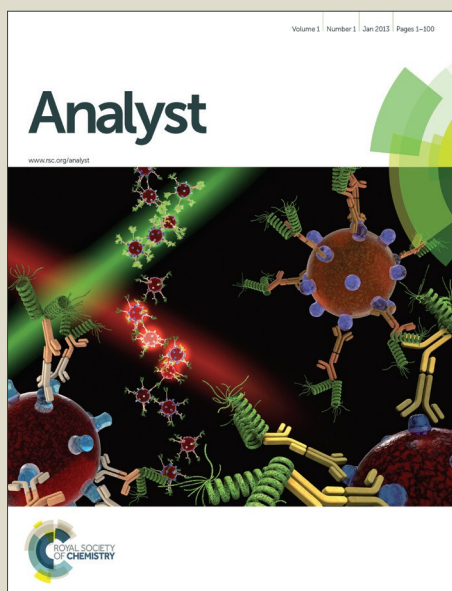


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Application of Surface Enhanced Raman Scattering to the Solution Based Detection of a Popular Legal High, 5,6-methylenedioxy-2-aminoindane (MDAI)

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Abstract

The ever increasing numbers and users of designer drugs means that analytical techniques have to evolve constantly to facilitate their identification and detection. We report that surface enhanced Raman scattering (SERS) offers a relatively fast and inexpensive method for the detection of MDAI at low concentrations. Careful optimisation of the silver sol, and salt concentrations was undertaken to ensure the SERS analysis was both reproducible and sensitive. The optimised system demonstrated acceptable peak variations of less than 15% RSD and resulted in a detection limit of just 8 ppm (5.4×10^{-5} M).

Introduction

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6 Recently there has been raised concern over the increased recreational usage of legal
7 highs.^{1,2,3} These synthetic derivatives of banned substances such as MDMA and
8 amphetamines have flooded the drugs market, often the derivatives provide a cheaper
9 alternative to illegal substances. Legal highs have the potential to cause major health risks,
10 due to little knowledge of the chemicals they contain and lack of information about human
11 consumption. Furthermore, whilst short-term side effects are easily documented little is
12 known about the effects of their long-term usage. Accessibility of the drugs over the internet
13 and in so called 'headshops', also makes them an attractive option for users, although sale of
14 the drugs is normally considered illegal under the medicines legislation act.²² This is why
15 many sellers of the 'highs' often advertise them as bath salts or plant food (not for human
16 consumption). In the UK alone it has been reported that 150 new legal highs were in
17 circulation in the three years between 2010 and 2013.²⁴ Popular 'highs' being sold over the
18 internet include 5-IAI (5-iodo-2-aminoindane),⁴ Benzofury (6-(2-aminopropyl)benzofuran)⁵
19 and MDAI (5,6-methylenedioxy-2-aminoindane)^{6,7} to name but a few. However, it is the
20 latter of these drugs, MDAI (Figure 1) which is of concern to this work. The widespread
21 availability and recreational use of MDAI is thought to have been created via the banning of
22 mephedrone, a cathinone derivative,^{8,9} which had caught the attention of the UK media
23 towards the end of 2009.^{10,11} Due to the growth in the number of users and documented ill-
24 effects, the substance was consequently categorised as a class B drug along with other
25 cathinone derivatives in April 2010.¹² Nichols at Purdue University first synthesised MDAI in
26 1990,¹³ the structural basis is similar to that of 3,4-methylenedioxy-*N*-methylamphetamine
27 (MDMA) with the only difference between the two being that the methylpropan-2-amine
28 moiety of MDMA is replaced with a 2-aminoindane group. MDAI has been shown to have a
29 indistinguishable pharmacology to MDMA (Figure 1) whose primary mechanism is to act as
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3 a selective serotonin releasing agent.^{14,15} It is therefore evident that like most amphetamines
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5 MDAI is taken for its entactogenic effects, which include increased levels of intimacy,
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7 consciousness and euphoria, but these effects often contradict online blogs written by the
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9 drugs users who demonstrate mixed reviews about the drug's effect.⁸ Most seriously
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11 however, it is believed that the first death caused by an MDAI overdose was recorded in the
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13 Isle of Man on the 15th of April 2011.¹⁶ The development and optimisation of new and
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15 existing laboratory analytical methods is essential in order to remain up-to-date with the rapid
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17 changes in drugs culture.¹⁷ However, little analytical work to detect and establish the limit of
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19 detection of MDAI has been carried out. There are very few studies in the literature that
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21 report analytical methods for the detection of MDAI. One article reports on how
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23 microcrystalline analysis of MDAI is capable of establishing a LOD of 0.2g/L,¹⁸ whilst
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25 another described the use of GC-MS, NMR and FT-IR for the characterisation of MDAI and
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27 other structural analogues.¹⁹ It is therefore evident that much work is needed in the detection
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29 of the synthetic legal high. Raman spectroscopy is an attractive option for the drugs analysis.
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31 However, whilst it generates a unique spectrum of interrogated analytes, the inherent lack of
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33 sensitivity and fluorescence based problem that severely affects its usefulness at detecting
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35 compounds at low concentrations. Both of these issues can be overcome using SERS (surface
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37 enhanced Raman scattering). Here, the analyte of interest is brought into close proximity to
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39 metal nanoparticles whose plasmon coupling with laser irradiation is responsible for
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41 enhanced Raman scattering effects which shows a great increase in analyte detection
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43 sensitivity over conventional Raman.^{20, 21} Optimisation of SERS systems is crucial to ensure
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45 that the reproducibility of signal and low detection limits are achieved. SERS analyses in
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47 solution are often highly dynamic so control of variables can often be difficult. The main
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49 component of a solution based system are the metal colloid, aggregating agent and analyte,
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51 all of which need to be optimised. Here it is demonstrated that the optimisation of a SERS
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3 system can be accomplished using a systematic approach for the rapid detection of MDAI at
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5 very low concentrations. The portability of Raman instruments coupled with the added
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7 sensitivity of SERS makes this methodology much more amenable than other analytical
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9 techniques to on-site sampling; that is to say within a club or home setting.
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11 12 13 **Experimental**

14 15 **Materials**

16 Silver nitrate (99.9999%) and trisodium citrate were purchased from Sigma Aldrich (Dorset,
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18 U.K.). A 100 mg capsule of MDAI (5,6-methylenedioxy-2-aminoindane) sold as ‘Sparkle’
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20 was purchased from a ‘headshop’ (Dr Herman’s, Manchester, U.K.). The drug contained
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22 within the capsule had a white flaky appearance. The purity of the drugs was verified via
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24 melting point tests, mass spectrometry (MS) and nuclear magnetic resonance spectroscopy
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26 (NMR). All solvents used were of analytical grade and water was HPLC certified.
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29 30 31 **Methods**

32 33 **Drug Purity Verification**

34 Although the drug was advertised as being supplied in 100 mg amounts, the weight of the
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36 drug without the capsule was only 73 mg therefore the analytical techniques used to verify
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38 purity had to be selected carefully. Both MS and NMR were used to derive the structure of
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40 the drug and to ensure that no other impurities were present. The melting point tests gave an
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42 accurate idea of the purity due to the sharpness and temperature at which the sample melted.
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44 The values obtained from these analyses could be directly compared to the original synthesis
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46 values.¹³
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49 50 51 **Mass Spectrometry**

52 The samples were analysed using electrospray ionisation mass spectrometry (ESI-MS)
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54 operating in positive mode. Two peaks were identified in the spectrum at m/z of 161 and 178,
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56 relating to MDAI minus the protonated amine moiety and protonated MDAI respectively.
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¹H NMR (200 MHz, D₂O)

δ 6.73, (s, 2, ArH), 5.84 (s, 2, CH₂), 4.05 (m, 1, CH), 3.19 (dd, 2, 2xCH, *J* = 15.4 Hz, 6.6 Hz), 2.84 (dd, 2, 2xCH, *J* = 15.4 Hz, 5.2 Hz), 2.15 (s, 2, NH₂)

¹³C NMR (300 MHz, D₂O)

Bracketed numbers relate to the positions of the carbons outlined in Figure 1.

δ 146.7 (1), 131.9 (2), 105.4 (3), 101.1(6), 52.1 (5), 36.9 (4)

Melting Point Test

Five replicate melting point tests were carried out on ~3mg of the drug per test. The melting point was sharp and averaged 275 °C only 1°C lower than the original synthetic value.¹³

Synthesis of Silver Colloids

All glassware was cleaned using aqua regia to remove any residual trace metals. After 1 h of treatment the flasks were then washed with copious amounts of methanol, dried under a stream of nitrogen then rinsed with water. To ensure all the solvents had evaporated, the flasks were placed in a temperature-controlled oven (60°C) for 20 min. Silver nanoparticles were synthesised using the Lee and Meisel method.²³ Initially AgNO₃ (90 mg) was dissolved in 500 mL of water and brought to the boil. Under vigorous stirring a 1% solution of trisodium citrate (10 mL) was added. The solution/sol was left to boil for 1 h, the formation of nanoparticles was verified when the previously transparent solution developed a milky green hue. The method was replicated for the synthesis of five batches of silver colloid.

UV-visible (UV-vis) Absorption/Extinction Nanoparticle Characterisation

In order to determine the position of the plasmon band λ_{\max} it was essential to characterise the nanoparticles using UV-vis spectrophotometry. Samples were prepared by combining 1 part

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3 silver colloid with 9 parts water. 1 mL of the dilute nanoparticle solution was then pipetted
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5 into a quartz cuvette and inserted into a sample holder of a Thermo Biomate 5 (Thermo
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7 Fisher Scientific Inc., Massachusetts, USA). A spectrum was collected for each of the 5
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9 colloidal batches. Figure 2 shows the typical UV-Vis spectra obtained whilst the table details
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11 the λ_{max} and full width, half maximum (FWHM) for each batch of colloid.
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14 **SERS Analyses**

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17 Raman spectra were collected using a DeltaNu Advantage benchtop Raman spectrometer
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19 (Intevac inc, California, USA). The instrument is equipped with a 633 nm HeNe laser with a
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21 power output of 3 mW at sample. Spectra were collected over a range of 200– 3400 cm^{-1} with
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23 a spectral resolution of 10 cm^{-1} . Solution samples were placed in an 8 mm diameter glass vial
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25 and subjected to laser irradiation once loaded into the sample cell attachment. The instrument
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27 was calibrated to determine the optimum distance from the laser to the glass vial using
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29 toluene and polystyrene. Raman spectra of the solid MDAI sample were also taken on the
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31 same system but unfortunately the signal response was a broad and featureless (see an
32
33 example spectrum in Figure S1). It was due to the lack of specific vibrational features and
34
35 strong fluorescent background that analyses had to be carried out in solution using SERS,
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37 which is known to quench fluorescence and also enhance the Raman signature.
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43 **Optimisation of Aggregation**

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46 The control of such a dynamic system present in solution based SERS is essential to ensure
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48 maximum reproducibility. One of the ways of managing reproducibility is to optimise the
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50 aggregation time. Variation in SERS signal can result from differing batches of colloids
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52 therefore 5 batches of silver colloids were synthesised and tested along with differing
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54 concentrations of KNO_3 aggregating agent (0.5 M and 1.0 M) with a set analyte concentration
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56 of 500 ppm (2.8×10^{-3} M). Although many different aggregating agents could have been used,
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3 previous experiments carried out in the group found that systems including KNO_3 gave the
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5 best SERS response (data not shown). To reduce any Raman/SERS signal variability as a
6
7 result of differing the volume of components, the colloid and analyte volume were kept at
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9 200 μL and aggregating agent at 50 μL , resulting in 450 μL of experimental solution being
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11 interrogated in total. The order in which the individual components were added to the glass
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13 vial was also kept constant. Initially the colloid was added followed by the MDAI solution
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15 then the aggregating agent. To allow time for the nanoparticles and analytes to equilibrate in
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17 solution a 40 min lag phase was included before the aggregating agent was added. Raman
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19 spectra was collected on each of the samples over a period of 40 min with each spectra
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21 generated over a 30 s interrogation period. This resulted in 80 spectra being collected for each
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23 sample. A total of 10 samples were scrutinized. A definition of what the proposed optimum
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25 aggregation time is and the methodology for its discovery is given in the results and
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27 discussion section.
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32 **Reproducibility Studies**

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36 Once the optimum aggregation time had been established for each experimental system,
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38 replicate sample reproducibility was tested. In order to do this, samples were made up exactly
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40 as outlined in the aggregation study, whilst varying the time that the samples were left to
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42 aggregate for their identified aggregation period before collecting a SERS spectrum of the
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44 sample for 30 s. Five replicate samples from each batch of sol at the differing salt
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46 concentrations were used to assess reproducibility. A total of 50 samples were interrogated (5
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48 replicates x 5 colloidal batches x 2 salt concentrations).
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Limit of detection (LOD) studies

Once the reproducibility of the SERS systems had been assessed, the best system was used to evaluate the LOD of the drug. The drug concentrations analysed ranged from 500 ppm (2.8×10^{-3} M) to 1 ppm (5.6×10^{-6} M). Five replicate samples were analysed at each concentration, and as before the optimised aggregation times were used. Analyte concentration did not affect the optimum aggregation time used.

Results and Discussion

Optimisation of Aggregation Time

Aggregation of nanoparticles is essential to produce 'hot-spots' from which the Raman signal of an analyte is enhanced. The addition of a salt is a common method of inducing aggregation, however the clustering of nanoparticles needs to be controlled if the enhanced signal is to be reproducible.

The initial challenge was thus to identify the optimum aggregation. The word optimum in this instance defines the time at which the SERS signal plateaus, yielding the most reproducible SERS response. The 40 min lag phase introduced before the addition of an aggregating agent was to allow the maximum number of MDAI molecules to associate with the nanoparticles and displace citrate molecules used to stabilise the metal entities, by doing this it was hoped that little variation and shifting of the SERS peaks would arise. Spectra generated on the DeltaNu Raman spectrometer were saved and exported in a .spc format. Data were analysed using Matlab version 2011a (The MathWorks, Inc., Natick, Massachusetts, USA). Once the spectra had been collected from the 10 SERS systems, the 80 spectra representative of each individual system were averaged, to elucidate the peak positions. A staggered plot of mean

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3 spectra for each of the colloidal batches is shown in Figure 3 (A shows spectra generated
4 when 0.5 M aggregating agent was used and B shows spectra generated when 1.0 M
5 aggregating agent was used). Every single peak that was present in the mean spectra was
6 assigned a maximum and given defined start and end points (minimums). The peaks were
7 then extracted and baseline corrected using an asymmetric least squares method. Although at
8 this stage in the analysis it was not possible to clarify whether the bands present were the
9 result of citrate scattering, MDAI scattering or in fact a combination of both.
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19 Spectra collected from colloidal batches 2 and 3 generated no spectral features when
20 combined with salts 1.0 M KNO₃ and 0.5 M KNO₃ respectively, repeat collections of the
21 spectral data sets also proved unsuccessful, so these systems were omitted from further
22 analysis. To interpret the optimum aggregation time for the systems, plots of peak area vs.
23 time (s) were generated for each identified peak and manually assessed, with the objective of
24 verifying the time at which the SERS signal reached a plateau. The time which was identified
25 at the centre of the plateau region was designated as the optimum aggregation period.
26 Identified times were then averaged across all the peaks identified in the individual systems.
27 To generate values for peak area trapezoidal integration was used. This method splits the area
28 under a peak into multiple trapezoidal components, from which the individual areas are
29 calculated then summed giving an overall area value between the specified minima of a
30 single peak. An example aggregation plot, demonstrating the position of the plateau region
31 can be seen in Figure 4 together with a table summarising the optimum aggregation times for
32 all the colloidal batch variations.
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50 51 **Reproducibility Studies** 52

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55 To study the reproducibility of SERS signal 5 replicates of each SERS system were used. In
56 addition to the 40 min lag phase, each of the systems were allowed to aggregate for the
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3 defined optimum aggregation period after the salt solution was introduced. Spectra were
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5 analysed in a similar way as described previously except this time the relative standard
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7 deviations (RSDs) of each peak area were calculated and used to assess reproducibility
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9 between the different batches of sol. All peaks with a RSD <15% were deemed reproducible
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11 and were tallied for each system. As the number of peaks present in the spectra of each of the
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13 systems appeared to vary quite significantly, the number of peaks <15% were calculated as a
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15 percentage of the total number of peaks visible. One reason for the presence of so many
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17 different peaks in the spectra and the inability to assign them all to MDAI is due to the
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19 dynamic nature of the solution phase system. As the citrate stabilised nanoparticles are mixed
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21 into a solution of MDAI, the citrate and MDAI undergo rapid exchange on the surface. It is
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23 expected that MDAI would have a higher affinity for the silver due to it containing an amine
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25 group, therefore it is also expected that greater numbers of MDAI will eventually reside on
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27 the nanoparticles, than citrate. However, it must be remembered that neither citrate nor
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29 MDAI molecules are covalently bound to the nanoparticles, but instead are loosely
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31 associated; this 'association' along with the dynamic exchange also means that the orientation
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33 of the molecules on the surface is constantly changing. One advantage of carrying out these
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35 experiments in solution is the averaging effects achieved from Brownian motion and a large
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37 laser sampling volume; however, this does not mean that the individual systems will display
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39 exactly the same numbers of spectral features. Table 1 shows the results of the reproducibility
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41 testing and it is evident that colloidal batch 1 with 0.5 M and 1.0M KNO₃ demonstrates the
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43 best reproducibility of all the batches with 71% and 69% of the peaks present displaying
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45 RSDs of <15%. Colloidal batch 4 combined with 1.0M of salt can be seen to have the worst
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47 reproducibility with only 1 peak in 16 having an RSD less than 15%. Therefore the system
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49 which consisted of Batch 1 and 0.5M salt was used to establish the LOD of MDAI. Although
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51 it was initially thought that higher concentrations of KNO₃ would hasten the aggregation time
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3 and effect reproducibility in some way, from this study no conclusions could be drawn as the
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5 effect this increase has on the overall MDAI signal. The levels of variation in SERS signal
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7 seen with the differing batches of colloid are due to the difference in size and distribution of
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9 the synthesised nanoparticles. Work carried out previously within the group highlighted the
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11 variability of colloidal batches and in doing so also emphasised the need for careful
12
13 assessment of the nanoparticles.²⁵
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16 17 **Limit of detection (LOD) studies**

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19 When the concentration of MDAI was lowered it became evident which seven peaks in the
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21 spectra were representative of the analyte. Figure 5 shows a mean blank spectrum (200 μL of
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23 colloidal batch 1, 200 μL of water and 50 μL of 0.5 M KNO_3) overlaid with a 500 ppm SERS
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25 spectrum of MDAI (200 μL of colloidal batch 1, 200 μL of 500 ppm MDAI and 50 μL of 0.5
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27 M KNO_3). The red bands highlight the peaks in the plot from which the LOD of MDAI was
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29 established. Table 2 shows the LOD and band assignments for each of the peaks analysed.
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31 The LOD was estimated using the baseline corrected peak intensities at each concentration
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33 and by applying Equation 1. Where SD is the standard deviation of the colloidal blank, c the
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35 intercept and m the gradient. The SD of the colloidal blank was estimated using the same
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37 peak positions and intensities used when MDAI was present.
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$$43 \quad LOD = \frac{((3 * SD \text{ of blank}) \pm c)}{m} \quad (1)$$

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49 The LODs calculated for the 7 peaks identified ranged from ~ 20 to 6 ppm (1.42×10^{-5} M to
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51 3.19×10^{-4} M). The average LOD estimated from all the peaks was 8 ppm (5.4×10^{-5} M).
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53 Example LOD plots are displayed in Figure 6. Figure S2 shows mean spectra collected at
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55 each MDAI concentration down to the LOD (including a colloidal blank)
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Conclusion

In this study it has been demonstrated that SERS can be used to detect the presence of MDAI in solutions at concentrations lower than previously reported methods of 200 ppm by microcrystalline testing.¹⁸ It has also been shown that signal variations can occur between different batches of silver sol synthesised using the same preparative methods, and this emphasises the need for optimisation in order to improve signal reproducibility. A technique's reproducibility and sensitivity ultimately influences its widespread usage in the analytical field, so optimisations like the ones carried out here are important. The optimised SERS method was also demonstrated to be clearly quantitative and the typical limit of detection for MDAI was 5.4×10^{-5} M. Overall a cheap, facile, sensitive and reproducible method for the detection of a substance has been produced. Moreover, due to the ready portability of Raman spectroscopy and SERS this approach could be deployed in the field. Further work would involve the evaluation of the discriminatory properties of such an optimised system, which we have previously reported for other drugs containing the same structural moieties using chemometric methods.²⁶

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Supporting Information

In the supporting information accompanying this manuscript there is a Raman spectrum of MDAI powder (Fig S1) and the SERS spectra of MDAI collected at different drug concentrations (Fig S2)

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Figures

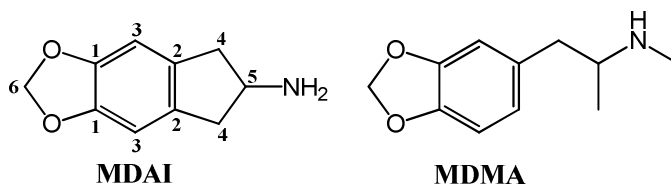
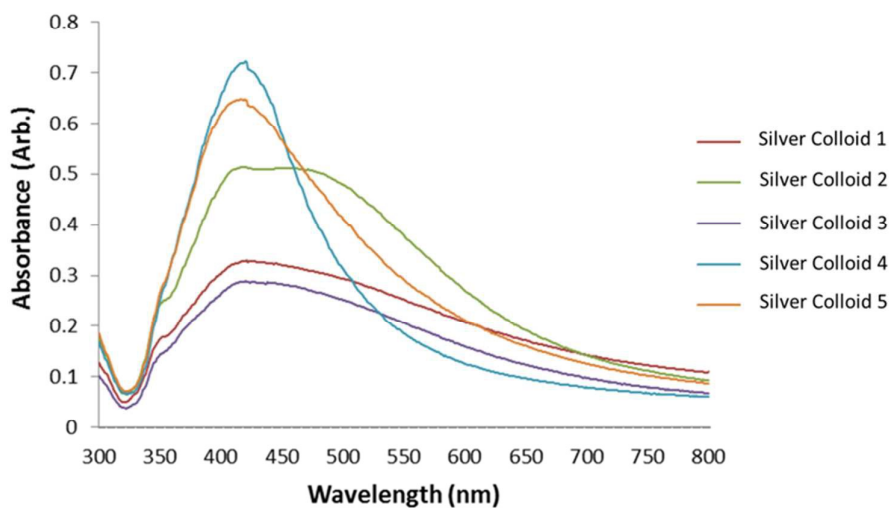


Figure 1 The structure of MDAI with numbers for NMR assignment and the amphetamine

MDMA



Colloidal Batch N ^o	λ_{\max} (nm)	FWHM (nm)
1	421	312
2	418	257
3	420	268
4	420	122
5	417	173

Figure 2 UV-vis spectrophotometry results for the five silver colloidal batches.

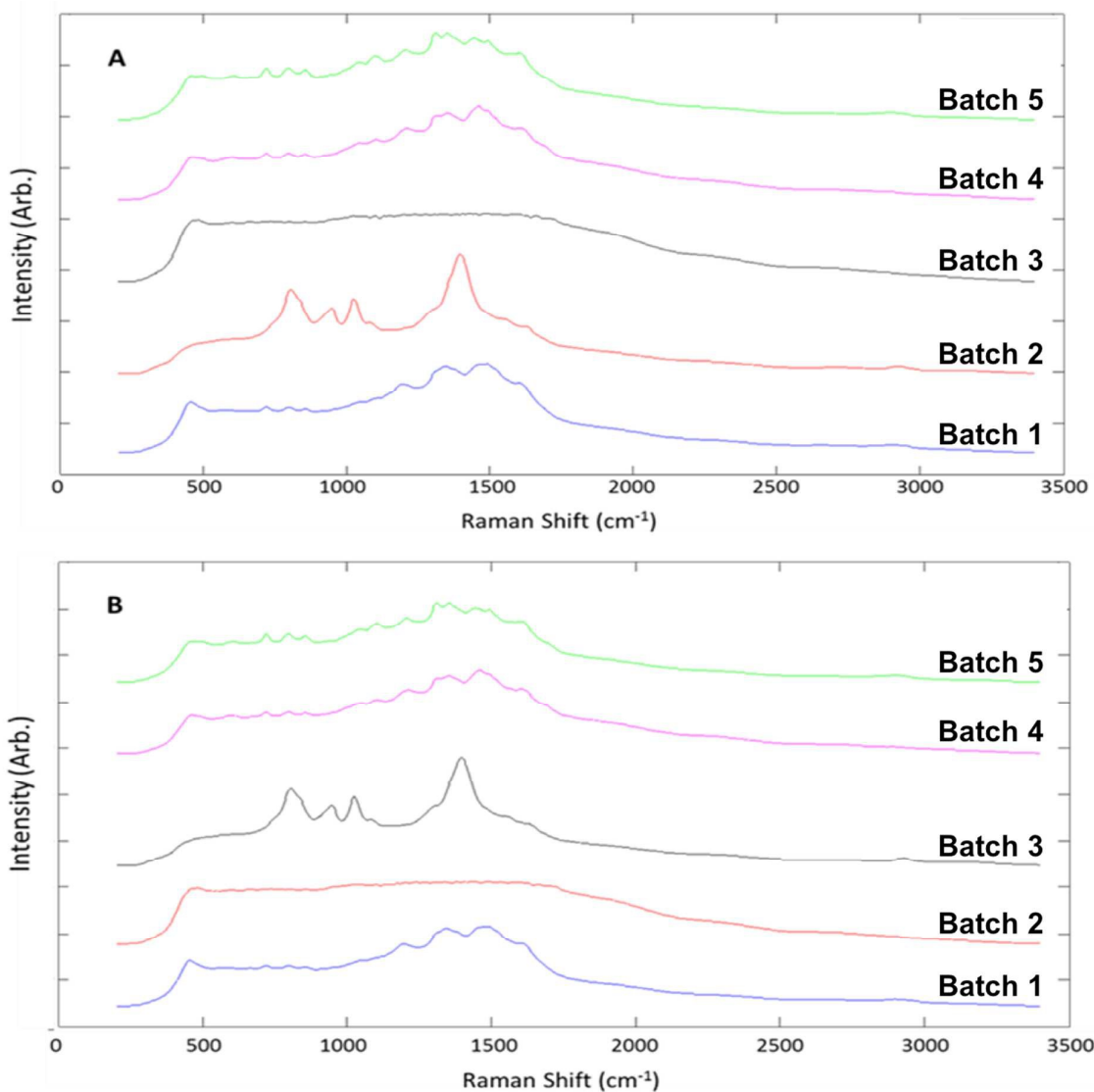
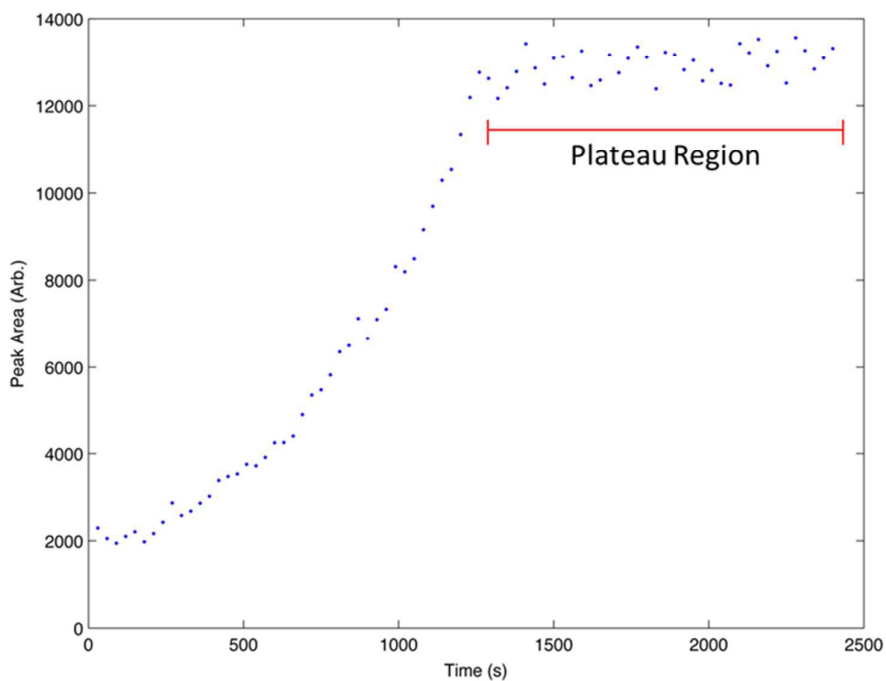


Figure 3 Average scaled spectra of each batch of colloid generated through the optimisation of aggregation experiment. (A) Represents spectra collected using 0.5 M aggregating agent (KNO₃) and (B) Represents spectra collected using 1M aggregating agent (KNO₃). Each spectrum was taken in the presence of 500 ppm (2.8×10^{-3} M) MDAI.



Colloidal Batch N ^o	KNO ₃ (M)	Optimum Aggregation Time (s)
1	0.5	1800
1	1	1650
2	0.5	1860
3	1	1000
4	0.5	1700
4	1	1600
5	0.5	1500
5	1	1400

Figure 4 An example plot of peak area *versus* time for the determination of optimum aggregation time. The plot was generated using the peak area at 1609 cm⁻¹ using colloidal batch 1 and 0.5 M KNO₃. The red line outlines the plateau region where the standard deviation relating to peak area is at its minimum. The time at the centre of this plateau region is estimated to be 1800 s or 30 min, this is defined as the optimum aggregation time. Optimised aggregation times are detailed in the adjoining table for the different colloidal batches and respective salt concentrations..

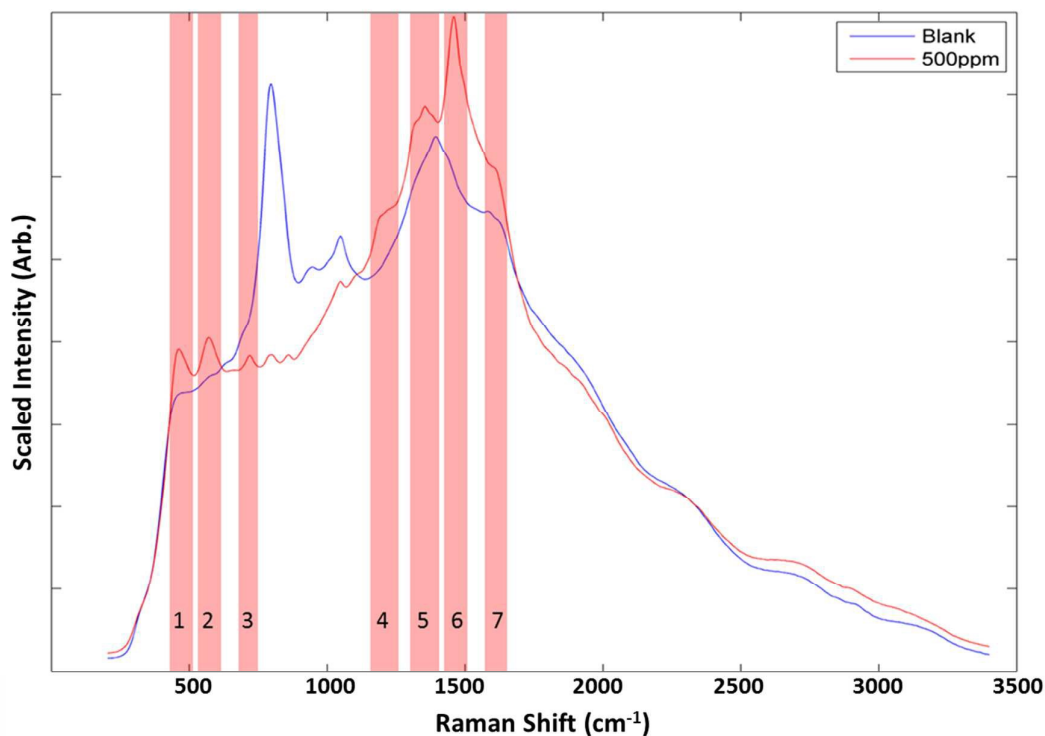


Figure 5 The plot shows a scaled overlay of SERS spectra for the optimised blank colloidal system in blue (colloidal batch 1, 0.5 M KNO₃ and no MDAI) with the optimised colloidal system containing MDAI in red (colloidal batch 1, 0.5M KNO₃, and 500 ppm MDAI). The peaks used for the LOD studies are highlighted by the red bands numbered 1-7. The peaks are positioned at 456 cm⁻¹, 565 cm⁻¹, 715 cm⁻¹, 1190 cm⁻¹, 1353 cm⁻¹, 1459 cm⁻¹ and 1609 cm⁻¹. At low concentrations of MDAI, the citrate peaks from ~1100-1650 cm⁻¹ appear more prevalent making it difficult to assign the MDAI peaks in this region.

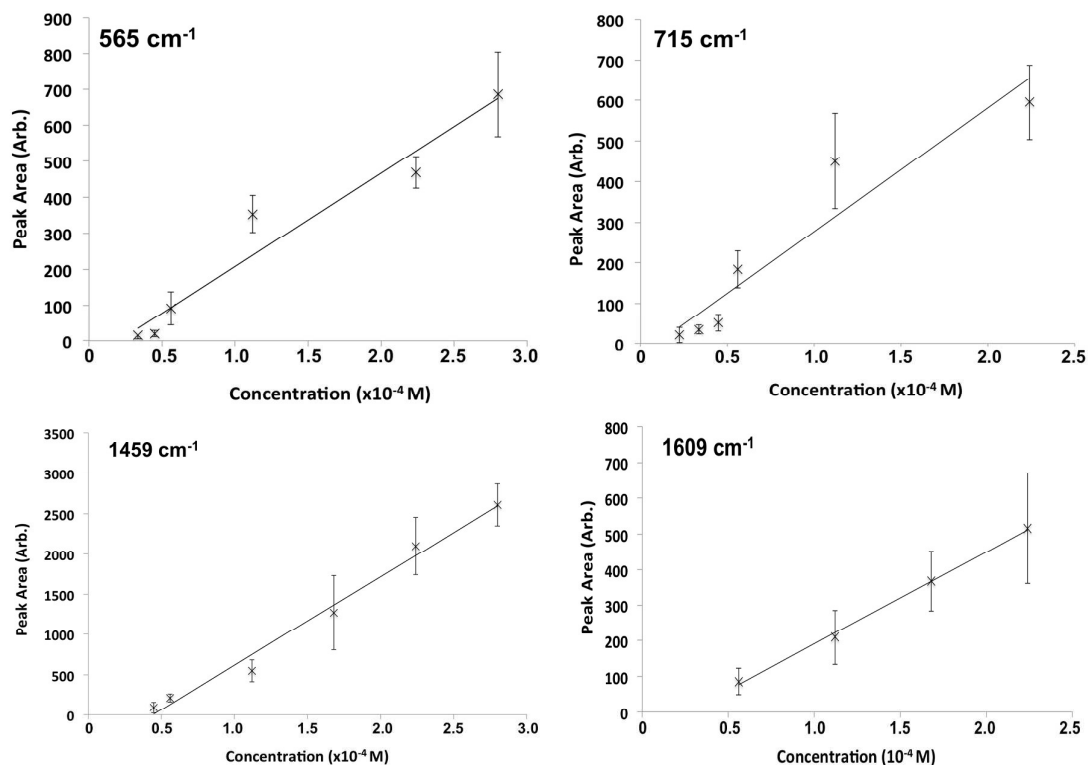


Figure 6 Example plots of peak area *versus* concentration for the four of the seven identified MDAI peaks. The y axes represent peak area (Arb.) whilst the x axes represent the concentration of MDAI ($\times 10^{-4}$ M). Whilst higher concentrations of MDAI were easy to distinguish, it is believed that the drug at these concentrations completely saturated the colloid surface, making the SERS signal non-linear with respect to concentration, therefore only the linear range is plotted.

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Tables

Table 1 The reproducibility of the peaks present in each of the systems is assessed to find the best system. RSDs are assessed using the peak areas calculated for every single peak present in the 5 replicate spectra collected. Peaks with an RSD <15 were deemed acceptable.

Colloidal Batch N°	KNO ₃ (M)	Peaks Present in Spectra		
		Total N°	N° with RSD <15	% of peaks with RSD <15
1	0.5	14	10	71
1	1	13	9	69
2	0.5	14	4	29
3	1	16	5	31
4	0.5	12	7	58
4	1	16	1	6
5	0.5	18	6	33
5	1	16	8	50

Table 2 Tentative SERS vibrational assignments for the 7 peaks identified for MDAI.

Peak Position (cm ⁻¹)	Assignment	Estimated LOD (M)
456	Unassigned	3.30x10 ⁻⁵
565	Unassigned	3.19x10 ⁻⁵
715	Substituted benzene deformation	3.80x10 ⁻⁵
1190	C-N stretch or dioxolane ring vibration	1.42x10 ⁻⁴
1353	Unassigned	4.53x10 ⁻⁵
1459	1,2,4,5-tetrasubstituted benzene vibration	5.31x10 ⁻⁵
1609	C=C aromatic stretch	3.75x10 ⁻⁵

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