analysis is to calculate 
them all and put them in a matrix, called the covariance 
matrix. Since the covariance matrix is square, we can 
calculate its eigenvectors and eigenvalues. These are 
important, as they provide useful information about the 
patterns of variations in the data. In general, once 
eigenvectors are found from the covariance matrix, the next 
step is to order them by eigenvalues, highest to lowest. This 
gives us the components or dimensions in the order of their 
significance. With this ordering, we can decide to ignore the 
components of lesser significance. It lost some information, 
but if the eigenvalues are small, it is not lost too much. In 
this paper, it was used the first three principal components.

When it was applied PCA to the data presented above 
pertaining to the visible spectra of gold nanoparticles 
several clusters appeared. Each cluster corresponds to a 
gold nanoparticle spectral group of a particular size. 
Plotting principal component values as a function of gold 
nanoparticle size, we find that we can obtain a function that 
can be used to determine GNP size based on the principal 
component of its visible spectrum extract from PCA applied 
to a set of standard suspensions.

3. Results and Discussions

Applying the Principal Component Analysis to the 45 
visible absorption spectra it was obtained for these nine 
gold nanoparticle diameters from 5 to 80 nm.

The main distinguishing information obtained from by 
PCA is well represented by the first few components.

In figure 3 is showing the behavior pattern based on GNP 
size the first three PCs, where is clearly distinguish ed and 
easily identifies particle size in a GNP suspension by its 
visible spectrum.

This plot suggests that PCA is an excellent technique that 
could be used to discriminate spectroscopically particles of 
different sizes. For the characterization, it was used the 
principal component 1 (PC1), already that presented a 
favourable contribution between the principal component 2 
(PC2) and principal component 3 (PC3).

The above behavior can see in Figure 3 (a, b), there is a 
clear relation between PC1 and the diameters of the gold 
nanoparticles. This is, it is observed that the value of the 
PC1 increases as the size of the gold nanoparticle.

![Figure 3. PCA plots show the patterns for different GNP sizes. The main information obtained from the PCA is described by the first three principal components. a) Principal Component 1 vs. Principal component 2. b) Principal Component 1 vs. Principal component 3.](image)

![Figure 4. The first principal component as a function of the GNP size.](image)

In the figure 4 is shown the first principal component as a 
function of the GNP diameter (red solid line) to 
equation (1). It was found that this behavior in the first 
principal component is congruent with the diameter (dPCA) 
of the nanoparticle by a polynomial function ($R^2 = 0.99$)
The first principal component value. The fit parameters, \( a_1 = 43.72 \), \( a_2 = 2.27 \), \( a_3 = -0.09 \) and \( a_4 = 0.002 \) and the result of fitting to a polynomial of degree 3 between the first principal component and the diameter of the GNP.

In the figure 5 is shown the value graph for the dominant component vs. the position of the Surface Plasmon Resonance peak in the spectrum of each suspension, it is found a systematic variation. This one can be fitted to a third order polynomial function for wave length in nm (\( R^2 = 0.99 \)).

\[
x = b_1 + b_2 \lambda_{SPR} + b_3 \lambda_{SPR}^2 + b_4 \lambda_{SPR}^3
\]

Where \( \lambda_{SPR} \) is the position of the SPR peak and the fit parameters, \( b_1 = 213526.69 \), \( b_2 = -1199.61 \), \( b_3 = 2.24 \) and \( b_4 = -0.0014 \).

Finally, in figure 7 is shown the first principal component 1 vs. the concentration of GNPs, listed in Table 1. The fitting for above description, according to equation (3) is also shown in figure 7.

\[
c = 5.18 + 6.53 \exp\left(-\frac{x}{3.24}\right)
\]

Table 2. Comparisons between GNP diameters: using the surface plasmon resonance peak in the fit function reported by Haiss et al.[17], and principal component's analysis

| GNP Diameter (nm) | \( \lambda_{SPR} \) (nm) | \( d \) (nm) | \( d_{PCA} \) (nm) |
|-------------------|--------------------------|-------------|-----------------|
| 5                 | 514.83                   | --          | 5.6             |
| 10                | 518.47                   | --          | 8.84            |
| 15                | 520.00                   | --          | 12.83           |
| 20                | 523.33                   | 25.53       | 19.66           |
| 30                | 524.63                   | 30.52       | 31.14           |
| 40                | 526.38                   | 36.53       | 39.72           |
| 50                | 531.29                   | 50.14       | 51.75           |
| 60                | 534.88                   | 58.04       | 58.17           |
| 80                | 549.92                   | 81.43       | 81.21           |

Finally, it was shown that the method of principal components analysis can be considered as an alternative tool in order to determine the diameter of GNPs. Diameter, surface plasmon resonance peak and concentration of particles can be calculated as a function of the first principal component. It has also compared these results with a method relying on the assignment of SPR peak position, obtaining excellent results. Unlike this approach, in which the estimate of
diameter depends on the choice of a single spectral position, the present principal component analysis examines absorbance over the full range of the spectrum. In addition, our method determines the particle diameter over the full the range from 5 to 80 nm. Finally, this model will support largely standardized protocols for manufacturing gold nanoparticles and it is considered that it is easily exported to others nanoparticle characterization.

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