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Use of flow field-flow fractionation and single particle inductively coupled plasma mass spectrometry for size determination of selenium nanoparticles in a mixture†

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Various analytical techniques have been used for size analysis of selenium nanoparticles (SeNPs). These include flow field-flow fractionation (FIFFF), single particle inductively coupled plasma mass spectrometry (SP-ICP-MS), dynamic light scattering (DLS) and transmission electron microscopy (TEM). For hydrodynamic diameter estimation, the FIFFF technique was used and the results were compared with those analyzed by DLS. For core diameter estimation, the results obtained from SP-ICP-MS were compared with those from TEM. Two types of FIFFF channel were employed, *i.e.*, symmetrical FIFFF (Sy-FIFFF) and asymmetrical FIFFF (Asy-FIFFF). Considering the use of FIFFF, optimization was performed on a Sy-FIFFF channel to select the most appropriate carrier liquid and membrane in order to minimize problems due to particle membrane interaction. The use of FL-70 and 10 kDa RC provided an acceptable compromise peak quality and size accuracy for all samples of SeNPs which were coated by proteins (positively charged SeNPs) and sodium dodecyl sulfate (negatively charged SeNPs). FIFFF always yielded the lower estimate of the hydrodynamic size than DLS as a reference method. The results obtained by SP-ICP-MS were consistent with the TEM method for the core diameter estimation. The results from FIFFF and the DLS reference method were significantly different as confirmed by paired *t*-test analysis, while the results provided by SP-ICP-MS and the TEM reference method were not significantly different. Furthermore, consecutive size analysis by SP-ICP-MS for the fractions collected from FIFFF was proposed for sizing of SeNP mixtures. The combined technique helps to improve the size analysis in the complex samples and shows more advantages than using only SP-ICP-MS.

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Introduction

Nanoparticles have been used in wide applications owing to their unique chemical and physical properties such as large surface area and nanoscale size.¹ Selenium nanoparticles (SeNPs) have gained attention due to their functions of having anticancer,² and antimicrobial properties,³ as dietary supplements,⁴ and in food packaging.⁵ The synthesis and characterization of various functional inorganic nanoparticles by different techniques to provide chemical composition, morphology, and size have been reported in the literature.^{6–10} According to the European Commission, not only the size of particles but also the size distribution should be considered when dealing with the hazard or risk of nanomaterials to

human health and environment.¹¹ Therefore, various instruments have been used to provide the particle size and size distribution of NPs such as dynamic light scattering (DLS),¹² transmission electron microscopy (TEM),^{13,14} and field-flow fractionation (FFF) coupled with inductively coupled plasma mass spectrometry (ICP-MS).^{15,16}

Each characterization technique provides a different meaning of “diameter”. For example, DLS is used to determine the average intensity of brownian nanoparticles in colloidal suspension and recorded as “hydrodynamic diameter (d_h)”.¹⁷ This technique is generally used for engineered nanoparticles characterization including synthesized SeNPs characterization to control the growth of NPs from agglomeration.¹⁸ Transmission electron microscopy (TEM) is known as a powerful imaging method of NPs presented in the sample. By using computer imaging analysis software, the particles can be counted for size characterization. The core diameter information is obtained. However, the correctness of size analysis by TEM depends on the contrast of the TEM image.¹⁹ The area-equivalent size obtained by TEM is also limited by the number of measured particles. By using this technique, the

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aggregation of particles contributes to the erroneous size analysis.¹⁶ Therefore, an alternative technique for size based separation and quantification such as FIFFF or single particle ICP-MS is required.

Flow field-flow fractionation (FIFFF), a separation-based method, has been reported for hydrodynamic size determination of nanoparticles or colloidal particles, both as synthesized and as in samples containing complex matrices.^{4,20,21} Two different types of FIFFF are available including symmetrical flow field-flow fractionation (Sy-FIFFF) and asymmetrical flow field-flow fractionation (Asy-FIFFF). FIFFF deserves special attentions for nanoparticles characterization due to their high separation performance.^{22,23} In order to provide robust protocol and high reliability for inter-laboratory comparison of NPs size measurement, size characterization of SeNPs was tested by using two different techniques of FIFFF including Sy-FIFFF and Asy-FIFFF.

With FIFFF, the retention time is used to calculate the particle size (hydrodynamic diameter, " d_h ") of nanoparticles according to the Stoke's law theory. Nonetheless, the retention time of nanoparticles might be shifted due to the occurrence of particle membrane-interactions which is being a challenge in FIFFF, resulting in invalid size calculation of nanoparticles. This phenomenon has been investigated depending on some factors such as type of membrane material, carrier liquid, stabilizing agent of particles and operating condition, for example cross flow rate.^{24–30} Saenmuangchin and Siripinyanond²⁶ estimated the hydrodynamic diameter of AuNPs with different types of surface coating agent by using Sy-FIFFF. The effect of various types of surface coating agents: *i.e.*, electrostatically stabilizing agent, *e.g.*, tannic acid (TA) and citrate (CT); and sterically stabilizing agents, *e.g.*, polyethylene glycol (PEG), polyvinylpyrrolidone (PVP), and branched polyethylene imine (BPEI) in different membrane materials and carrier liquids to the retention behavior of AuNPs was observed because of the strong attractive interactions between particles and membrane as discussed by Meisterjahn *et al.*²⁴ The polarity and hydrophobicity of coating agents have a critical influence on the variability of retention time and relative recovery.³¹ Hence, two types of coating agents (protein and SDS), which resulted in SeNPs of different charges, were used to examine the performance of FIFFF as to expand the function of FIFFF in wide range of nanoparticles analysis.

Single particle inductively coupled plasma mass spectrometry (SP-ICP-MS) is known as an emerging technique for characterization, quantification and identification of nanoparticles.^{32,33} SP-ICP-MS offers the information on the number of particles detected per time and their mass concentration.³⁴ The diameter information from this technique is recorded as " d_{mass} ". This technique has been applied for SeNPs at low concentrations in commercial Se-rich yeast.³⁵ For complex samples, particle size analysis is still a challenge due to the presence of unaccounted forms of metal. Therefore, SP-ICP-MS has been used in combination with another analytical approach such as Asy-FIFFF as reported by Loeschner *et al.*³⁶ for analysis of AgNPs in chicken meat. As the analysis of the particle

size in the complex sample such as synthesized NPs is still a challenge, an alternative technique is required.

Overall aims of this work are focused on the application of Sy-FIFFF under optimum condition to determine the hydrodynamic diameter of various types of SeNPs compared with Asy-FIFFF and SP-ICP-MS to determine the core diameter of SeNPs; and SP-ICP-MS analysis after size fractionation by Sy-FIFFF for core diameter determination of SeNPs in the mixtures. TEM and DLS are selected as reference methods because both methods are known as routine technique for nanoparticle size analysis. Use of combined techniques of Sy-FIFFF and SP-ICP-MS was demonstrated for size analysis of synthesized SeNPs size mixtures as a model of complex sample.

Experimental

Reagents and materials

The deionized water used in this study was obtained from a water purification system (18.2 M Ω cm⁻¹) (EASY pure@II model D7031, Barnstead Thermolyne Corporation, Dubuque, Iowa, USA). Two protocols were used for SeNPs synthesis. In the first protocol, the following chemicals were used: 0.01 mol L⁻¹ of sodium dodecyl sulfate (SDS) (Fluka Steinheim, Germany) as stabilizing agent; 520 mmol L⁻¹ solution of sodium thiosulfate (Sigma-Aldrich, St.Louis, Missouri, USA) as reducing agent; and 5.2 mmol L⁻¹ of selenium(IV) dioxide (Alfa Aesar, Massachusetts, USA) as precursor. In the second protocol, the reagents for preparation of SeNPs (protein as stabilizing agent) was 30 mmol L⁻¹ of sodium selenite as precursor, 2.5% (w/w) of β -lactoglobulin from bovine milk and 2.5% (w/w) albumin from bovine serum as stabilizing agents. All chemicals used in this second protocol were purchased from Sigma-Aldrich, St.Louis, Missouri, USA. 300 mmol L⁻¹ of L-ascorbic acid (Fisher Scientific UK Limited, Leicestershire, UK) was used as reducing agent. FL-70 was purchased from Fisher Scientific UK Limited, Leicestershire, UK (composition: 1.4% tetrasodium ethylenediaminetetraacetate, 0.5% sodium oleate, 0.1% sodium bicarbonate, 2.7% sodium carbonate, 3.8% triethanolamine oleate 3.8%, 88.8% water and others). Sodium azide (NaN₃) was purchased from Merck, Darmstadt, Germany. Commercial gold nanoparticle standards (AuNPs) 20 nm, 60 nm, and 80 nm (Nanocomposix, Inc., San Diego, California, USA) were used to check the sensitivity of SP-ICP-MS and test the performance of Sy-FIFFF system. Commercial polystyrene (PS) nanoparticle standards 20 nm, 40 nm, 60 nm and 100 nm (Sigma-Aldrich, St.Louis, Missouri, USA) were used for checking the performance of Asy-FIFFF system. 0.1% HNO₃ (Merck, Darmstadt, Germany) was used for preparation of selenium standard solution in solution mode ICP-MS measurement. The pure 99% methane gas (S.I.Technology Co.Ltd., Bangkok, Thailand) was used as a reaction gas.

SeNPs synthesis

SeNPs were prepared by using chemical reduction method with two different protocols, as shown in Fig. 1. For the first protocol which is SDS stabilized, sodium thiosulfate was used as the reducing agent. Both reducing agent (520 mmol L⁻¹ sodium



thiosulfate) and precursor solution (5.2 mmol L⁻¹ selenium dioxide) were prepared in the 0.01 mol L⁻¹ SDS. SDS was used not only as stabilizing agent but also as diluent. Various amounts of sodium thiosulfate were added in selenium(IV) dioxide, as summarized in Fig. 1. At room temperature, the selenium solutions converted from colorless to various colors. Stirring was carried out up to 5 hours. Then, the resulting SeNPs was stored at room temperature. All detailed reactions are mentioned elsewhere.¹⁴

SeNPs by protein stabilized (second protocol) was prepared by adding ascorbic acid into protein solution, as illustrated in Fig. 1. The solutions were allowed to react for 15 minutes to ensure the high yield of selenium nanoparticles. To prevent SeNPs from aggregation, SeNPs were stored at 4 °C.

Instrumentation

Nano zetasizer based on dynamic light scattering. Hydrodynamic diameter (d_h) of the synthesized SeNPs were determined by using nano zetasizer (Malvern Instrument Nano Series ZS, Worcestershire, UK) based on dynamic light scattering (DLS). Freshly synthesized SeNPs were measured at 25 °C in triplicates for which SeNPs stabilized by SDS was diluted in 0.01 mol L⁻¹ SDS and SeNPs stabilized by protein was diluted in water as similar to the medium used in the synthesis. Size was determined based on the average intensity after checking of instrument accuracy by using 100 nm standard polystyrene (PS) nanoparticles diluted in water. The measurements of zeta potential of SeNPs were performed by using zetasizer (Malvern Instrument Nano Series ZS, Worcestershire, UK) and all samples were prepared as same as the sample preparation for

DLS measurements. The synthesized SeNPs was put in the capillary zeta cell for the zeta potential measurement.²⁶ The measurements of zeta potential was carried out in triplicates for each sample.

FIFFF systems. Two different flow field-flow fractionation systems were employed for sizing of SeNPs. The first type of FIFFF is symmetrical flow field-flow fractionation (Sy-FIFFF) (Model PN-1021-FO; Postnova Analytics, Landsberg, Germany) with the dimension of 27.7 cm long × 2.0 cm wide × 250 μm thick. The system was equipped by a rectangular shape of 250 μm spacer which was clamped between two parallel blocks. 1 kDa and 10 kDa regenerated cellulose RC; and 1 kDa and 10 kDa polyethersulfone (PES) membranes (Postnova Analytics) were used to observe the effect of membrane materials. This system was connected to two HPLC pumps (Model PN 2101, Postnova Analytics) to drive the carrier liquid which were 0.02% FL-70 + 0.02% NaN₃ and 0.02% SDS + 0.02% NaN₃. All carrier liquids were prepared in de-ionized water. Those carrier liquids were used to study the effect of carrier liquids on the retention behavior of SeNPs. Then, samples were introduced through the injector valve and detected *via* UV/VIS detector (Model SPD 20AV, Shimadzu, Japan) at 410 nm for small particles (sample A–E; sample H and sample I) and 550 nm for large particles (sample F and sample G). More details about Sy-FIFFF optimum conditions are summarized in Table S1.† The optimum conditions were selected based on the good peak quality and percentage relative recovery of carrier liquid and membrane material (see ESI Section 1 and Fig. S1–S3†).

Another FIFFF called Asy-FIFFF (AF2000, Postnova Analytics, Salt Lake City, USA) with the dimension of 33.5 cm long × 4.0 cm

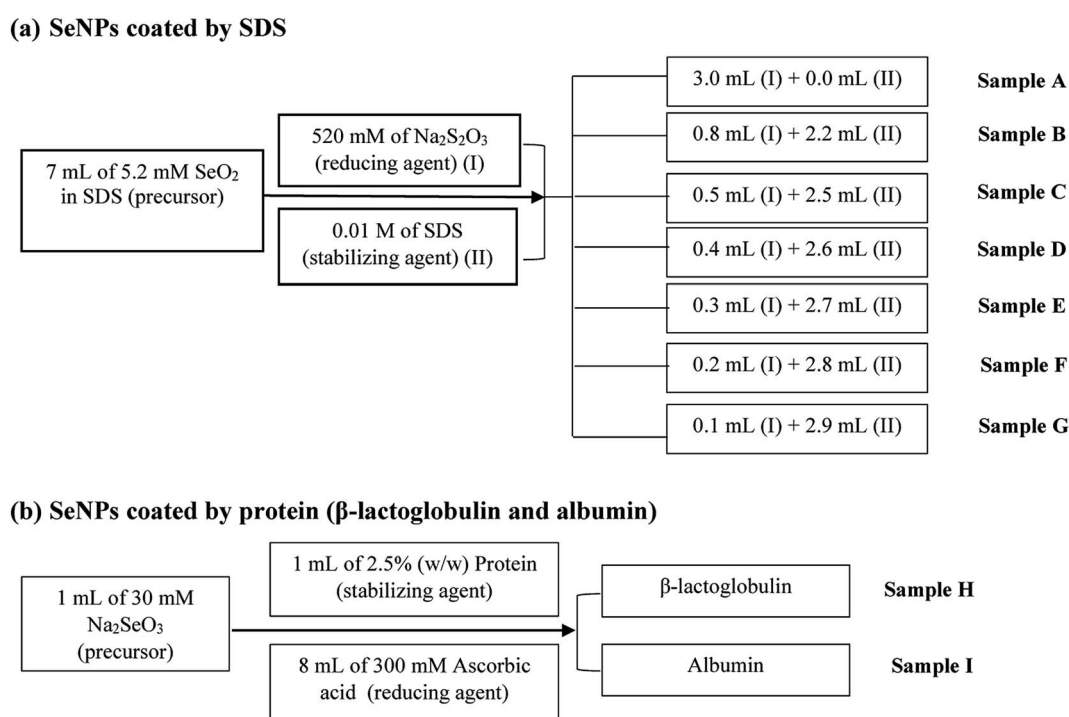


Fig. 1 Schematic diagram showing the synthesis protocol of SeNPs stabilized by (a) SDS and (b) proteins.



wide \times 250 μm thick was employed. A trapezoidal shape of 350 μm spacer with 10 kDa RC membrane (Postnova Analytics) was used. The Asy-FIFFF system used in this study was coupled with UV detector SPD-20A (Postnova Analytics). The detection wavelength at 410 nm was used for small particles (sample A–E; sample H and sample I) and 550 nm for large particles (sample F and sample G). Carrier liquid (0.02% FL-70 + 0.02% NaN_3) was delivered by two HPLC pumps (Postnova Analytics). To determine the channel thickness, 100 nm PS standard particle was introduced into the FIFFF. Then, the hydrodynamic diameters of synthesized SeNPs were determined. PS standards of 20 nm, 40 nm, and 60 nm were injected into the Asy-FIFFF for checking the sensitivity of system. The details of running condition are shown in Table S2.†

Transmission electron microscope. The morphology of all samples was analyzed by TEM (JEOL, JEM-2100 Electron Microscope, USA). 500 μL of fresh SeNPs suspension was dropped on to the copper grid and dried for a few minutes, then the copper grid was placed on the sample holder and introduced into the system. The data conversion from TEM image to the size distribution was carried out by using ImageJ launcher, version 1.4.3.67. Area of each particle in the TEM image was converted into diameter (d_{area}) following eqn (1) and then eqn (2).

$$A = \pi r^2 \quad (1)$$

$$d = 2 \times r \quad (2)$$

SP-ICP-MS. A PerkinElmer NexION 2000 ICP-MS was used in a single particle mode for size characterization of the synthesized SeNPs. The sample flow rate was determined by weighing the mass of water before and after taken up into the system by peristaltic pump for 5 minutes. In this work, the transport efficiency was determined based on the standard particle number concentration³⁷ of AuNPs 60 nm (5 mg L^{-1}) with the particle number concentration of 3.2×10^4 particles per mL. Theoretically, the transport efficiency was calculated following eqn (3):

$$f = C_{\text{STD}} \times \eta_{\text{STD}} \times Q_{\text{sample}} \quad (3)$$

where f is pulse frequency, C_{STD} is mass concentration (μg per event), η_{STD} is transport efficiency, and Q_{sample} is sample flow rate (mL min^{-1}). Following the method performed by Motellier *et al.* (2017),³⁸ the ionic standard solution of gold in the 0.1% nitric acid was used for checking the sensitivity of SP-ICP-MS to perform the size accuracy of particles. Then, three different sizes of AuNPs (20 nm, 60 nm, and 80 nm) were used for checking the system sensitivity. All AuNPs used were prepared in deionized water. Pulse intensity was converted into the size distribution by following eqn (4):

$$W = C_{\text{STD}} \times \eta_{\text{STD}} \times Q_{\text{sample}} \quad (4)$$

Pulse intensity was converted into the mass flux, which is further converted into diameter. The calibration curve was

constructed by using 1, 3, and 5 $\mu\text{g L}^{-1}$ of ionic selenium standard solution. All individual samples were diluted at: $500\times$; $50\times$; and followed by $25\times$ in deionized water before introduction into the SP-ICP-MS. The dilution processes of the samples were supposedly optimized to avoid coincidence events. The Syngistix™ nano application software was used for data evaluation. The instrumental operating conditions are listed in Table S3.†

Consecutive size analysis by SP-ICP-MS after Sy-FIFFF. Two sample mixtures of SeNPs-SDS were tested. Those include $60.5 \pm 1.4 \text{ nm}$ (sample B) and $80.8 \pm 2.0 \text{ nm}$ (sample C); and $60.8 \pm 1.0 \text{ nm}$ (sample B) and $130.1 \pm 2.3 \text{ nm}$ (sample E). DLS was used to confirm those sizes of individual SeNPs (sample B; sample C; and sample E). The mixtures of sample B and sample C; and sample C and sample E were diluted as same as the individual SeNPs as explained earlier and were sonicated prior to SP-ICP-MS analysis. Then, both mixtures were introduced into Sy-FIFFF under optimum conditions (see Table S1†). The fractions of both mixtures were collected from the left side (fractions 1 and 3) and the right side (fractions 2 and 4) of the nanoparticle peak (see more details in the result and discussion part). 50 μL of the fraction collected from Sy-FIFFF was introduced into the ICP-MS in single particle mode with a $100\times$ dilution.

Data evaluation of hydrodynamic diameter and core diameter from FIFFF and SP-ICP-MS

In Sy-FIFFF and Asy-FIFFF system, peak area of each sample was analyzed by using OriginPro 8.6 software (OriginLab Corporation, MA, USA). The retention time (t_r) of peak elution for all samples was obtained to determine the hydrodynamic diameter (d_h), then was calculated using eqn (5) below:

$$d_h = \frac{2V_0kT}{\pi\eta V_C w^2 t_r} \quad (5)$$

where V_0 = volumetric channel flow rate, k = Boltzmann's constant, T = temperature, V_C = volumetric cross flow rate, η = viscosity of carrier liquid and w = channel thickness. V_0 and V_C values were chosen from the optimization considering from the well-defined elution peak, good resolution, and short running time. Channel thickness determinations were carried out from the retention time of 0.1% solid of 100 nm standard PS nanoparticle based on a peak breakthrough technique proposed by Giddings *et al.* (1993).³⁹ For SP-ICP-MS, data evaluation was performed by plotting all the histograms from the Syngistix™ nano application software which is pulse intensity (counts per dwell) vs. frequency reading.

Statistical analysis (paired *t*-test method) for comparison of the results by each technique

In order to compare results obtained by two different techniques, paired *t*-test analysis was used for checking if there is any significant difference between two techniques. The *t* calculation (t_{cal}) was compared with the *t* critical or theory (from *t*-table). The t_{cal} was calculated from the data of experiment as shown in eqn (6):



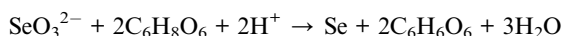
$$t_{\text{cal}} = \frac{\bar{d} - \mu_d}{S_d} \times n^{1/2} \quad (6)$$

where \bar{d} is the sample mean difference, S_d is the standard deviation of differences, μ_d is the true mean difference, and n is number of paired sample data of the test. In our experiment, nine samples were tested resulting in 9 paired data (n). If the t_{cal} is smaller than t_{crit} , there is no significant difference between the two methods.⁴⁰

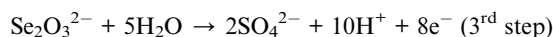
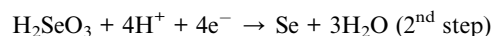
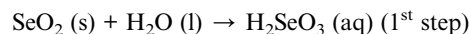
Result and discussion

DLS for hydrodynamic diameter determination of SeNPs

SeNPs with different types of stabilizing agents were successfully synthesized (Fig. 2a). These include sodium dodecyl sulfate (SDS) stabilized SeNPs (sample A–G) (Fig. 2a from A to G) and proteins stabilized SeNPs including β -lactoglobulin (Fig. 2a, H) and albumin (Fig. 2a, I). These coating agents were used to prevent SeNPs from aggregation.⁴¹ SeNPs were prepared by ascorbic acid ($\text{C}_6\text{H}_8\text{O}_6$) reduction of selenite (SeO_3^{2-}) to selenium with zero oxidation state.⁴



For the synthesis of SeNPs coated by SDS, various amounts of sodium thiosulfate as reducing agent were used. The formation of SeNPs occurred when sodium thiosulfate as reducing agents was added into the solution containing selenium dioxide as precursor and SDS as stabilizing agents. According to Lin *et al.* (2002),¹⁴ selenite in the second step was rapidly reduced to selenium with zero valent in the presence of sodium thiosulfate.



As the amount of sodium thiosulfate decreased, the orange color changed from light to dark (from A to F) except SeNPs with the lowest amount of thiosulfate (G). The lowest amount of thiosulfate resulted in the largest particle size and the highest possibility to undergo aggregation as it happened in the SeNPs (G). According to Lin *et al.* (2002),¹⁴ at higher concentrations of thiosulfate reducing agent the nucleation process was faster than the growth of the particle, resulting in smaller particles. Considering the absorption spectra of SeNPs in Fig. S4,†

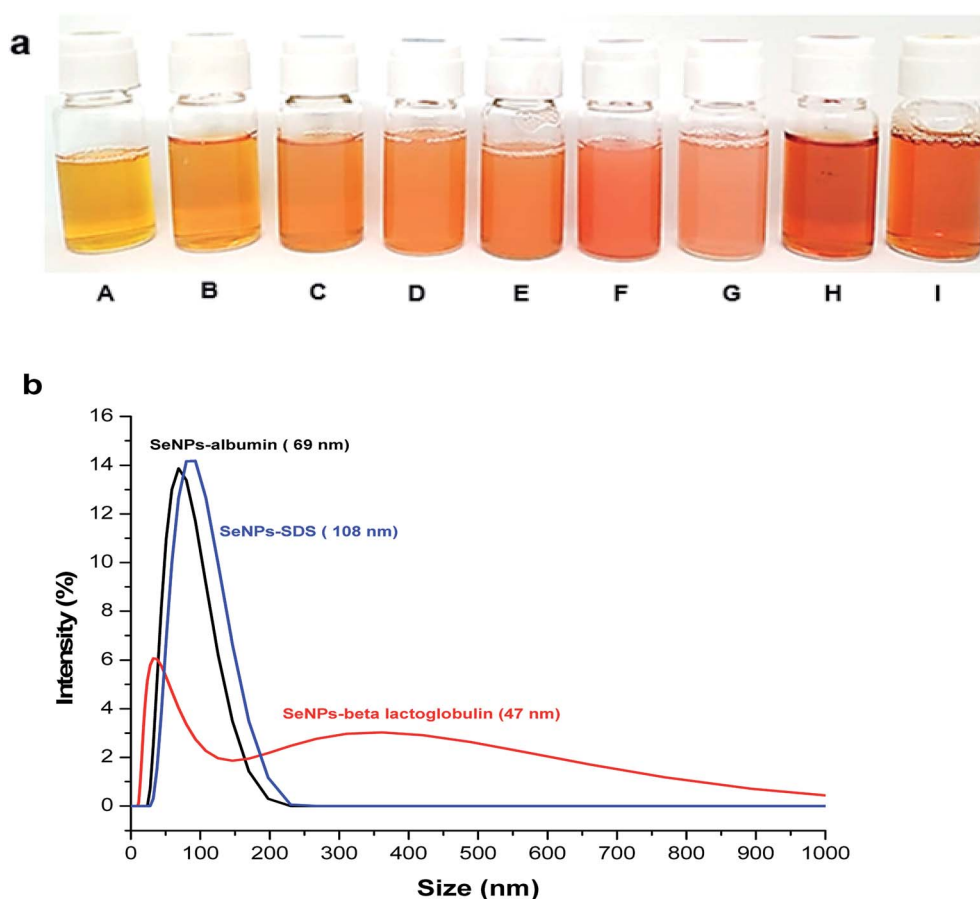


Fig. 2 (a) Image of SeNPs synthesized by using SDS as stabilizing agent with various amounts of sodium thiosulfate (sample A–G) and protein as stabilizing agents (sample H and I) (b) size distribution observed from DLS for SeNPs stabilized by SDS (sample D); SeNPs stabilized by β -lactoglobulin; and SeNPs stabilized by albumin ($n = 3$).



decreasing the amounts of thiosulfate resulted in the red-shift of SeNPs absorption, indicating the larger particle size.

Different types of stabilizing agents provided different zeta potential values. The zeta potential values of SeNPs coated by proteins are positive (23.3 ± 0.5 mV) for SeNPs stabilized by β -lactoglobulin (sample H) and (35.8 ± 1.5 mV) for SeNPs stabilized by albumin (sample I) and that of SDS is negative (see Table 1) which is due to the different functional groups of each stabilizer. The zeta potential values of the sample should be known before FIFFF experiment to ensure a good selection of carrier liquid and membrane materials.³⁸

The information of d_h is also helpful prior to FIFFF experiment for monitoring of the behavior of NPs.³⁸ Hence, DLS was used for hydrodynamic diameter determination of SeNPs. Table 1 summarizes the hydrodynamic diameters of all SeNPs by using DLS. Considering the hydrodynamic diameters of SeNPs stabilized by proteins and SDS with the smallest size, SeNPs coated by proteins (sample H and sample I) showed larger size than SeNPs coated by SDS (sample A). This is due to the differences between molecular weight of each stabilizing agent which are 288.4 g mol^{-1} (~ 0.28 kDa) for SDS, 18.4 kDa for β -lactoglobulin and 66.5 kDa for albumin. However, the size of the NPs does not depend only on the molecular weight of coating agent but also the concentration of stabilizing agent and reducing agent.¹⁴

In order to better evaluate the stability of SeNPs coated by protein before introduction into FIFFF, the size distribution of those SeNPs was monitored. Focusing on the results from DLS for size distribution of SeNPs, SeNPs coated by β -lactoglobulin does not show a monomodal peak such as for those SeNPs coated by albumin and SDS, as illustrated in Fig. 2b. The hydrodynamic diameter at the larger value around 350 nm was observed for SeNPs stabilized by β -lactoglobulin and contributed more than 50% of intensity. The zeta potential value of SeNPs coated by β -lactoglobulin is less than 30 mV, implying that agglomeration from SeNPs coated by β -lactoglobulin is more likely as compared to other NPs mentioned. The bimodal peak of β -lactoglobulin observed from DLS is due to that at

pH = 2, β -lactoglobulin is presented as dimers and other larger aggregates. Above pH 7, the dimers of β -lactoglobulin can be dissociated into monomers.⁴² Adjusting the pH above 7, the zeta potential of SeNPs coated by β -lactoglobulin changed from positive to negative charge.^{4,15} In this case, the stabilizing agents or ligands used for NPs stabilization may affect the diffusion behavior of nanoparticles such as the shifting of peak intensity towards the larger value.^{38,43} It should be noted that challenge still remains for analysis by DLS, particularly for nanoparticles coated by proteins due to their possibility for aggregation as it was also previously mentioned in another literature for β -lactoglobulin characterization.⁴⁴ To reduce the aggregate formation of SeNPs coated by β -lactoglobulin by using DLS, adjusting the pH and setting the right temperature is highly recommended.

FIFFF for hydrodynamic diameter determination of SeNPs

For accurate hydrodynamic diameter estimation of SeNPs by Sy-FIFFF, the optimization of Sy-FIFFF system is necessary. In order to minimize particle membrane interaction, types of membrane materials and carrier liquids were carefully chosen to obtain good peak quality. All details of the condition optimization were performed and the optimum condition was selected as explained in ESI (see Sections 1–3 in ESI; Fig. S1–S3[†]). All SeNPs samples were introduced into the Sy-FIFFF by using FL-70 and sodium azide as carrier liquid (approximately pH 8–9) and 10 kDa RC as membrane material. In order to evaluate the efficiency of Sy-FIFFF system, we also introduced all SeNPs-SDS samples into Asy-FIFFF system under the same conditions. The fractograms of SeNPs obtained from two different systems of FIFFF are illustrated in Fig. 3.

The fractograms of SDS stabilized SeNPs from Sy-FIFFF are shown in Fig. 3a, and those of protein stabilized are illustrated in Fig. 3b. The fractograms of SeNPs obtained from Asy-FIFFF are displayed in Fig. 3c and d. It is interesting to have a closer look at the elution peak of SeNPs stabilized by β -lactoglobulin. As mentioned earlier by using DLS method, bimodal peaks were

Table 1 The hydrodynamic diameter (at peak) of synthesized SeNPs by using Sy-FIFFF and Asy-FIFFF compared with DLS ($n = 3$) and the core diameter (at peak) of synthesized SeNPs by using TEM and SP-ICP-MS^a

SeNPs synthesized	Hydrodynamic diameter			Core diameter		Zeta potential (mV)
	Sy-FIFFF	Asy-FIFFF	DLS	TEM	SP-ICP-MS	
SeNPs coated by SDS with various amounts of thiosulfate						
Sample A	33.6 ± 0.5	28.1 ± 0.3	36.1 ± 1.0	26.9 ± 3.5 ($N = 172$)	27.6 ± 1.8	-67.8 ± 1.5
Sample B	52.3 ± 1.2	61.7 ± 1.1	58.7 ± 0.2	32.0 ± 3.1 ($N = 212$)	35.1 ± 2.2	-60.2 ± 2.7
Sample C	82.7 ± 1.8	78.3 ± 0.9	88.3 ± 0.2	70.1 ± 3.2 ($N = 157$)	67.9 ± 1.4	-31.5 ± 1.6
Sample D	100.9 ± 1.8	107.5 ± 0.6	108.9 ± 0.8	93.0 ± 1.6 ($N = 78$)	90.3 ± 0.2	-31.2 ± 1.1
Sample E	127.7 ± 1.2	123.7 ± 0.3	134.9 ± 0.9	107.9 ± 2.6 ($N = 51$)	110.9 ± 3.3	-32.5 ± 1.1
Sample F	164.6 ± 2.4	156.1 ± 0.2	175.7 ± 0.7	149.4 ± 3.1 ($N = 22$)	143.9 ± 1.9	-37.2 ± 3.6
Sample G	188.4 ± 1.5	189.1 ± 0.4	210.6 ± 1.9	180.9 ± 2.4 ($N = 40$)	180.8 ± 3.2	-37.5 ± 2.4
SeNPs coated by proteins						
Sample H	44.8 ± 1.4	50.2 ± 0.6	47.3 ± 1.0	31.8 ± 2.5 ($N = 101$)	33.0 ± 1.4	23.3 ± 0.5
Sample I	75.2 ± 2.3	56.6 ± 0.9	68.7 ± 0.4	63.6 ± 4.6 ($N = 105$)	61.8 ± 2.2	35.8 ± 1.5

^a Note: N is the total numbers of particles detected.



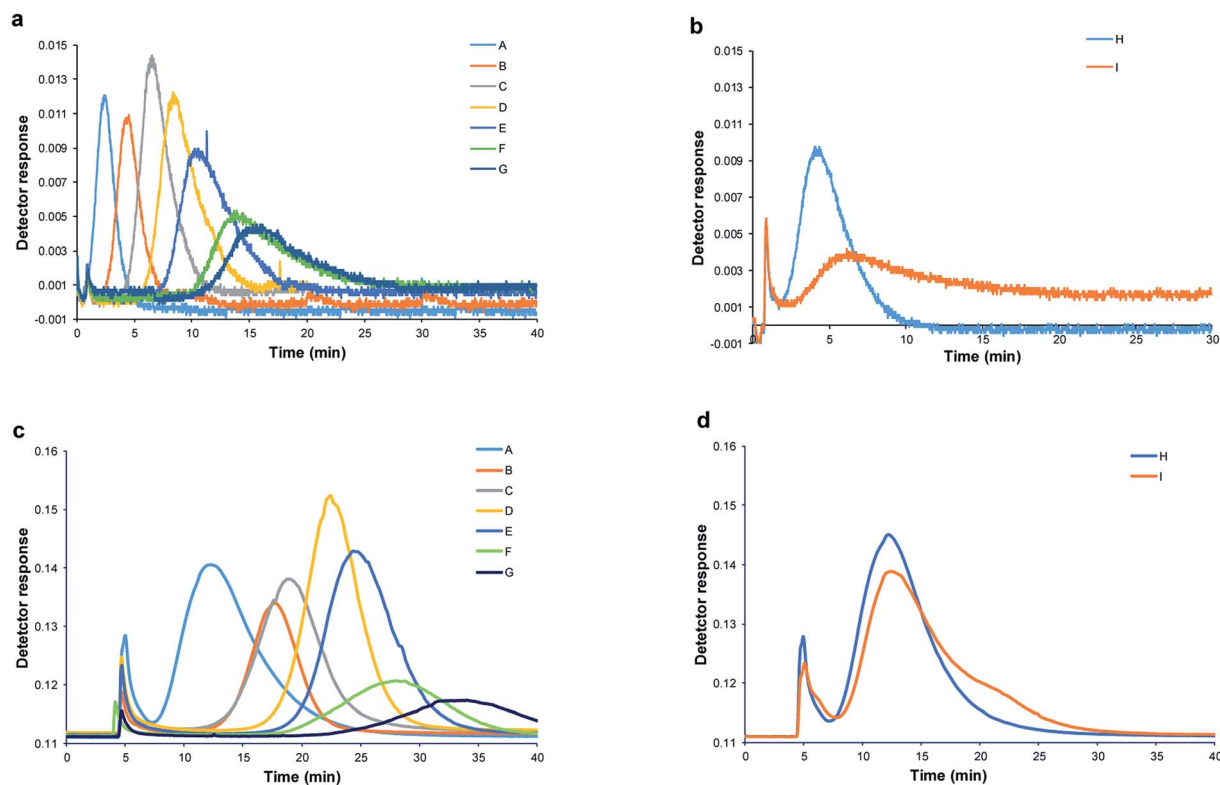


Fig. 3 Overlaid fractograms of SeNPs synthesized with various types of stabilizing agents from Sy-FIFFF (a and b) and Asy-FIFFF (c and d). (a) and (c) SeNPs stabilized by SDS with various amounts (mL) of thiosulfate as reducing agent [sample A–G represents the shifting retention time of SeNPs synthesized by reduced the amount of thiosulfate]. (b) and (d) SeNPs stabilized by protein [β -lactoglobulin (sample H) and albumin (sample I)] with the use of 0.02% FL-70 and 0.02% NaN_3 and 10 kDa RC membrane ($n = 3$).

observed. With Sy-FIFFF and Asy-FIFFF, however, only one peak was observed for SeNPs coated by β -lactoglobulin. By converting the fractograms into particle size distribution shown in Fig. S5(a and b),[†] the size detectable was in the range of 50–100 nm under this operating condition.

The hydrodynamic diameter (d_h) estimation from Sy-FIFFF and Asy-FIFFF was calculated based on the retention time observed from the fractogram and converted into the diameter. Following method by Giddings *et al.* and Dou *et al.* (2015),^{39,45} we calculated the hydrodynamic diameter of SeNPs based on the channel thickness measurements from the known diameter of standard PS particle which is a suitable procedure for all types of FIFFF. But this method is also challenging due to the uncertainties from membrane compression. The type of standard such as PS nanoparticles are suitable to use for channel thickness determination. PS standard particles are recommended to use because it is inert avoiding particle membrane interaction and it is rather temperature independence.⁴⁵ According to the FIFFF retention theory (see eqn (5)) following the method by Saenmuangchin and Siripinyanond in 2018 for AuNPs²⁶ by using Sy-FIFFF, the size calculation of NPs can be directly determined based on the retention time if the value of the channel thickness is known. The channel thickness (w) is important to be measured by using standard nanoparticle at the same condition as for sample measurement.⁴⁵ Hence, 100 nm standard polystyrene particle was selected for channel thickness determination.

According to the results from statistical analysis (see eqn (6)), the hydrodynamic diameters obtained from Sy-FIFFF and Asy-FIFFF were not significantly different as t_{cal} (0.72) was less than t_{crit} (2.31) as shown in Table 1. It shows that the different channel construction of FIFFF does not affect on the size measurements of NPs but affects on the peak widths. As Asy-FIFFF channel has only one bottom frit and lower surface area than that of Sy-FIFFF due to its trapezoidal shape, it can minimize the particle membrane interaction and decrease peak widths.⁴⁶ As shown in Fig. 3b and d for SeNPs stabilized by protein, the peak elutions of SeNPs (sample I) are different. For sample I, SeNPs coated by albumin showed two peaks due to the aggregation of particles (Fig. 3d). By using this technique, the determination of the hydrodynamic diameter based on retention time is very dependent on the particles type and coating. In summary, under optimum conditions Sy-FIFFF has ability to estimate the hydrodynamic diameters of SeNPs as the results were similarly to those from Asy-FIFFF.

Concerning the size comparison of SeNPs from both techniques with DLS by using paired t -test method, the t calculation (t_{cal}) from the experiment is 3.29 for Asy-FIFFF and 3.08 for Sy-FIFFF, whereas the t critical (t_{crit}) is 2.31 (at $P = 0.05$). Therefore, the hydrodynamic diameters from both techniques (FIFFF and DLS) are significantly different as t_{cal} was higher than t_{crit} . The difference of both FIFFF techniques from the DLS was mainly due to the possibility of DLS to provide larger particle size than



FLFFF. According to the light diffusion theory, a particle having an “ n ” times larger diameter diffuses “ n^2 ” times more light. The larger particles show more scattering than the smaller particles, which can affect to the average mean diameter of NPs. As observed in the earlier study by other investigators for SiO₂ NPs,⁴⁷ DLS measured the z-averaged diameter whereas FLFFF with UV detection offered the information on the volume-averaged diameter, causing the DLS to show high sensitivity for larger size ranges and low sensitivity for smaller size ranges. Another plausible explanation for larger size observed from DLS especially for SeNPs coated by SDS was the extension of the electrical double layer of SDS micelles⁴⁸ around SeNPs, and thereby reducing the diffusion speed of particles which resulted in the larger experimentally observed hydrodynamic diameter.

TEM for core size determination of SeNPs

The information on core diameter is obtained by using TEM. With TEM, size distribution of NPs could be analyzed by using NIH-image/ImageJ program. This program can be used for counting the number of representative particles in the image.

The interpretation of results by using this technique was reported by Woehrle *et al.*¹⁹

According to the TEM image (see Fig. 4), the size of SeNPs increased as a result of decreasing the amount of thiosulfate. Particle size distributions from TEM are shown in the Fig. 5A (sample A–C) and ESI (see Fig. S6† (sample D–I)). The particle size distribution as shown in those figures was obtained by data conversion from the area to diameter (see eqn (1) and (2)) of each image of SeNPs (A–I), and was converted to the particle size distribution by using OriginPro 8.6 software. It is clearly seen from those figures that the population of nanoparticles decreases as the size increases. For the large particle, agglomeration is a main cause for decreasing numbers of particles, as shown in Fig. S6(f)† for sample F. For SeNPs coated by protein (sample H and I), the agglomeration of particles occurred (Fig. 3(H) and (I)). As shown in Fig. S6(h) and (i),† the particle size distribution detected in this sample was until around 180 nm which indicated the aggregation of samples. By using only this technique without confirmation by another technique for all SeNPs samples, the largest particle detected as shown in the particle size distribution

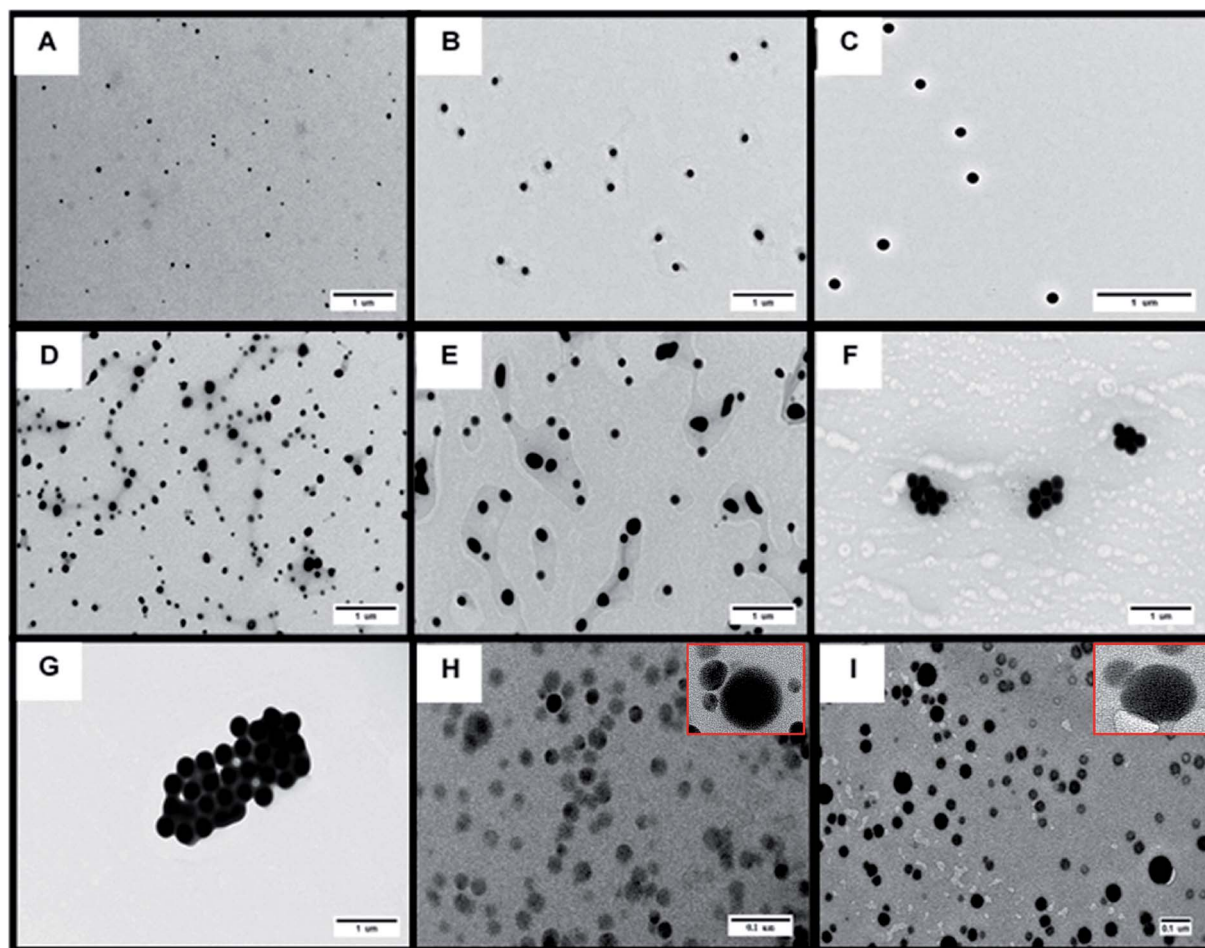
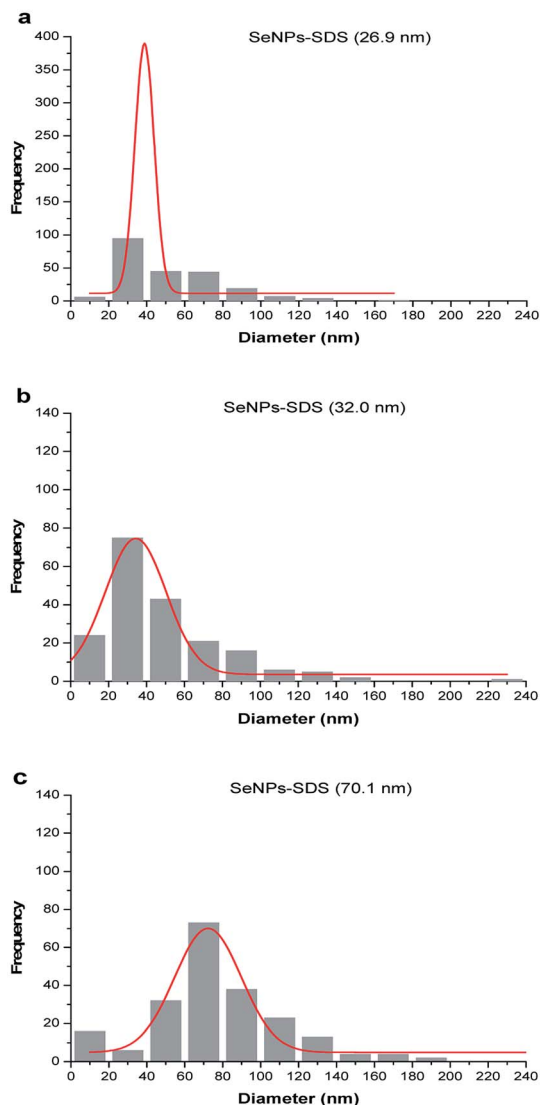


Fig. 4 TEM images of SeNPs. Image (A)–(G) represent the SeNPs stabilized by SDS with decreasing amount of thiosulfate with the mean diameter of 26.9 ± 2.5 (sample A), 32.0 ± 2.1 (sample B), 68.4 ± 0.2 (sample C), 93.0 ± 2.6 (sample D), 114.5 ± 1.6 (sample E), 147.2 ± 3.1 (sample F), and 179.9 ± 2.4 nm (sample G). Image (H) and (I) represent the sample by using protein as stabilizing agent with the mean diameter of 32.3 ± 1.6 (sample H) and 62.5 ± 2.5 (sample I).



(A) TEM



(B) SP-ICP-MS

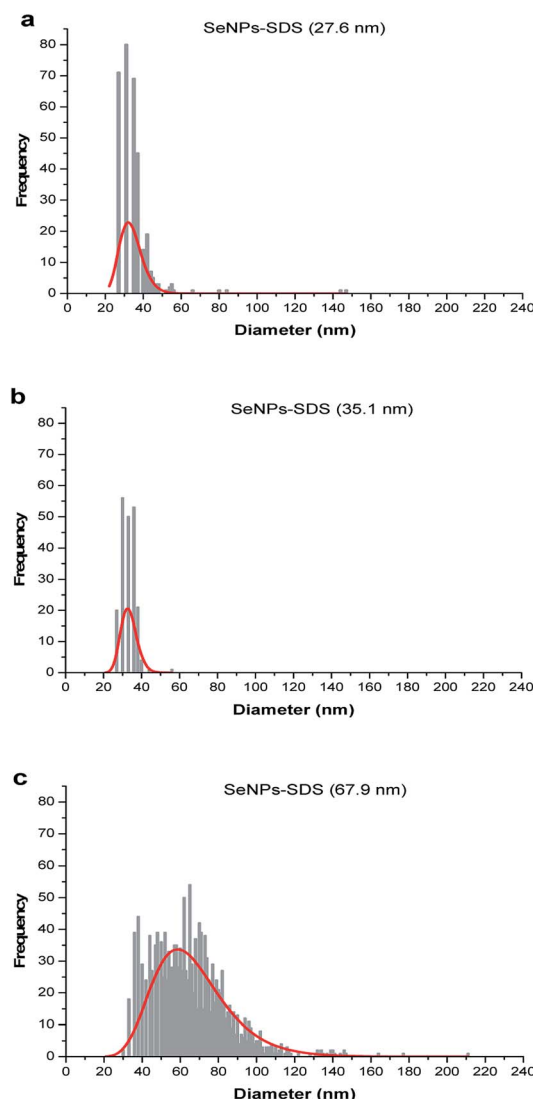


Fig. 5 Particle size distribution of SeNPs-SDS (sample A; sample B; sample C) from (A) TEM ($n = 5$) and (B) SP-ICP-MS ($n = 2$). The number of particle selected for area-equivalent diameter measurements from TEM was 20 to over 100 particles in 5 TEM image for one sample.

(Fig. S6†) can not be distinguished due to the agglomeration of SeNPs presented in the suspension or during the preparation on the TEM grid, as the sample was prepared by drying for few minutes before introducing into the TEM as also mentioned by Loeschner *et al.* (2013).³⁶ But, this technique allowed the detection of particle size < 10 nm in the sample as shown in Fig. 5A(a–c) for small particles of SeNPs coated by SDS. In contrast, the actual number concentration of nanoparticle population cannot be provided by TEM (quantification analysis). Therefore, SP-ICP-MS is used for providing specific information on number of particles and their mass concentration.

SP-ICP-MS for core size determination of SeNPs

The differences between SP-ICP-MS and TEM is the sample concentration needed for the analysis. SP-ICP-MS allows the

concentration in ppb–ppt level to be analyzed, meaning less sample consumption. In contrast, TEM can detect the smallest particle size down to few nanometers (see Fig. 5). The slight differences in the particle size for SP-ICP-MS are shown in the Fig. 5B of (a–c) sample (represents sample A–C), the particle size below 10 nm was not detected as compared with TEM technique (see Fig. 5A). The smallest particle size detectable by SP-ICP-MS as shown for sample A–C (Fig. 5B(a–c)) is between 20 to 25 nm. Based on this result, TEM can be used for sample with size lower than 10 nm but not for sample with low concentration and small amount of samples. In this case, SP-ICP-MS is a well-suited method to perform the particle size of NPs at low number concentration of sample, but high sample dilution is required which could cause particle instability and more attention should be paid. Although the number of classes in SP-ICP-MS size distribution was not the same for all samples, we diluted



all samples to contain between 1.0 to 4.0×10^4 particles per mL to ensure reliable particle size analysis.

Table 1 shows the particle size of SeNPs by SP-ICP-MS and TEM. The results obtained from SP-ICP-MS are in good agreement with those from TEM which ranged from 30 nm to 180 nm. There was no significant difference between the two techniques, as checked by paired *t*-test method. The t_{cal} (0.5) is less than t_{crit} (2.31) at $P = 0.05$. Eventhough TEM and SP-ICP-MS techniques provide similar results, the apparent difference is the number population of nanoparticles present when plotted as size distribution which is shown in Fig. S6 and S7.†

In the following experiment, the mixed size of SeNPs-SDS was also introduced into SP-ICP-MS to test the feasibility of system for more complex sample such as synthesized NPs. The mixed size sample B and sample C; and sample B and sample E were observed as shown in Fig. 6a and b. It is not clear how

many size presents in the suspension as shown by the red arrow. For further investigation, SP-ICP-MS size analysis was carried out for the fractions collected from Sy-FIFFF since the system showed the best performance for SeNPs separation based on their different sizes under optimum condition of membrane material and carrier liquid.

Comparison of the diameter information obtained from various techniques

As described earlier, the difference in particle size values obtained from various techniques is due to the different principles of each technique. According to the data in Table 1, DLS and FIFFF yielded larger sizes than those from TEM and SP-ICP-MS. In theory, DLS and FIFFF measure the hydrodynamic diameter, which refers not only to the core diameter of particle but also

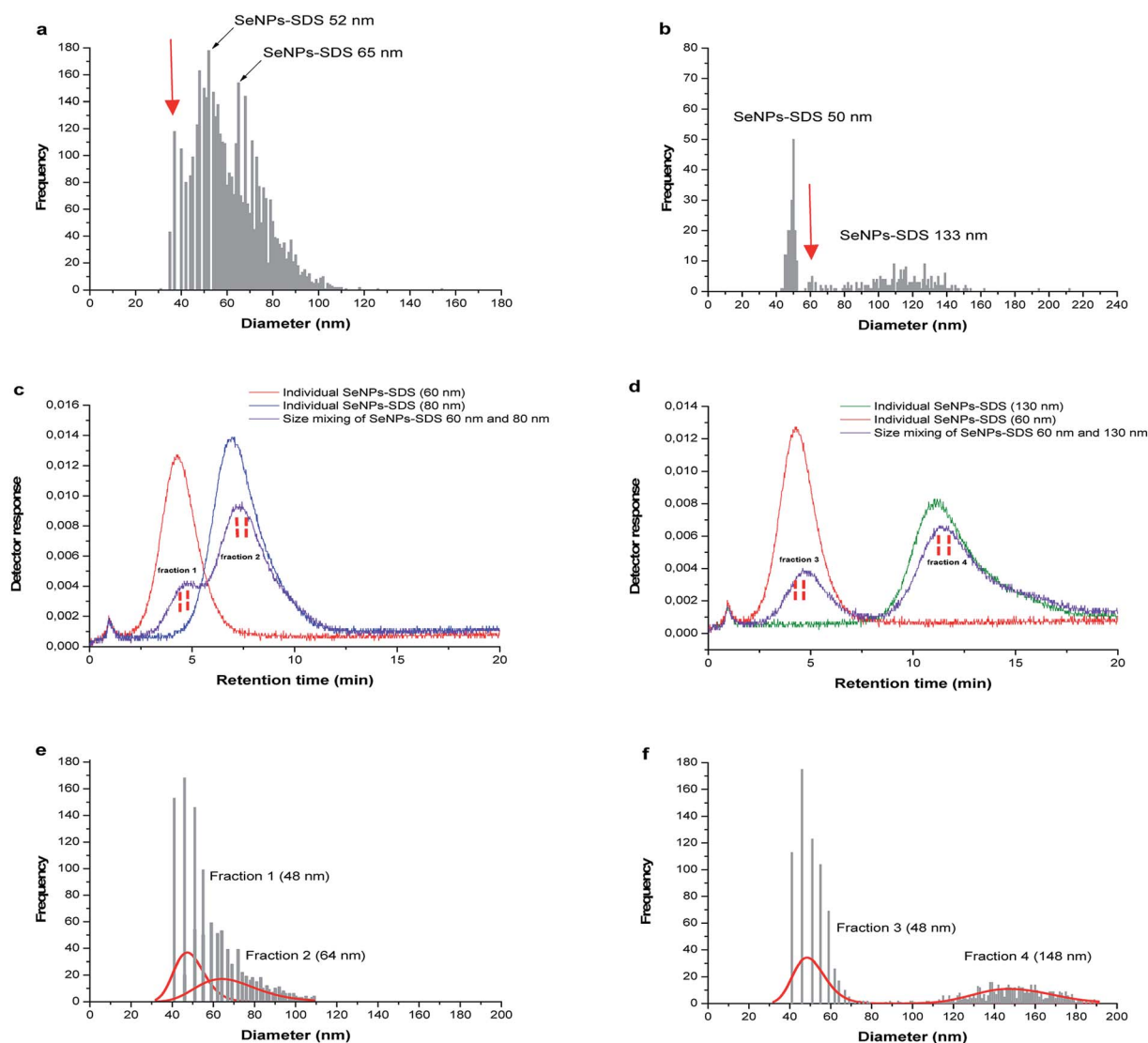


Fig. 6 (a) and (b) Direct measurement of size mixture of 60 nm (sample B) and 80 nm (sample C); and 60 nm (sample B) and 130 nm (sample E) (DLS analysis) into SP-ICP-MS. (c) and (d) Sy-FIFFF fractograms of individual SeNPs of 60 nm, 80 nm and 130 nm and fractions of SeNPs 60 nm and 80 nm; and 60 nm and 130 nm collected for size analysis by SP-ICPMS. (e) and (f) Number size distribution of SeNPs fractions collected from Sy-FIFFF ($n = 3$).



Table 2 The size comparison of using SP-ICP-MS with or without coupling with FIFFF for size mixture^a

Mixtures of SeNPs	Mean size of SeNPs in mixtures (nm)		
	Obtained from FIFFF ($n = 3$)	Obtained from SP-ICP-MS ($n = 2$)	Obtained from SP-ICP-MS after fraction collection from FIFFF ($n = 2$)
Sample B + sample C (1 : 2 volume ratio)	56.7 ± 1.2 ^a /79.1 ± 1.8 ^b	52.4 ± 4.8/64.6 ± 3.2	48.1 ± 1.0 ^a /64.0 ± 2.9 ^b
Sample C + sample E (1 : 2 volume ratio)	58.2 ± 1.7 ^c /130.2 ± 1.2 ^d	52.4 ± 4.8/133.9 ± 2.4	48.7 ± 1.1 ^c /148.5 ± 1.9 ^d

^a a = fraction 1; b = fraction 2; c = fraction 3; d = fraction 4.

everything around the particle core including stabilizing agents. DLS detection is based on the translational diffusion of particles which depends not only on the size of the particle but also the surface structure including type of ions in the medium. Changes in the surroundings of the particle affect on the diffusion speed of NPs, and thereby changing in the hydrodynamic diameter.⁴⁹ According to the results in Table 1, different medium and surface structure of stabilizing agents between SeNPs coated by protein (dissolved in water) and SeNPs coated by SDS (dissolved in SDS) affected on the stability of NPs which was shown by the zeta potential values. All SeNPs coated by SDS exhibited monomodal peak but not with SeNPs coated by protein (β -lactoglobulin, Fig. 2b) which had the zeta potential lower than ± 30 mV.

Considering the results from TEM and SP-ICP-MS, both techniques gave similar results according to the paired *t*-test analysis. From Table 1, TEM showed the largest values of standard deviation as compared to the other techniques. This might be due to the aggregation of NPs during the sample preparation process as the samples need to be dried under ultrahigh vacuum before TEM analysis. As shown in Fig. 4 for β -lactoglobulin and albumin stabilized SeNPs, the overlap of larger particle with smaller particle was observed. Without careful adjustment of the TEM image contrast in ImageJ software, the apparent reading is not assigned to a single particle. As the agglomeration was observed as shown in Fig. 4, the size distribution in Fig. S6† broadened and shifted to larger size, which was mentioned by Zook *et al.* in 2011 for AuNPs agglomeration.⁵⁰ The ability of TEM to observe the agglomeration of NPs is considered as an advantage of TEM over SP-ICP-MS. Size distributions obtained by SP-ICP-MS were shown to be narrow with the size limit detection of approximately 20 nm (Fig. 5 and S6†). However, the average size values obtained by SP-ICP-MS are in acceptable agreement with TEM analysis.

Consecutive size analysis by SP-ICP-MS after Sy-FIFFF for SeNPs mixtures

As mentioned in the previous part, two mixtures of different size SeNPs stabilized by SDS (sample B) and (sample C); ((sample B) and (sample E)) were introduced into Sy-FIFFF. Then, the fractions were collected from Sy-FIFFF with different interval times for further size observation by SP-ICP-MS. Fractions were

collected from Sy-FIFFF in the certain area of elution peak as shown in the dashed line in Fig. 6c (sample B and sample C) and Fig. 6d (sample C and sample E). The retention times observed for the particles in the mixture were slightly different from the retention times when individual size was introduced into the system. This is due to the different stability of SeNPs as individual and in the mixture.

All the fractions collected from Sy-FIFFF were introduced into SP-ICP-MS. Fig. 6e and f shows the size distributions by SP-ICP-MS of fractions collected from Sy-FIFFF. Fraction 1, 2, and 4, which corresponds to the individual particle of sample B, sample C, and sample E, shows a different size value from individual particle (see Fig. S8 in ESI†), especially fraction 4 (see Table 2). This might be due to the fact that larger particles are easier to agglomerate due to the high dilution by using SP-ICP-MS as reported by Loeschner *et al.*³⁶ for AgNPs fraction. The amount of particles was reduced after fractionation by FIFFF which is correlated with the number pulse counts of particle. This might be due to some particle loss during fractionation in Sy-FIFFF.

Conclusion

Different techniques were exploited to determine the hydrodynamic diameter and core diameter of synthesized SeNPs. For implementing FIFFF with the goal to provide hydrodynamic size information of nanoparticles, the selection of carrier liquid and membrane are fundamental to obtain accurate particle size results. For complex samples, the different sizes obtained by FIFFF and DLS depend on the properties of stabilizing agents and the possibility of particle agglomeration that affect to the reliability of size information. TEM yields a good visualization of sample, but high particle concentration is required. Furthermore, SP-ICP-MS is a well-suited method to provide number-based size distribution with the use of very low concentration and short time analysis. For the analysis of mixtures containing more than one size, fractionation by FIFFF was carried out prior to size analysis by SP-ICP-MS. This approach is highly recommended for samples with mixed size. Finally, it is clearly presented that no single technique provides high size accuracy and high reliability of sizing analysis for complex samples. SP-ICP-MS can not be applied in a stand-alone manner when the suspension composed more than one



size. The capability of the combined technique of FIFFF and SP-ICP-MS should be further explored for characterization of NPs in the complex sample. For future work, the developed method can be applied to detect NPs in complex matrices such as in the environment. Attention will be paid on careful optimization of sample dilutions to provide accurate size results.

Conflicts of interest

There are no conflicts to declare.

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