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Measuring the relative concentration of particle populations using differential centrifugal sedimentation†

The factors that affect the accuracy and precision of differential centrifugal sedimentation (DCS) for the analysis of nanoparticle concentration are described. Particles are separated by their sedimentation rate and detected using light absorption. In principle, the relative concentration of particles in different populations can be found, but the uncertainty in such measurements is unclear. We show that the most appropriate measurement of particle concentration using this technique is the mass concentration, rather than the number concentration. The relative mass concentration of two discrete populations can be measured with reasonable precision, usually without resorting to complicated data analysis. We provide practical approaches to find the relative mass concentrations for two cases: spherical particles of different materials and agglomerated particles of the same material. For spherical particles made of different materials, naïve analysis of the results can provide relative mass concentrations that are many orders of magnitude in error. Correction factors can be calculated that reduce the error to less than 50%. In the case of agglomerated particles we show that errors of less than 20% are possible and demonstrate, in the case of gold particles, that a combination of UV-visible spectroscopy and DCS enable practical values of mass and number based particle concentrations to be obtained.

Introduction

The unique advantages of nanoparticles can only be realised with sufficient specification of shape, size, concentration, aggregation state and surface chemistry. It is critically important that dispersity in these parameters is understood and controlled. The measurement of absolute concentration is particularly important because this parameter is important to ensure repeatability in processing, performance and, in biological settings, dose. Relative concentrations within a particle population directly impact the measurement of average particle size and agglomeration. Therefore, analytical methods which are capable of measuring both the absolute¹ and the relative concentrations of particles within discrete populations require particular attention and development. Such measurements are used to establish the size distribution of particles, where the

distribution can be thought of as a combination of many discrete populations. Whilst many analytical methods provide a representation of the size distribution of nanoparticles, it is unclear whether they are accurate due to material, size and shape dependent biases.²⁻⁶

Many commonly employed methods are able to provide accurate measurements under the ideal conditions of perfect sphericity, homogeneous composition and monodispersity, but struggle to produce meaningful results if these conditions are not met. For example, the ability to distinguish two populations of different spherical diameters from a monodisperse population of rod-like particles is a particular issue for a number of routine population-based approaches. Another rather common problem in the preparation of particles for biological and sensing applications is the aggregation of particles during processing steps.⁷

One of the most appropriate strategies to overcome this analytical problem is to separate the particles prior to detection and analysis for example, using field-flow fractionation⁸ or analytical ultracentrifugation⁹ with a variety of detection or measurement methods. One of the critical issues is the translation of the detection signal into a useful measure of relative concentration.

If the particle separation process provides highly resolved particle populations and the sample is sufficiently understood then the detection method can be rather simple. In the case of

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differential centrifugal sedimentation (DCS), the typical relative resolution in particle diameter is 5% which, in the case of well controlled particles, is more than sufficient to separate distinct populations of particles. The detection method is usually based upon light attenuation which, provided the refractive indices of the particle components and fluid are known, along with the shape and internal structure of the particles, provides sufficient information to measure the relative concentration of particles in different populations. Whilst the requirement for detailed understanding of the optical properties of the particles seems overwhelming, in many cases these can be reasonably approximated without incurring significant error. Here, we show that the absolute measurement of particle concentration could be rather inaccurate with DCS, but the measurement of relative concentrations between different populations are quite reasonable. If the particle populations are different shapes and sizes of the same material then, usually, the accuracy in the relative mass concentration of the populations can be within 20%.

Theory

In a DCS experiment, the instrument records the intensity of light at a given wavelength transmitted through a liquid medium as a function of time. Here, we consider a particle population with low dispersity (<10%) in size and shape such that useful mean values can be found for parameters such as diameter and extinction cross sections.

The time scale is converted into a particle diameter, D, based upon various densities, viscosities and instrumental parameters that are either measured or assumed. The intensity scale is converted into a total mass per diameter step, $M/\Delta D$, scale using eqn (1), which is justified in the ESI S.1.†

$$\frac{M}{\Delta D} = \frac{1}{\delta} \left[\frac{8\pi R_{\rm f}^2 \ln(R_{\rm f}/R_0)}{3} \right] \left\{ \frac{\pi D^2}{4\sigma} \right\} \rho_{\rm P} A = \frac{\gamma}{\delta} \rho_{\rm P} \frac{A}{Q}$$
(1)

where: A is the measured absorbance; O is the extinction efficiency obtained from Mie theory (the reciprocal of the terms in curly brackets); ρ_P is the particle density; γ is an instrument factor which depends upon the radii of the liquid surface, R_0 , and the detector, R_f , and; σ is the extinction cross section for an individual particle in the fluid at the detector position. Here, we do not consider the effects of multiple scattering, nor the finite collection area of the detector, although the latter effect is taken into account in the manufacturer's software as described in the ESI.† The term δ is a correction factor due to changes in fluid densities and viscosities in the gradient. For dense particles it is approximately the ratio of the viscosity at the detector position to the 'average' viscosity between R_0 and R_f , *i.e.* typically ~ 1.2 . This could be important for absolute measurements of concentration, but for the relative concentrations of two populations both γ and δ generally cancel and are therefore not discussed further.

The total mass of a monodisperse particle population is obtained by integrating the mass scale over its diameter range in the data. By equating ΔD with the infinitesimal $\mathrm{d}D$, eqn (1) can be integrated to provide eqn (2):

$$M \propto \frac{D^3 \rho}{\sigma} A \propto \frac{V \rho}{\sigma} A \tag{2}$$

here, σ is used rather than Q because, as described in ESI S.2,† in the case of non-spherical particles it simplifies discussion to consider the volume of the particle, V, and the extinction cross section.

Eqn (2) is valid if the particle density and extinction cross sections are known. In principle, to correct the measured mass $M_{\rm m}$ into the true mass, $M_{\rm m}$ a conversion factor is required, this is defined in eqn (3), and is also given in ESI S.2.†

$$\frac{M}{M_{\rm m}} = \frac{\sigma_{\rm x} V \rho}{\sigma V_{\rm m} \rho_{\rm x}} = \frac{\sigma_{\rm x} D^3 \rho}{\sigma D_{\rm m}^{\ 3} \rho_{\rm x}} \tag{3}$$

where the subscript 'x' refers to the assumed value, the subscript 'm' is the measured value and unsubscripted terms represent accurate values.

If two particle populations are present in the data, it is possible to determine the relative mass or number concentration by adapting eqn (3). The two populations are designated 'A' and 'B' and their relative mass in the sample $w = (M_A/M_B)$ can be found from their measured relative mass M_m in a single experiment. The relationship is expressed in eqn (4).

$$\frac{w}{w_{\rm m}} = \frac{\sigma_{\rm xA}\sigma_{\rm B}}{\sigma_{\rm xB}\sigma_{\rm A}} \frac{V_{\rm mB}V_{\rm A}}{V_{\rm mA}V_{\rm B}} \frac{\rho_{\rm A}}{\rho_{\rm R}} \tag{4}$$

Different materials

We consider a simple case of two monodisperse populations of spherical particles made of different materials and the data analysed using the properties of particle A. Mixtures of different nanomaterials have been used to generate novel functionality, ¹⁰⁻¹² and the measurement of relative concentration in the mixture is therefore important. In this case, $\sigma_{xA} = \sigma_A$ and $V_{mA} = V_A$. If the Mie cross sections are $\sigma(n_f, n, k, D)$ then, provided these can be calculated, eqn (4) becomes eqn (5), which may be used to convert the measured relative fraction into the actual mass fraction.

$$\frac{w}{w_{\rm m}} = \frac{\sigma \left(n_{\rm f}, n_{\rm B}, k_{\rm B}, D_{m\rm B} \sqrt{\frac{(\rho_{\rm B} - \rho_{\rm a})}{(\rho_{\rm A} - \rho_{\rm a})}} \right)}{\sigma \left(n_{\rm f}, n_{\rm A}, k_{\rm A}, D_{m\rm B} \right)} \left[\frac{(\rho_{\rm B} - \rho_{\rm a})}{(\rho_{\rm A} - \rho_{\rm a})} \right]^{1.5} \frac{\rho_{\rm A}}{\rho_{\rm B}}$$
(5)

in which n is the refractive index of the fluid at the detector position (subscript f) or the type of particle(subscript A or B), k is the relevant extinction coefficient and ρ_a is the average density of the fluid between the liquid surface and the detector.

Dimeric particles

It is commonly observed in DCS that nominally monodisperse particles contain some level of aggregation, evident as distinct populations at defined diameters larger than the primary population. Here we focus upon the relative concentration of dimeric particles (*i.e.* two particles stuck together), and call this population A. If the primary particle population, B, is spherical and the density and refractive index are known then eqn (4) simplifies to eqn (6).

Paper

$$\frac{w}{w_{\rm m}} = \frac{\sigma_{xA}}{\sigma_{\rm A}} \frac{V_{\rm A}}{V_{mA}} \tag{6}$$

As described in ESI S.2,† the shape of the particle changes the frictional force by a factor, χ , known as the dynamic shape factor, which is generally greater than 1. For DCS this results in $V_{\rm A} = \chi^{1.5} V_{m\rm A}$. The measured diameter is different to the equivalent volumetric diameter by a factor $1/\chi^{0.5}$. In DCS, dimers are typically observed at $D_{mA} = 1.20(\pm 0.01)D_{mB}$ implying $\chi \approx$ $1.10(\pm 0.02)$.

These dimeric particles can have a profound effect on nanoparticle size measurements because, in the Rayleigh limit, they have a fourfold higher scattering cross section than primary particles and their apparent size in techniques such as dynamic light scattering (DLS) and particle tracking analysis (PTA) is approximately a factor χ larger than the volumeequivalent diameter, i.e. an apparent diameter ~1.4 times larger than the primary population.

The value of σ_{xA} can be calculated using Mie theory from the apparent diameter of the dimers, D_{mA} . It is not straightforward to obtain σ_A . However, it is possible using, for example, the superposition T-Matrix method13 as shown shortly. Certain limiting cases can be established without such methods.

For non-plasmonic particles in the Rayleigh limit (much less than 50 nm diameter) the orientation-averaged extinction cross section is rather independent of particle shape and σ is proportional to V^y , where y = 1 for particles which absorb light and y = 2 for particles that scatter light. Therefore, in this limit, $w = \chi^{1.5(1-y)} w_{\rm m}$ and the measured mass fraction for particles which absorb light, since (1 - y) = 0 and $w = w_m$, is accurate. For particles which only scatter light, the measured mass fraction of dimers to monomers should be multiplied by $\gamma^{-1.5} = 0.87$ to obtain the correct value, *i.e.* in the data, the relative mass concentration of the dimer population is overestimated by ~15%. Note that if this relative mass concentration were converted into a relative number concentration using the measured volume ratio, $V_{\rm B}/V_{\rm mA}=0.58$, rather than the actual volume ratio, $V_{\rm B}/V_{\rm A}=0.5$, the relative number concentration of dimers to monomers would be overestimated by \sim 15% for absorbing particles and \sim 30% for scattering particles.

For non-plasmonic particles with strong absorbance that are much larger than the wavelength of light, the extinction cross section is proportional to the average area of the particle projected onto a plane. The orientation-averaged projected area of a dimeric particle is 1.165 times larger than its volumeequivalent sphere and eqn (6) becomes $w = (\chi^{0.5}/1.165) =$ 0.90w_m. In this limit, the relative dimer mass fraction is overestimated by \sim 11%.

These analyses suggest that the simplest method of reporting the DCS fractional concentrations of different populations of the same type of nanomaterial is as the mass fraction. In the limits analysed above, the error in using this approach is <20%. Conversion to a relative number concentration requires caution because of general uncertainties in diameter resulting from the unknown shape factor, χ .

To confirm the general applicability of this approach outside the limiting cases described above, accurate extinction cross sections for dimeric particles were calculated using the T-Matrix approach for randomly oriented dimers.13 Fortran code available from a NASA website [http://www.giss.nasa.gov/staff/ mmischenko/t_matrix.html] was translated into MATLAB and eqn (6) evaluated for particles with a range of refractive indexes, n, extinction coefficients, k, and primary particle diameters, $D_{\rm B}$. The MATLAB script is provided in ESI S6.† The wavelength of light was selected to be 405 nm, consistent with some DCS instruments, and the refractive index of the fluid, $n_{\rm f}$, was assumed to be 1.34 at this wavelength and typical fluid composition. These choices are not important for the general conclusions. The T-Matrix results were assessed for convergence and numerical stability by changing the number of expansion functions and confirming that there was a negligible change in the resulting cross sections. The effect of particle separation in the dimer was assessed by performing the calculations at 1 nm, 2 nm and 5 nm separations. For the particles considered here this separation distance has a small, <2%, effect. However for plasmonic particles (typically, n < 0.5 and k >1) it is, as may be anticipated,14 a very serious issue.

In Fig. 1 selected results are shown for particles with $n/n_{\rm f} =$ 1.05, 1.25 and 1.45 (n = 1.41, 1.68 and 1.94 respectively) and k/n_f = 0, 0.01 and 1 representing scattering particles, weakly absorbing particles and strongly absorbing particles. The plots are over the range $D_{\rm B}=10$ nm to 1000 nm and show $w_{\rm m}/w$, i.e. the factor by which the directly measured relative mass concentration of population A to population B is different to the theoretical relative mass concentration.

Fig. 1A shows that this method works well for absorbing particles, with an error of less than 10% and limiting values of $w_{\rm m}/w = 1$ at small sizes and $w_{\rm m}/w = 1.11$ at large sizes, as expected. There is some structure in the curves which depends upon the particle refractive index, but this is a rather small (<5%) effect.

Fig. 1C shows that the method works less well for transparent particles. Here the detailed structure of the scattered intensity causes very large errors in some cases. At small sizes $w_{\rm m}/w = 1.15$ as expected and with $D_{\rm B} < 100$ nm the bias is nearly identical for all particles considered. In this region the curves are closely described by $w_{\rm m}/w = 1 + 0.14 \cos(\pi D_{\rm B}/110)$. For particles which have refractive indexes close to that of the medium, the error remains less than 20% over a wide range of diameters. For the particles with n = 1.68 the error exceeds 20% at $D_{\rm B} = 850$ nm and for n = 1.94 the error exceeds 20% at $D_{\rm B} =$ 500 nm. Therefore, for transparent particles with both a large size and refractive index, relative mass concentrations are highly inaccurate.

Fig. 1B represents weakly absorbing particles. It is clear that these represent a case intermediate between the extremes shown in the other two panels. Scattering cross sections are small at low values of $D_{\rm B}$ and, in this regime, even weak absorption has a strong effect. At large values of D_B the scattering cross section for particles dominates and, especially for particles with a high value of n, the values of $w_{\rm m}/w$ are similar to those in panel Fig. 1C. This extends a previous study of weakly

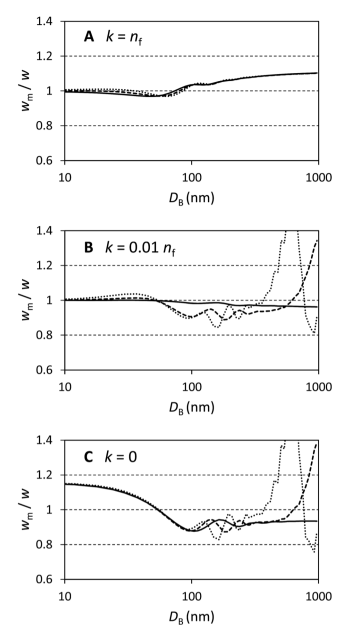


Fig. 1 Ratio of the measured dimer to primary particle relative mass concentration to the true value in DCS for spherical primary particles as a function of primary particle diameter D_B . Solid lines are for particles with n = 1.41, dashed lines n = 1.68, and dotted lines n = 1.94. (A) Materials which strongly absorb light. (B) Materials which weakly absorb light. (C) Transparent materials

absorbing particles with a small refractive index difference to the medium.¹⁵ In that work it was also shown that, for large particles, the error in using volume equivalent sphere scattering cross sections for dimeric particles can be of the order of 10% to 20% and, additionally, that the error increased with the number of particles in the aggregate.

It is important to understand these errors in the context of other contributions to uncertainty encapsulated in eqn (3). Because the volume ratio, $V_A/V_B = 2$, is known, the dynamic shape factor, χ , is known to within 2%, and the assumed density cancels, the important source of uncertainty is the extinction

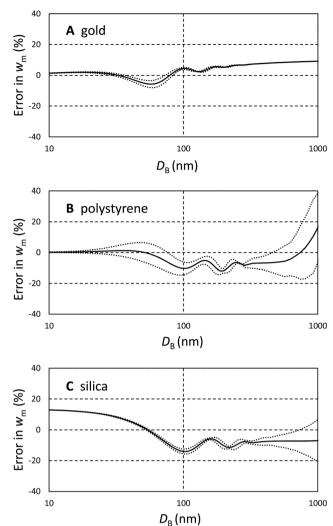


Fig. 2 Error in measured relative dimer to primary mass concentration, $w_{\rm m}$ for three typical materials. The bias due to shape effects is shown as solid lines and the estimated errors due to incorrect size and refractive index are shown as dotted lines indicating ranges either side of the bias. (A) Gold. (B) Polystyrene. (C) Silica.

cross section, σ . Errors in the input parameters to the Mie calculations may, or may not, be important in determining w. One potential source of error is the uncertainty in particle size, possibly resulting from incorrect particle or fluid densities. This error becomes increasingly significant as the density of the particle approaches that of the fluid. The other sources of error are the assumed optical constants for the particles and fluids.

Since these errors relate only to the assumed values of σ , they may be analysed using Mie calculations to find $(\delta \sigma/\delta D)$ and $(\delta \sigma/\delta D)$ δn) using small changes to D and n. To estimate the uncertainty we propagate the resulting values of σ through eqn (3) for both types of particle and establish the change in $w_{\rm m}$, $(\delta w_{\rm m}/\delta D)$ and $(\delta w_{\rm m}/\delta n)$. To estimate the uncertainties due to these factors, the change in $w_{\rm m}$, $\Delta w_{\rm m}$, for a finite change in D, ΔD , is taken to be $\Delta w_{\rm m} = (\delta w_{\rm m}/\delta D)\Delta D$, and similarly for changes in n. We use a 5% error in both n and D and combine the resulting fractional changes in $w_{\rm m}$ in quadrature to provide a description of the Paper

potential magnitude of these sources of error in comparison with the bias calculated above.

The magnitude of these estimated errors are shown in Fig. 2 plotted as dotted lines around the bias. As may be anticipated from Fig. 1A, for strongly absorbing particles such as gold the dominant error is the bias due to shape effects. The errors in diameter and refractive index tend to cancel. We note here that if longer wavelengths (>480 nm) are used to analyse gold, plasmonic effects cause the scattered intensity and absorption to be strongly dependent upon shape and the wavelength of light. 14,16 These prohibit an easy form of quantitative analysis. For polystyrene and silica, errors in diameter and refractive index are more significant, and increase with the refractive index of the particle. So, for polystyrene it is questionable whether a correction for the bias induced by shape is justifiable. However, for silica it is clear that this is still the dominant effect. We note that, for these particles, the error due to refractive index is the dominant effect and is typically 5 times that of a similar relative error in diameter. If an error in diameter arises from an incorrect density difference then, because diameter scales with the square root of density difference, it is clear that even a 10% error in density will usually result in less than 1% error in relative mass concentration. However, it is important to note that here we consider two populations separated in size by only 20% and the errors will be more significant when the populations have a greater size difference.

Materials and methods

Gold nanoparticles with nominal size of 80 nm and 100 nm were purchased from BBI International (Cardiff, UK). The particle manufacturer declared a size of the particles of 78.8 nm and 98.0 nm respectively, as measured by Transmission Electron Microscopy (TEM), and a concentration of 1.1×10^{10} NPs per mL and 5.6×10^9 NPs per mL respectively.

Polystyrene nanoparticles of nominal size 300 nm were synthesized by emulsion polymerization of styrene using sodium dodecyl sulphate (SDS) as surfactant. The polymerization reactions were carried out in a 1 L five-neck reactor equipped with a condenser, a mechanical stirrer, a thermometer and inlets for nitrogen and styrene. 500 mL of ultrapure water eventually containing 0.3 g SDS were introduced into the reactor at room temperature with a stirring rate of 300 rpm, then 50 mL of styrene was added dropwise. The mixture was purged with nitrogen, and nitrogen was fluxed during the entire polymerization procedure. The reactor was heated to 80 °C, then a potassium persulfate aqueous solution (5.0 mL, 0.74 mmol) was added, and the mixture was reacted for 24 h. The obtained latex was purified from surfactant and unreacted monomer by

repeated dialyses against ultrapure water (cellulose membrane, molecular weight cut-off 12 kDa). The particle concentration was determined gravimetrically. In detail, five aliquots of 1.00 mL of nanoparticle suspension were placed in pre-weighed aluminum dishes and dried in a vacuum oven at 80 $^{\circ}\mathrm{C}$ for 24 hours, then weighed to quantify the residual. The solid content is the average of the five measurement and the relative standard deviation is ${\sim}3\%$.

Silica nanoparticles of nominal size 100 nm were provided by Anthoine Thill, CEA. The number concentration was measured by particle tracking analysis (PTA) using an NS500 instrument (Malvern Panalytical, UK) equipped with a 405 nm laser and EMCCD camera. Particles were individually tracked and counted using the manufacturer's software NTA3.2. The estimated uncertainty in the particle number concentration is \sim 10%. The particle concentrations were in good agreement with inductively coupled plasma mass spectrometry (ICPMS) measurements based upon the total silicon content of the particles.

Samples with mixed populations of particles were produced by mixing the original particles samples to obtain the nominal concentrations indicated in Table 1. The nominal concentrations for gold are known to $\sim\!20\%$ accuracy from UV-vis measurements¹⁷ and those of the PS particles are estimated gravimetrically.

The aggregation of the gold particles of nominal size 80 nm was induced by using a surface biotin-streptavidin strategy. One batch of the particles, batch 1, was incubated in a 28.5 µM solution 2,5,8,11,14,17,20-heptaoxadocosane-22-thiol (mPEG-thiol, $M_{\rm w}$ 356.5, Polypure, Oslo, Norway) in 1 mM 3-[4-(2-hydroxyethyl)-1-piperazinyl]propanesulfonic acid (EPPS, Sigma). Another batch of the particles, batch 2, was incubated in a 1 mM EPPS containing 22.8 µM mPEG-thiol and 6.42 µM biotinPEG-thiol ($M_{\rm w}$ 788.0, Polypure, Oslo, Norway). The particle batches were incubated for 2 hours under gentle shaking. The excess thiols were removed via centrifugation. 4 cycles of 1 hour centrifugation of the samples at 500 rcf were followed by gentle removal of the supernatant and redispersion of the particles in fresh 1 mM EPPS buffer at pH 7.8. The two batches of particles were analysed by DCS and the distributions are shown in S4 of the ESI.† Batch 1 particles remain unagglomerated and a low level of aggregation is observed for the batch 2 particles.

Batch 2 was subsequently incubated in Neutralised Chimeric Avidin (NCAvd, $M_{\rm w}$ 14 324) to induce further agglomeration of the particles. The NCAvd was based on a previously described thermostable avidin form, ¹⁸ which was developed by applying charge-neutralized mutations. ^{19,20} The production and characterization of the NCAvd are detailed elsewhere. ^{21,22} For the agglomeration to take place, the particles were centrifuged for 1

Table 1 Name and composition of the samples with mixed particle populations (NPs = nanoparticles)

Sample	Component A	Component B	Nominal [A] (NPs per mL)	Nominal [B] (NPs per mL)
Mix 1 Mix 2	100 nm Au NPs 300 nm PS NPs	100 nm SiO_2 NPs 100 nm SiO_2 NPs	$4 imes 10^9 \ 4 imes 10^8$	$1 \times 10^{12} \\ 1 \times 10^{12}$

Table 2 Description of the composition of the samples exhibiting increasing level of agglomeration

Sample name	Batch 1 (nominal)	Batch 2 (nominal)	Batch 1 (measured)	Batch 2 (measured)	[C] _{NCAvd}
Agg1	100%	_	100%	0%	_
Agg2	67%	33%	64%	36%	0.14 μM
Agg3	33%	67%	34%	66%	0.29 μM
Agg4	_	100%	0%	100%	$0.44~\mu\text{M}$

hour at 500 rcf in presence of 0.87 µM NCAvd. After centrifugation, the sample was gently shaken to disperse the particles.

Four samples exhibiting increasing level of aggregation were produced by mixing the two batches of particles in EPPS buffer in different ratios according to Table 2. The final volume of the particles was the same as the initial volume of the unmodified particles. The total particle concentration is therefore expected to be close to the nominal particle concentration of 1.10×10^{10} NPs per mL, although the centrifugation steps for the removal of excess PEG molecules may have caused some loss of particles. The excess NCAvd was left in solution.

DCS

DCS was performed using a CPS 24000 disc centrifuge (CPS Instruments Inc., Stuart, Florida, USA) equipped with an LED laser emitting light with wavelength between 385 nm and 425 nm and with spectral intensity peak at 405 nm. The instrument was operated at 20 000 rpm with a typical 14.4 mL 8% to 24% (w/w) sucrose gradient of in water (average gradient density $\rho_{\rm f} = 1.064~{\rm g~cm^{-3}})$ if not otherwise stated. This was generated by injections of decreasing sucrose concentration, followed by a final addition of 0.5 mL dodecane as an evaporation barrier. A period of 30 min was allowed prior to measurement acquisition to facilitate thermal equilibrium. A calibration of the instrument was performed before each sample injection by using polyvinyl chloride (PVC) calibration particles with nominal size of 237 nm and density $1.385 \mathrm{~g~cm}^{-3}$ provided by the instrument manufacturer. The uncertainty of these values was independently measured to be 5% and 3.5% respectively for a confidence level of 95%.23 The injected sample volume was measured by weighing the syringe containing the sample before and after each injection and assuming a density of the sample solution of 1 g cm⁻³. Table 3 summarises the values of materials density, refractive index and absorptions used for the data analysis.

Table 3 Material parameters used in the analysis of binary mixtures and agglomerates of nanoparticles. The first fluid values are for the agglomerates and for the Au: SiO2 mix and the second for the PS: SiO₂ mix

Materials	Density ρ (g cm ⁻³)	Refractive index n (405 nm)	Absorption k (405 nm)
Gold (Au) Polystyrene (PS) Silica (SiO ₂) Fluid	19.3 1.052 1.91 1.064, 1.018	1.62 1.594 1.457 1.357, 1.340	1.95 0.02 0

UV-visible spectra

UV-visible spectra were acquired in triplicate using quartz cuvettes using a LAMBDA 850 spectrophotometer (Perkin Elmer Inc., MA, USA). Samples were analysed over the wavelength range 250 nm to 800 nm.

DLS

DLS was used to measure the hydrodynamic radius of the particles and particle agglomerates. These were obtained in triplicate on a Zetasizer Nano ZS 3600 (Malvern Instruments Ltd., Malvern, UK) equipped with a max. 4 mW He-Ne laser, emitting at 633 nm. The measurements were performed in disposable capillary cuvettes (DTS1070, Malvern Instruments Ltd., UK) at 23 °C and preceded by a 3 min equilibration time. The scattered-light intensity-weighted size distributions were expressed in terms of their Z-average (Z-ave) calculated by the instrument software by applying the cumulant method.

Results

Samples with mixed populations

We illustrate that our approach to the analysis of samples containing different materials is justifiable using data from two binary mixtures of particles. These data are shown in Fig. 3 and comprise, in the first case, a mixture of gold (Au) and silica (SiO₂) particles of approximately equal diameter.

In Fig. 3A the data have been analysed using the material properties of gold and in Fig. 3B with those of silica. The second sample is a mixture of polystyrene (PS) and silica particles. In Fig. 3C the data have been analysed using the material properties of PS and in Fig. 3D with those of silica. The various material properties used in this analysis are provided in Table 3. In both cases there is a clear separation of the two populations and the particles have the correct size when their densities are used in the analysis. The results of the analysis of both data sets using, in each case, the two different particle properties are provided in Table 4.

Here, the measured diameter, mass and calculated extinction cross section for particle type 'A' is provided and highlighted in bold. The results of selecting the other type of particle as type 'A' are shown in the next column. Because these particles are close to spherical and the optical properties known, we can expect that data analysis using the correct parameters (i.e. particle 'A') will yield a precise measurement of the injected mass. However, we note that the measured DCS mass is lower than the expected mass of injected particles for silica (100 µg), PS (0.6 μ g) and Au (3.0 μ g). The measured values are a factor of 2 or 3 lower than expected, showing reasonable consistency in

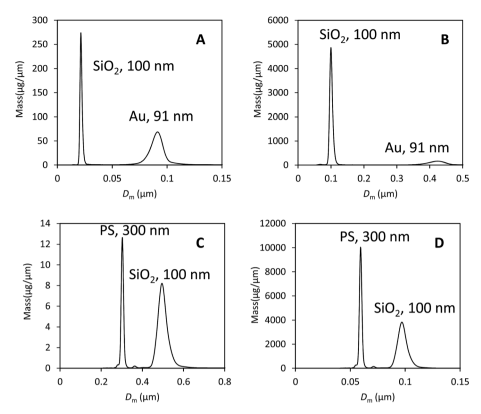


Fig. 3 Binary mixtures of particles analysed by DCS. (A) Gold and silica particles of approximately equal diameter analysed using parameters for gold. (B) Data as panel (A) but analysed using parameters for silica. (C) Polystyrene (PS) and silica particles analysed using parameters for PS. (D) Data as panel (C) but analysed using parameters for silica.

error despite the large difference in particle densities and optical properties. Also in Table 4 are the incorrect measured diameters, masses and calculated cross sections for particle type 'B'.

There are two methods of obtaining the true mass fraction, w, of the different populations. Having analysed the data using two different material parameters, the ratio of the measured injected masses, $M_{\rm A}$, is the most directly useful in a practical context. Alternatively, in each individual experiment, the value of $w_{\rm m}$ can be converted into w using eqn (5). These should give similar values, but are not expected to be identical because the size dispersity of the particle populations has not been accounted for in the second method. In Table 5 the methods are compared and show a reasonable accord in the mass fraction values, w. Comparison of these to the values obtained for $w_{\rm m}$ clearly show that the error can exceed two orders of magnitude

 Table 4
 Single analysis results and Mie extinction cross sections

	Mixture	9						
	SiO ₂ an	and Au SiO ₂ and PS			d PS			
Particle A, B	Au	SiO_2	SiO ₂	Au	PS	SiO ₂	SiO ₂	PS
$D_{\mathbf{A}}, D_{m\mathbf{B}} (\mathbf{nm})$	91	22	100	420	300	500	97	59
$M_{\rm A}$, $M_{m\rm B}$ (µg)	1.0	0.66	54	11	0.18	0.43	39	28
$\sigma_{\rm A}$, $\sigma_{x\rm B}$ (nm ²)	20 000	280	48	25 000	52 000	340 000	42	3.5

if the material densities and optical properties are incorrect. In comparison, the precision in both the modal diameter and $w_{\rm m}$ in DCS is better than 1% in repeat measurements. This is illustrated in S3 of the ESI.†

The method of reanalysing the data for each material is to be preferred, but the correction provided in eqn (5) can be useful if, for example, the raw data are not accessible.

Samples with agglomerated populations

By applying a binding strategy based on the strong affinity of biotin to Avidin we succeeded in producing a set of samples with a defined and stable level of agglomeration. This is shown in S5 of the ESI,† where the high resolution light extinction-based size distributions of the samples measured by DCS are shown as well as DLS data results and UV-visible spectra. As described in S5, standard methods of interpreting the data from UV-visible and DLS data show that the agglomerated nature of samples cannot easily be identified.

The Agg1 sample demonstrates a monomodal DCS peak, enabling the concentration of this sample to be measured with $\sim\!20\%$ accuracy directly from the UV-vis data as 7.3×10^9 NPs per mL. This is approximately two thirds of the starting concentration, implying a significant loss of particles in the preparation procedure. For 80 nm diameter particles, this number concentration translates to 38 μg mL⁻¹ of gold, which may be compared to the integrated DCS value of 15 μg mL⁻¹.

Table 5 Nominal and measured mass fractions, w_m and w

SiO ₂ : Au	w
Nominal	33
$M_{\rm A}$ (SiO ₂ analysis) : $M_{\rm A}$ (Au analysis)	54

	$w_{ m m}$	$w/w_{\rm m}$ (eqn (5))	w (eqn)	(5))
Au analysis	0.66	58	38	
SiO ₂ analysis	4.9	7.9	39	
SiO ₂ : PS			1	w
Nominal			1	170
M _A (SiO ₂ analysis	$: M_{\rm A}$ (PS ana)	lysis)	2	220

	$w_{ m m}$	$w/w_{\rm m}$ (eqn (5))	w (eqn (5))
PS analysis	2.4	110	260
SiO ₂ analysis	1.4	200	280

The disparity is significant, and similar to that noted above in the mixed samples. It may be thought, in this case, to arise from the presence of the surface coating on the gold particles which, in DCS, reduces the density, increasing the sedimentation time and causing the diameter of the particle to be underestimated²⁴ as 71 nm. However, in this case, the extinction cross section of gold is approximately proportional to the volume and eqn (2) indicates that the total mass of gold should be unchanged. The

effect of a coating on the gold also increases the local refractive index around the gold and this will lead to a change in extinction cross section of particles. We note, however, that this will affect both the UV-vis and the DCS results in a similar fashion. Inspection of eqn (1) demonstrates that the DCS value has high uncertainty due to a number of instrumental details (such as detector position) and experimental factors (such as the viscosity gradient) that may bias the results. There may also be other losses during the injection procedure. Therefore the UV-vis results are thought to be more accurate in this case. However, on the basis of the results from mixed particles, it seems reasonable to assume that the relative concentrations found by DCS are representative of the sample.

The concentration of the most agglomerated sample, Agg4, may be analysed in a similar fashion. Although the particle size is not well defined, the value of the UV-visible absorbance at 450 nm, A_{450} , will be approximately proportional to the mass concentration of gold in the sample. Similarly, the total mass of gold may be obtained directly from the DCS analysis and the relative value to the Agg1 sample compared. Here it is assumed that the bias in the DCS results is consistent for all samples. In Fig. 4A and B we compare both of these methods to the values obtained for the monomodal Agg1 sample and the mixed samples.

In Fig. 4A the value of A_{450} is divided by that of the sample Agg1. If the two particle populations do not interact we expect

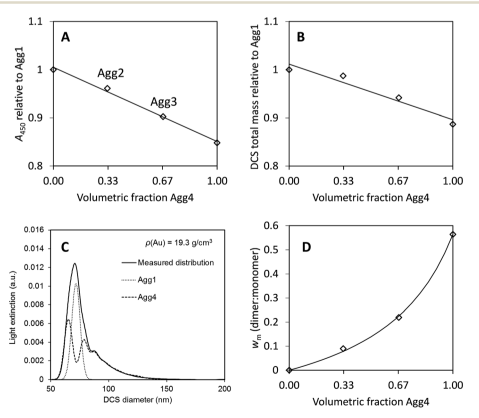


Fig. 4 Results from agglomerated samples. (A) UV-vis absorbance at 450 nm, A_{450} , normalised to the monomodal sample, Agg1. (B) Total mass of particles from DCS normalised to the monomodal sample, Agg1. (C) Fit to DCS data for sample Agg3, using descriptions of Agg1 and Agg4 data. (D) Relative mass of dimers to monomers from DCS, $w_{\rm m}$. Lines are fits to the data, in (A) and (B) they are linear, for D eqn (7) is used with a=3.1.

 Table 6
 Population analysis of Agg4 sample using DCS

Population	Primary	Dimer	Trimer	Tetramer	Pentamer	6+-mer
D_{Veq} (nm), estimated	85	107	123	135	145	154+
χ, observed	1	1.10	1.16	1.18	(1.17)	_
Fractional mass (%)	39	22	15	7	7	10
Fractional number (%)	65	19	8	3	2	<3

this absorbance to be linear with, φ , the volume fraction of Agg4 in the mixture. Here, there is excellent precision and this result suggests that the relative concentration of gold in Agg4 is 85% \pm 2% of that in Agg1. A similar analysis of DCS data shown in Fig. 4B provides a similar result with lower precision, 89% \pm 4%. In both cases, there is some additional uncertainty associated with the concentration of higher agglomerates because their extinction cross sections are unknown.

A further method of analysing the DCS data enables the relative concentration of monomeric particles in the two samples to be assessed. Essentially, we use the Agg1 particles as an internal calibrant in the Agg2 and Agg3 mixtures to find the concentration of the Agg4 sample.

We demonstrate in the ESI, S5,† that the DCS data for Agg4 can be closely matched by a fit describing different aggregated populations. A summary of the relative mass and number concentrations of populations of particles in Agg4 is provided in Table 6. The equivalent volume diameters, D_{veq} , for each population are based upon an estimate of the primary particle diameter of 85 nm, which takes account of the TEM core diameter (79 nm) plus the additional coatings. We note that the monodisperse Agg1 sample, which has a thin PEG coating, has a DLS diameter of 81 nm and the NCAvd molecule is \sim 4 nm in size,²² but is unlikely to completely coat the particles, therefore 85 nm is a reasonable value for the primary particle diameter. The relative diameters in DCS enables measurement of the dynamic shape factor, χ , with reasonable certainty up to the tetramer, beyond this the populations are no longer distinct and therefore the fit is uncertain.

In these fits to the DCS data the primary particle and doublet peaks are sufficiently resolved to measure $w_{\rm m}$ (dimer:monomer) for this sample. Analysis of the data using a variety of different input densities to account for the effect of the coating reveals that these have a negligible effect on the result and we find $w_{\rm m}=0.56\pm0.01$, where the dominant error arises from the quality of the fit.

For samples Agg2 and Agg3, the data can be described by a linear combination of the Agg1 and Agg4 results, as shown in Fig. 4C. These fits provide the measured results given in Table 2, which confirm the Agg1:Agg4 volumetric composition of the samples. Note here that the particle size is incorrect, but the relative mass of gold detected in each population should not be affected by this, as described previously. From the fits, the mass of gold found for the Agg1 population can be added to the monomer population in the Agg4 fit and the value of $w_{\rm m}$ found for each sample. The results are shown in Fig. 4D.

$$w_{\rm m}(\varphi) = w_{\rm m}(\text{Agg4}) \left[\frac{\varphi}{\varphi + a(1 - \varphi)} \right]$$
 (7)

The value of $w_{\rm m}$ can be described by eqn (7), where φ is the volume fraction of Agg4 mixed solution and a is the mass ratio of primary particles in the Agg1 suspension to that in the Agg4 suspension. By fitting the data in Fig. 4D, we find $a=3.1\pm0.1$. Because the mass of gold in the two primary particles are equal, and the UV-vis number concentration for Agg1 is accurate, the number concentration of primary particles in Agg4 is found to be 2.4×10^9 NPs per mL. The dominant error in this result is the ~20% error in the Agg1 number concentration from the UV-vis spectrum. Using the mass distribution in DCS, this permits measurement of the relative concentration of gold in Agg4 to that in Agg1, and we find a value of 84% \pm 2%, which is consistent with the UV-vis estimate.

From Table 6, it is possible to provide two important particle number concentrations for Agg4: The number concentration of primary particles is found from the relative mass fractions as 6.2×10^9 NPs per mL and the number concentration of discrete particles (counting clusters as single particles) as 3.7×10^9 NPs per mL. The former would be relevant to methods which have difficulty distinguishing aggregates, such as SAXS, and the latter would be relevant to techniques such as single particle ICPMS, particle tracking analysis and resistive pulse sensing which count agglomerated particles as single entities.

The average particle size can also be estimated, with the number-weighted average size being 97 nm and the mass-weighted average size as 110 nm. We note that the DLS *Z*-average size of Agg4 is 121 nm. Therefore, if the definition of a nanomaterial is that the average particle size should be less than 100 nm, this sample would be classified as a nanomaterial by some methods, but not by others.

Conclusion

In this paper we describe the factors that affect the accuracy of DCS photosedimentation for measuring the concentration of particles. We demonstrate that the accurate measurement of particle mass concentrations is feasible, but requires substantial knowledge of the instrumental and experimental details as well as the physical properties of the particles. We find that, in cases where there is a reliable independent measurement of particle concentration, that the DCS measurement underestimates the mass concentration by a factor of 2 or 3. The origin of this discrepancy requires further investigation. By assuming that this error remains constant during a single experiment

regardless of particle type, we show that the measurement of relative mass concentrations of different particle populations is feasible with reasonable accuracy. For particle populations of different types, it is possible to correct the data to provide a reasonable estimate of the relative mass concentrations and show, using example mixtures, that these estimates are consistent to within a factor of 2. This compares to order of magnitude errors without such corrections.

For different particle populations of the same type of material, we demonstrate that the relative mass concentrations are reliable. For particles that are weakly aggregated the relative mass concentrations of the dimeric population compared to the monomeric population have a typical error of less than 20% over a wide range of particle size. Furthermore, this measurement is insensitive to small (\sim 5%) errors in the assumed refractive indexes and densities of the particles. This analysis is demonstrated on an agglomerated gold sample, where calibration using a non-agglomerated internal reference enables an accurate measurement of the particle number concentration, both in terms of primary particle number and the number of discrete particles.

Conflicts of interest

There are no conflicts of interest to declare.

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