

1 Investigation of Multi-Angle Detection Schemes for Light 2 Scattering of Optically Trapped Asymmetric Colloidal 3 Particles

4 **Daniel Maciver^{a*}, Praveen Parthasarathi^a, Leo Lue^a, Jan Sefcik^a, Mark Haw^a**

5 ^aDepartment of Chemical Engineering, University of Strathclyde, 75 Montrose Street, Glasgow, G1 1XL,
6 Scotland

7 **Abstract.** Optical trapping is a well understood method for transduction and detection of forces on trapped
8 colloidal particles, these trapped entities can be further characterised using light-scattering, posing a two-fold
9 challenge: one experimental, concerning the optimal arrangement of detectors to gather data and minimise
10 signal noise, and the other theoretical, involving solving of the inverse scattering problem in order to interpret
11 light scattering data to determine size, shape, or orientation of the trapped object. Experimentally, combining
12 static light scattering techniques with optical trapping poses significant engineering challenges due to the
13 space constraints in a conventional optical trapping setup. We investigated a plausible scenario of detecting
14 scattered light from an optically trapped asymmetric particles using a novel, multi-angle, optical-fibre based
15 detection scheme, we show how a Bayesian inference-based analysis of the data, combined with a neural-
16 network trained on data simulated to mimic light scattering detection signals in such scenarios, can be used
17 for solving the inverse light scattering problem and characterising colloidal trapped entities. To demonstrate
18 the method, we discuss its application to measuring the instantaneous orientations of a trapped asymmetric
19 microsphere dimer and determine the minimum number of detectors required for a reliable estimation in
20 the presence of signal noise. This approach can be extended to determine any characteristics of the trapped
21 microstructure that influence the light scattering pattern, including size and shape of colloidal objects.

22 1 Introduction

23 Since their invention in the late 1980s, optical tweezers have found application in experiments
24 ranging from single molecule biophysics¹ to testing the fundamental assumptions of quantum
25 mechanics,² thanks to the Brownian dynamics of colloidal systems to the tweezer can
26 transduce and detect forces down to the order of a few pico-newtons. Going beyond forces,
27 further structural, dynamic and chemical characterisation of complex trapped entities could
28 provide useful information, as demonstrated in areas such as metrology³ and colloidal
29 science.⁴ Spectroscopic techniques such as Raman scattering⁵ have been used for chemical
30 characterisation of trapped objects, while dynamical characterisation has been demonstrated
31 using data from tweezer’s Quadrant Photo Detector (QPD) by following the centre-of-
32 mass Brownian motion of the trapped entity⁶ and measuring rotation of the centre-of-
33 mass.⁷ A recent work aimed at characterising trapped entities demonstrated how neural
34 networks can be trained to distinguish between optically trapped micro-beads of different
35 size and material by means of a principal component analysis of the forward scattered light
36 detected using a QPD.⁸ A more direct, albeit cumbersome attempt at detecting scattered
37 light from trapped biological cells also was attempted⁹ where the experimental cuvette was
38 placed inside an elliptical mirror that directed light scattered from the trapped biological-
39 cell onto a photodetector via a rotating aperture that helped select the scattering-angle.
40 Thus, past studies on optical trapping have focused on either on particle trapping to study
41 trapping dynamics, or on the characterisation of particles with relatively simple complex
42 trapping dynamics.

43 While both⁸ and⁹ demonstrate some ability to characterise trapped entities,⁸ is perhaps
44 best suited to characterise micron-sized particles with simple trapping dynamics, and⁹
45 describes an experimental setup that is difficult to adopt and suffers from a low bandwidth
46 that might not be best suited for monitoring dynamics. A light scattering detection scheme

47 built around an optical trap that is easier to implement and has the advantage of high
48 bandwidth was demonstrated by Safran and co-workers in,¹⁰ where a single-mode optical
49 fibre was aligned to detect the scattered light from a trapped bead and study its Brownian
50 motion, commonly now referred to as Localised Dynamic Light Scattering (LDLS). This
51 was later expanded upon¹¹ by collecting back scattered light to characterise the Stokes
52 friction coefficient as a function of trapping depth. While both papers provided dynamical
53 information, structural information about the trapped bead was not available as the scattered
54 light was only measured across a small angular range. Furthermore, the main drawback
55 to a LDLS is that it cannot be used to characterise asymmetric particles such as dimers
56 or more complex aggregates; in which case the Brownian motion is no longer simply
57 translational, but rotational about the centre of mass.²

58 One way of mitigating this is to remove translational fluctuations from the analysis;
59 by monitoring its scattering pattern, Cang and co-workers were able to spatially fix a
60 gold nanorod at the centre of trapping laser by moving the sample plane accordingly, to
61 an accuracy of 200 nm.¹² However, this has limited applications to anisotropic scatterers
62 such as biological matter where the internal structure of many cells makes them inherently
63 anisotropic.⁹ Thus, even if the Cartesian coordinates are fixed, characterisation of the
64 scattering pattern will have to separate the contributions of size, structure, and orientation.
65 As an example of the relatively sparse literature on measuring orientations of complex
66 trapped objects,¹³ employs imaging to study the orientation of trapped dimers: scattering
67 can give more quantitative and potentially more rapid time-resolved information, but only
68 of course if the scattering signals can be interpreted. In this work, we propose a novel
69 approach that expands on the previous approaches¹⁰¹¹ to detect scattered light simultaneously
70 at multiple angles (Figure 1), combined with a novel Bayesian inference-based analysis
71 technique, to enable interpretation of the resulting multi-angle data from an anisotropic,
72 asymmetric scatterer, and optimisation to provide maximal information from the detector

73 signals.

74 To demonstrate this approach, we study an anisotropic scattering entity i.e. an asymmetric
75 dimer, to determine dynamic and structural information about the trapped entity. As
76 a paradigmatic example of extracting information from the scatterer's scattering data,
77 we explore how to estimate the dimer's instantaneous orientation from the scattering
78 signals using Bayesian inference, as well as how to optimise the analysis by implementing
79 'prior knowledge' to obtain the most reliable estimate. We, first train a neural network
80 to effectively identify the mapping between scattering signals and dimer orientation, by
81 calculating the scattering signal from a simulated asymmetric dimer undergoing Brownian
82 motion in an optical trap and mapping to the 2 known instantaneous orientation of the
83 simulated dimer. We then show how Bayesian inference can be used to optimise our
84 estimation of the true dimer orientation from the light scattering signals. Furthermore,
85 we demonstrate how the model's performance when dealing with signal noise, a common
86 problem when analysing scattering behaviour. This approach can be extended to determine
87 any characteristic that influences the light scattering pattern of trapped colloidal objects
88 including: size, shape, and orientation.

89 2 Methodology

90 2.1 Orientation estimation from scattering measurements

91 Consider a dimer in the optical trap (Fig. 2a), we can define at any point in time a unit
92 vector \hat{s} pointing from the centre of the larger sphere to the centre of the smaller sphere.
93 A plane wave 'probe' laser, perpendicular to the trapping laser, is incident on the dimer,
94 generating a scattering pattern dependent on the dimer's orientation $I(\hat{s}, \theta)$ which can be
95 computed using software such as MSTM.¹⁴ To represent the experimental set up consisting
96 of a set of optical fibres recording scattered light, we choose four angles ($\theta_1, \theta_2, \theta_3, \theta_4$)

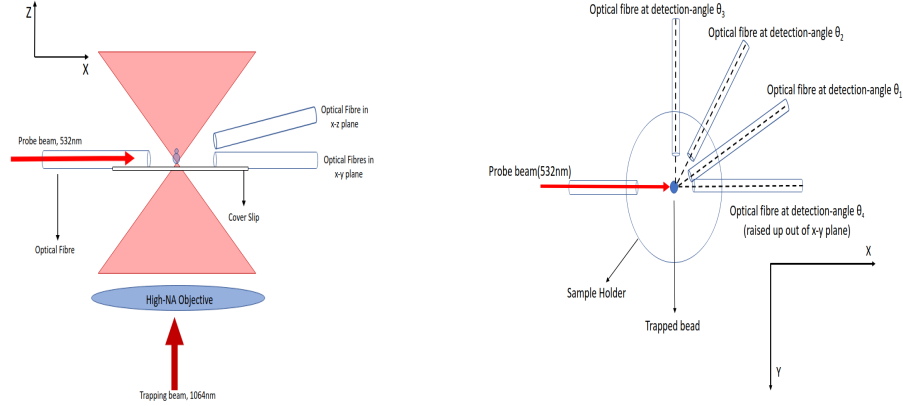


Fig 1: Proposed experimental set up for scattering measurements from an object in an optical trap. The probe beam for scattering measurements is incident perpendicular to the trapping laser propagation direction. a) Side view. b) Top view. Note that three of the detector fibres are co-planar with the incident probe beam, while the fourth detector is placed out of the plane (see Sec 3.1).

97 and record the calculated intensity at each angle θ_k , $I(\hat{s}, \theta_k)$.

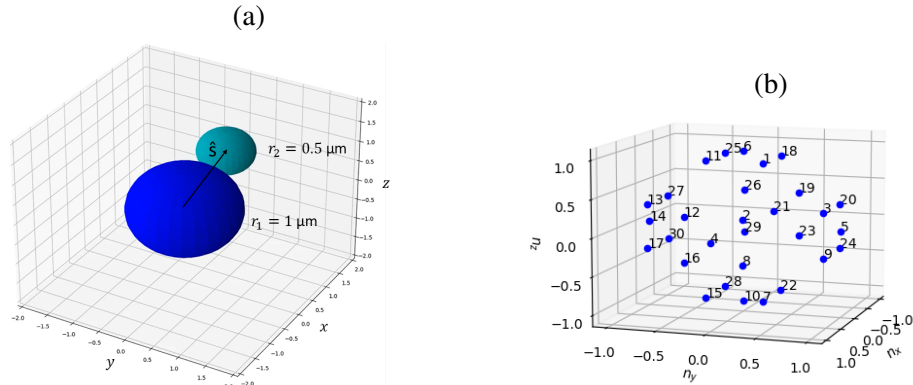


Fig 2: (a) Example dimer in orientation \bar{s} , (b) 30 Reference orientations represented by vectors pointing from $[0,0,0]$ to each point

98 Our goal is to determine the orientation of the trapped dimer based on the measured
 99 intensity $I(\hat{n}, \theta_k)$. Rather than aim immediately for an exact estimate of the dimer's
 100 orientation, for the purposes of interpretation of the scattering and optimisation of the
 101 measurement setup it is more convenient to discretize the possible orientation space into

102 a number of possible reference orientations, which we can then use as 'classification
 103 categories' in a neural network methodology to map scattering data to orientation (see
 104 below for further discussion). Here we choose $n_{ref} = 30$ reference orientations $\hat{\mathbf{n}}_\alpha$
 105 evenly distributed on a unit sphere¹⁵ (Figure 2b) leading to a maximum nearest-neighbour
 106 spacing between two neighbouring reference orientations of 0.895 radians. Using MSTM
 107 we compute the raw intensities at each of the measurement angles that would be generated
 108 by a dimer in each reference orientation, $I(\hat{\mathbf{n}}_\alpha, \theta_k)$. While the number and position of
 109 detection fibres is technically arbitrary there are several constraining factors that limit our
 110 ability to infer useful information from the trapped object, see Section 3.1 for a detailed
 111 breakdown of our choice of detection angles. The raw intensities are normalized according
 112 to:

$$y_k(\hat{\mathbf{n}}_\alpha) = \frac{I(\hat{\mathbf{n}}_\alpha, \theta_k) - \langle I(\hat{\mathbf{n}}, \theta_k) \rangle}{\langle I^2(\hat{\mathbf{n}}, \theta_k) \rangle - \langle I(\hat{\mathbf{n}}, \theta_k) \rangle^2} \quad (1)$$

113 where the denominator is simply the standard deviation across the set of values $I(\hat{\mathbf{n}}, \theta_k)$.
 114 The reference orientations, raw intensities, and scaled signals are given in Tables A1 and
 115 A2.

116 Note that the collected scattering signals are not necessarily simply related to their
 117 associated reference orientations: as is well known from such examples of the inverse
 118 scattering problem. While it is trivial to compute the light scattering pattern for any
 119 given particle with any particular characteristic (i.e. size, shape, or orientation), inferring
 120 the light scattering from a unknown particle to determine said characteristic is incredibly
 121 difficult due to complex mapping between scattering and said characteristic. Even if the
 122 orientation space is divided evenly between reference orientation the subsequent signal
 123 space ends up being appearing mixed making simple comparisons of signals useless for
 124 inferring information on the particle. Shown below is two clusters of orientation vectors

125 and there respective measured scattering signals - the points have been coloured based
 126 on their proximity to the centre of their respective cluster. While the orientation space
 127 appears tightly packed and ordered the signal space quickly spreads out in an asymmetric
 128 fashion. Furthermore as seen in Fig 3b the signal mapping can intersect itself which
 129 only further increases the complexity. While in some instances the mapping between one
 130 reference orientation and another is discrete, in other instances the mapping becomes far
 131 more complex to discern.

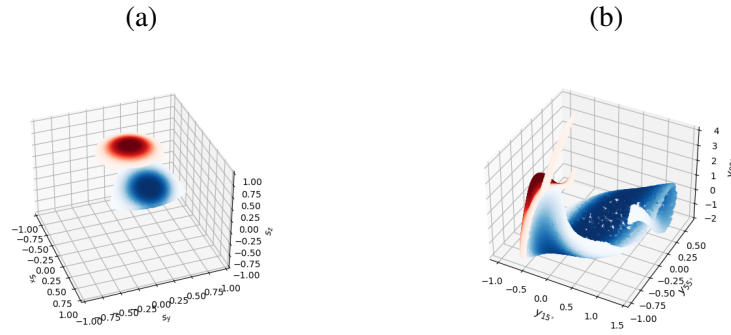


Fig 3: (a) Distribution of orientation vectors and (b) their respective scattering signals. Points are coloured according to their distance from the centre of each cluster (red points centred around $[0.00, 0.00, 1.00]$, blue points centred at $[0.71, 0.00, 0.71]$)

132 Nevertheless, at least where the uncertainty in signal measurements is low (see below),
 133 we can predict the orientation from the scattering by utilising computational techniques
 134 such as neural networks. We thus utilised the Python machine learning program *scikit-*
 135 *learn* to build a neural network for identifying the dimer's orientation from its light scattering
 136 signal. The network was trained by generating a database of random orientation vectors,
 137 calculating the corresponding light scattering signals, and then using the network to estimate
 138 the probability of a given signal coming from a dimer in a given reference orientation. The
 139 network's loss function was evaluated and used to improve the estimation, the network
 140 being trained until the improvement in the loss function was less than 0.0001. Importantly,

141 the estimation provided by the neural network can be improved further by accounting for
 142 any prior information we know about the dimer, utilising Bayesian inference to update the
 143 neural network's estimation:

$$p(\hat{\mathbf{n}}_\alpha | y_k(\hat{\mathbf{s}})) = \frac{p(y_k(\hat{\mathbf{s}}) | \hat{\mathbf{n}}_\alpha) p(\hat{\mathbf{n}}_\alpha)}{p(y_k(\hat{\mathbf{s}}))} \quad (2)$$

144 where $p(\hat{\mathbf{n}}_\alpha)$ and $p(y_1, y_2, y_3)$ are the prior estimates of the distributions of particle orientations
 145 and instantaneous signals, respectively. *Without* any prior evidence we must assume that
 146 the orientation prior of the dimer $p(\hat{\mathbf{n}}_\alpha)$ is uniform. However, inference about the dimer's
 147 possible current orientation from knowledge of previous measurements can be used to
 148 inform our estimate of $p(\hat{\mathbf{n}}_\alpha)$ (see Section 3.2). The latter prior $p(y)$ is the probability
 149 of measuring a signal (y_1, y_2, y_3) . This is given by taking the discrete integral over the
 150 collection of reference orientations:

$$p(y_1, y_2, y_3, y_4) = \sum_{\alpha=1}^{n_{\text{ref}}} p(y_1, y_2, y_3, y_4 | \hat{\mathbf{n}}_\alpha) p(\hat{\mathbf{n}}_\alpha) \quad (3)$$

151 From (2) we obtain the key result, a mass probability distribution denoting the probability
 152 that our dimer is in orientation $\hat{\mathbf{n}}_\alpha$ given a measured signal (y_1, y_2, y_3) , *i.e.* an estimated
 153 mapping from scattering measurement to orientation estimate.

154 2.2 Calculation of error

155 To evaluate the above estimation of dimer orientation from scattering signal, we use a
 156 Brownian simulation of a dimer in the optical trap (Section 2.3) to compare estimated
 157 most probable reference orientation, derived from the dimer's scattering through Eq. (2),
 158 with the dimer's known *actual* orientation $\hat{\mathbf{s}}$. MSTM provides calculated light scattering

159 from the simulated dimer $I(\hat{s}, \theta)$ and we use (1) to obtain normalized values at each
 160 measurement angle θ_k , $y_1(\hat{s})$, $y_2(\hat{s})$, $y_3(\hat{s})$, from which we obtain $p(\hat{\mathbf{n}}_\alpha \parallel y_1, y_2, y_3)$.
 161 Because we know the actual orientation \hat{s} we can measure the error in the model's estimate
 162 by comparing the reference orientation closest to \hat{s} , denoted as $\hat{\mathbf{n}}_{best}$, with the most probable
 163 predicted orientation from Eq. (2). An ideal result would be one where the probability
 164 distribution is 0 for every $\hat{\mathbf{n}}$ apart from $\hat{\mathbf{n}}_{best}$:

$$p_{best} = \begin{cases} 1 & \text{when } \hat{\mathbf{n}}_\alpha = \hat{\mathbf{n}}_{best} \\ 0 & \text{anywhere else} \end{cases} \quad (4)$$

165 In reality the distribution from Eq. (2) will assign some non-zero probability to every
 166 reference orientation, leading to some level 'confidence' in orientation prediction, which
 167 can be quantified by calculating the Kullback-Leibler divergence K_l between the two
 168 distributions:

$$K_{l,\#}(p_{best} \parallel p(\hat{\mathbf{n}}_\alpha | y_1, y_2, y_3)) = p_{best} \ln \left[\frac{p_{best}}{p(\hat{\mathbf{n}}_{best} | y_1, y_2, y_3)} \right] \quad (5)$$

169 where a larger value of K_l indicates that our model is less confident in its prediction of the
 170 dimer's orientation. The divergence K_l thus illustrates the 'spread' in the estimated dimer
 171 orientation probability — a distribution strongly peaked at some value would give us more
 172 confidence in that value than a near-uniform distribution where the scattering measurement
 173 could imply a wide range of possible orientations — but it does not directly indicate our
 174 estimates actual accuracy, that can be simply defined as the percentage of our estimations
 175 that are correct.

176 2.3 Brownian Simulation

177 We use the Brownian OT package developed by Fung *et al*¹⁶ to simulate the motion of an
178 asymmetric dimer (Figure 2a) within an optical trap. Brownian OT combines MSTM¹⁴
179 and “Optical Tweezer Toolbox” (*ott*)¹⁷ to simulate the motion of arbitrary shaped sphere
180 clusters. We simulate the motion of a dimer trapped in a highly focused Gaussian beam
181 by calculating the optical forces imparted by the laser, and the Brownian force due to the
182 surrounding fluid. MSTM provides the necessary T-matrix to compute the optical force
183 via *ott*. The Brownian force is found by computing the dimer’s diffusion tensor according
184 to the analytical solutions provided by Nir and Acrivos.¹⁸ We simulated a polystyrene
185 dimer ($n = 1.59$) in a suspension of water ($n_{med} = 1.33$) over the course of 1 s with a
186 simulation time step of 1×10^{-5} s. We placed the dimer 4 microns below the trap focus
187 at an angle 30° from the horizontal, the resulting trajectory is shown below in Sec 3.3.
188 We chose these initial parameters because it demonstrates our model’s performance in non
189 steady state conditions.

190 3 Results and Discussion

191 3.1 Minimal number of detectors

192 The exact number of detectors was initially assumed to be arbitrary, in that it made no
193 difference to our estimate whether we used 2 angles or 200. For practical purposes
194 it seemed beneficial that we demonstrate our method works for a minimal number of
195 detection angles, as geometric constraints come into play when trying to install a high
196 number of detection fibres for any optical tweezer set up.

197 When all of the detectors lie in the same plane the expected signal can appear identical
198 despite the dimer being in completely different orientations. This is shown in Figure 4
199 which plots the expected signals from 30 reference orientations, each point is labelled with

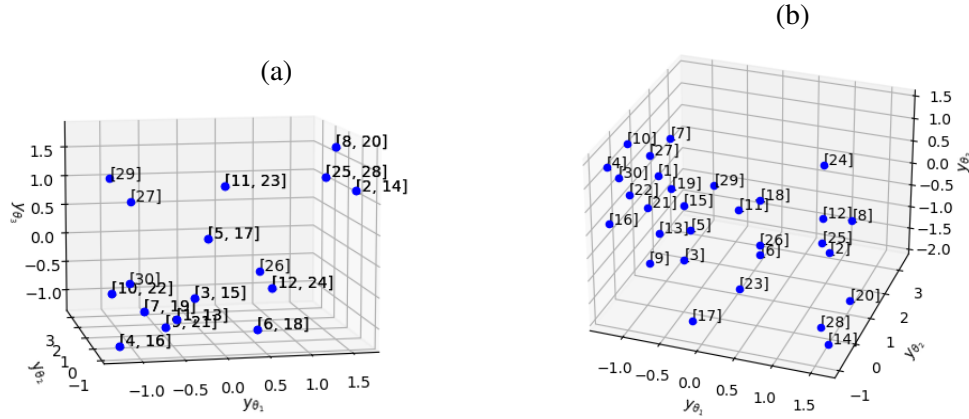


Fig 4: Expected scattered signals from reference orientations -see fig 2 - when: (a) all three detectors are in the X-Y plane, (b) when 1 detector is raised out of the X-Y plane.

its corresponding reference orientation, the fact that points have multiple labels is because the dimer's scattering is indistinguishable in these two reference orientation. It should be noted that these pairs are reflected in one or more axis which suggests that these are due to the arrangement of our detectors. More specifically, if the detectors are placed say in the x-y plane then only when the dimer is pointed nearly fully upright will the expected signal be entirely unique. This is illustrative of the difficulty behind the inverse light scattering problem; as one cannot always map a given signal to a particular parameter value.

To remedy this we raise the third detector out of the x-y plane; as such the expected signals from each reference orientation is unique. As seen between Figures 4a & b each reference orientation now has a unique scattering signal, though with only three detectors the difference in expected signals can appear insignificant. By adding a 4th detector we can differentiate signals more reliably, improving the neural networks performance. In line with our goal of making this method viable in a laboratory setting we decided not to increase the number of detectors further than 4.

214 3.2 Testing the Model

215 Using our simulation from Section 2.3 we simulated the motion of a silica dimer ($n =$
 216 1.45) trapped in water ($n = 1.33$) within a 5 mW optical trap. The trapping laser is 1064nm
 217 NIR focused through a 1.25 NA objective. The dimer is comprised of two tangent spheres
 218 with radii $1\mu m$ and $0.5\mu m$ respectively. We simulated the first 10 seconds of motion,
 219 calculating the orientation and position every 1 ms.

220 We applied Eq. (2), taking the reference orientation with the highest probability as our
 221 estimate of the dimer’s instantaneous orientation \hat{n}_{est} . To visualise the model’s performance
 222 we plotted the radial distance between our estimation \hat{n}_{est} and the dimer’s *actual* instantaneous
 223 orientation \hat{s} versus time. For comparison, we also plotted the radian distance between the
 224 dimer’s instantaneous orientation and the closest reference orientation, denoted \hat{n}_{best} . The
 225 dotted line indicates the maximum radian distance (0.896 radians) between two *neighbouring*
 226 reference orientations: if we are under this line then we know our estimate is at least
 227 neighbouring the best result. Assuming a uniform prior of the reference orientations $p(\hat{n}_\alpha)$
 228 the neural network’s predictions (\hat{n}_{est} from Eq. (2)) are at times reasonable, but there are
 229 significant large and random jumps away from the correct result (Fig. 5).

230 One reason we observe such large jumps in orientation estimated from scattering
 231 signals is that there is no simple correlation between the ‘distance in scattering space’
 232 between scattering signals from two different orientations, and their separation in orientation
 233 space: even a large change in orientation can involve a small change in scattering. Combining
 234 this fact with use of a uniform prior, indicating essentially no knowledge of how orientation
 235 should behave, there is no constraint on how much estimated orientation can change from
 236 time-step to time-step. To improve the estimation we can therefore use knowledge of the
 237 physical limitations of the object in the trap and its dynamics, imposing a more physically
 238 grounded prior, accounting in this case for the fact that the motion of the dimer is limited

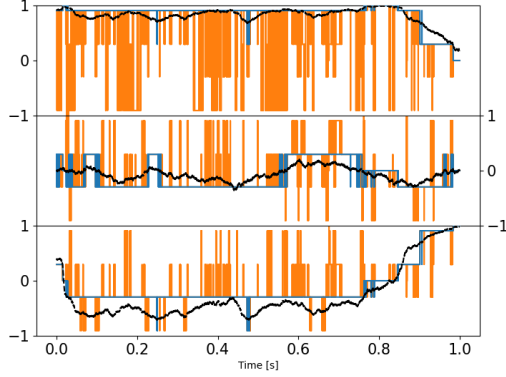


Fig 5: Model's estimation of dimer orientation over the simulation time, assuming uniform prior $p(\hat{\mathbf{n}}_\alpha)$, broken up into x, y, and z components for clarity. Blue line denotes the best result we can achieve (the reference orientation $\hat{\mathbf{n}}_{best}$ that is closest to the actual orientation), orange line denotes the result provided by eq 2: where the orange line is not visible, the model's prediction agrees with $\hat{\mathbf{n}}_{best}$. Dotted black line is the instantaneous orientation $\hat{\mathbf{s}}$.

239 due to the trap stiffness. Here the prior of the reference orientations $p(\hat{\mathbf{n}}_\alpha)$ was redefined at
 240 each time step as a Boltzmann distribution of the physical distance between the previous
 241 estimate $\hat{\mathbf{n}}_{est}(t - \Delta t)$ and each reference orientation $\hat{\mathbf{n}}_\alpha$. Put simply, we are reweighing our
 242 estimation based on the size of rotation required, with smaller movements being favoured
 243 over large movements:

$$p(\hat{\mathbf{n}}_\alpha) = \frac{e^{\beta(\hat{\mathbf{n}}_\alpha \cdot \hat{\mathbf{n}}_{est}(t - \Delta t))}}{\sum_{\alpha=1}^{n_{ref}} e^{\beta(\hat{\mathbf{n}}_\alpha \cdot \hat{\mathbf{n}}_{est}(t - \Delta t))}} \quad (6)$$

244 Here β is a weighting factor describing the dimer's freedom of motion within the trap. As
 245 shown in Figure 6 implementation of Eq (6) helps significantly reduce the large random
 246 excursions of estimated orientation away from the 'best' result.

247 The simulation data from Section 2.3 was used to evaluate our model's performance —
 248 covered in Section 2.2. By summing the divergence of each measurement across the entire

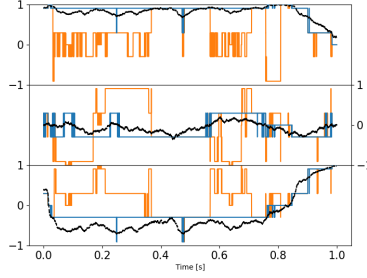


Fig 6: Estimation of dimer orientation with $p(\hat{\mathbf{n}}_\alpha)$ defined by Eq (6). Blue line denotes the best result we can achieve, orange line denotes the result provided by eq 2. Dotted black line is the instantaneous orientation $\hat{\mathbf{s}}$ (see Section 2.1).

simulation we get an evaluation of how well the model performed in estimating the dimer's orientation. To compare the effects of changing certain parameters on the performance of our model we compare our result of $K_{l,total}$ to a worst case scenario and evaluate how much it improves upon this, denoted as $F(K_l)$:

$$K_{l, total} = \sum_{\# = 1}^{timesteps} K_{l, \#} \quad (7)$$

$$K_{l, worst} = \sum_{\# = 1}^{timesteps} \ln \left[\frac{1}{1/n_{ref}} \right] \quad (8)$$

$$F(K_l) = \frac{K_{l, worst}}{K_{l, total}} \quad (9)$$

The worst case scenario is akin to randomly choosing a reference orientation at each time step. The greater the value of $F(K_l)$, the better our model's confidence is in characterising the dimer's motion. Because our model is dependent on several parameters we need to a sophisticated method for understanding how these parameters correlate with $F(K_l)$.

257 3.3 Asymmetric dimer dynamics

258 The Brownian OT software was used to simulate the motion of a trapped dimer ($a_1 =$
 259 $1\text{ }\mu\text{m}$, $a_2 = 0.5\text{ }\mu\text{m}$) over the first 1 seconds of entering the optical trap. The initial
 260 orientation was set at $s = (0.923, 0.0, 0.385)$. The dimer's position and orientation was
 261 recorded every $10\text{ }\mu\text{s}$ for using as a test dataset for our model.

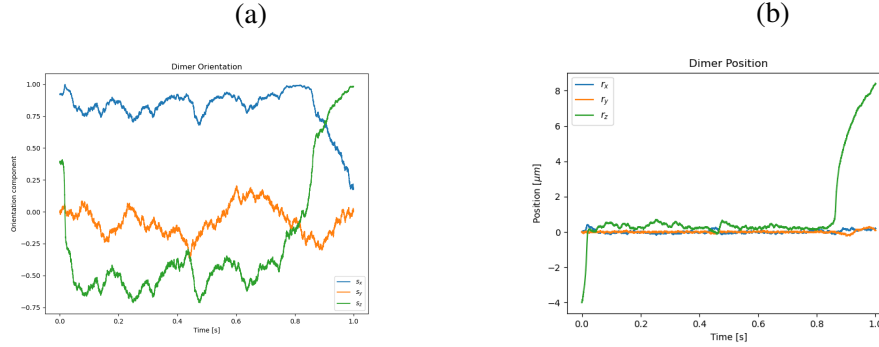


Fig 7: Simulation results of: (a) the dimer's orientation vector with time, (b) the dimer's [x,y,z] position with time.

262 In the simulations of Vigilante *et al.*,¹⁶ trapped symmetrical dimers were investigated;
 263 their findings showed that the optical torque on the dimer goes to zero while aligned
 264 vertically and is at its maximum in a horizontal alignment. However as seen in Figure 7
 265 asymmetric dimers demonstrate dynamics that do not immediately achieve steady state.
 266 We chose to use asymmetric dimers as our benchmark due to this fact, as its orientational
 267 motion is far more complex than a symmetric dimer. In the future we hope to further
 268 investigate the motion of asymmetric dimers.

269 3.4 Accounting for sources of error in light scattering measurements

270 When it comes to analysing light scattering from any size particle, error analysis becomes
 271 a significant factor. Typically this can be accounted for by averaging over long periods of
 272 time to get an assessment of the steady state conditions of the target particle. However in

our case where we wish to know the instantaneous orientation, we instead have to rely on our understanding of how uncertainty can effect our model’s performance. We identified two areas which are likely sources of error in our estimation: firstly, an incorrect modelling of the target particle, and secondly, signal noise arising from experimental factors. We highlight how we address these areas below.

Impact of incorrect dimer sizing One of the main limitations of our model is that we assume that the dimer being modelled in MSTM is accurate to the dimer being trapped in the optical tweezer. Sizing molecules accurately is a significant challenge for single particle analysis so there is bound to be some uncertainty with the measurements. We ran our model 3 times with the neural net being trained on a dimer of size ratio 1 : 1.95, 1 : 2.00 and 1 : 2.05.

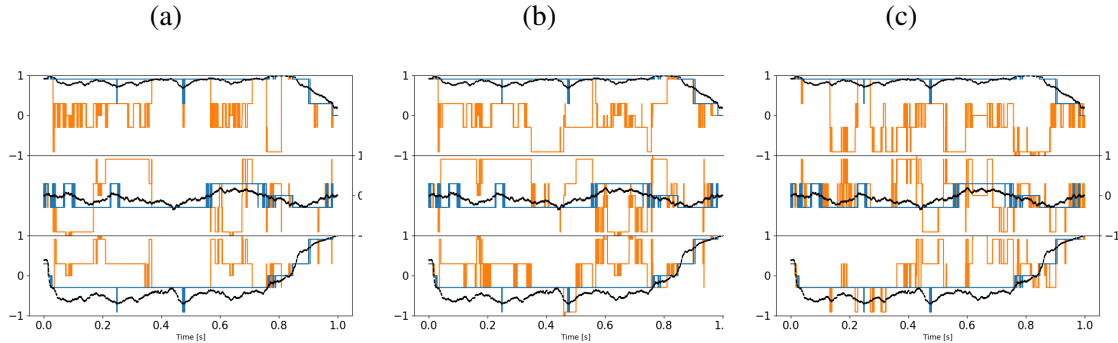


Fig 8: Model estimates of orientation when neural net has been trained on dimer of size ratio: (a) 1:2 [$F(K_l) = 9.456$], (b) 1:2.05 [$F(K_l) = 1.324$], (c) 1:1.95 [$F(K_l) = 1.325$] ($n_{refs} = 30$)

As can be seen from Fig 8 even the slightest change in size ratio makes a very significant difference to the performance of our model. This amounts to just over 100 nm in the dimer’s overall size, yet results in our model being correct from over 90 % of the time to now as low as 30 %. This highlights the importance of correctly sizing trapped entities

288 before performing any in depth analysis of the scattering pattern, as even the slightest
 289 deviation can have a serious impact. We addressed this by increasing the number of
 290 available reference orientations from 30 to 126 (following the same procedure as given
 291 by¹⁵ to evenly space out the coordinates) and increasing the weighting factor in Eq 6.
 292 While this didn't have a significant improvement on the overall accuracy of the model, in
 293 the worst case having a slight increase from 30.5 % to 40.3 %, it did help to significantly
 294 reduce the magnitude between our model's estimations and the dimer's motion as seen
 295 below in Fig 9.

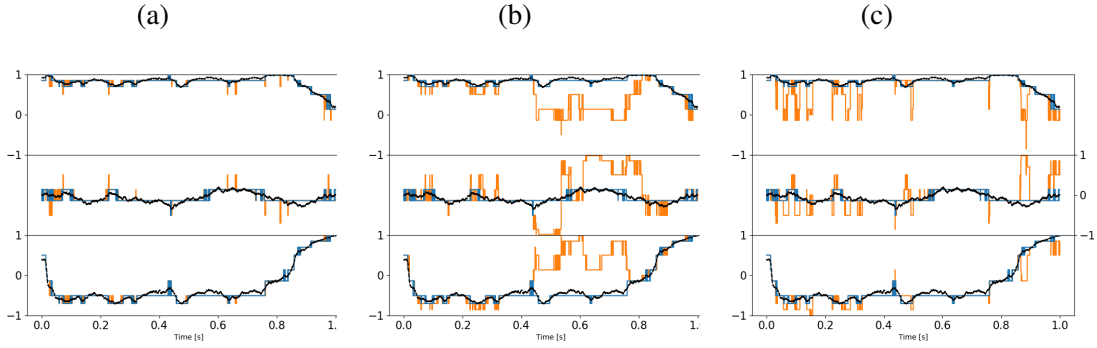


Fig 9: Model estimates of orientation when neural net has been trained on dimer of size
 ratio: (a) 1:2 [$F(K_l) = 11.756$], (b) 1:2.05 [$F(K_l) = 1.233$], (c) 1:1.95 [$F(K_l) = 2.128$],
 ($n_{refs} = 126$)

296 Notably the increasing the number of reference orientations had a greater effect when
 297 our neural network was trained on a 1:1.95 dimer than a 1:2.05 dimer. This suggests that
 298 overshooting our size estimate will be less detrimental to our estimation. Notably if the our
 299 sizing is off the neural network does not predict a smooth motion within the trap; instead
 300 predicting that the dimer is jumping back and forth between different orientations. This
 301 suggest that we can narrow down our estimate of the particle's size by assessing how the
 302 dimer is reorienting within the trap, as we should expect a smooth continuous prediction.
 303 Since we are working with a spherical dimer it also stands to reason that techniques such

304 as image analysis could be used in part to address this, so long as the trapped entity is
 305 sufficiently illuminated.

306 **Impact of measurement noise on model predictions** So far a key assumption of
 307 the neural network implementation is that the detected scattering signal has no uncertainty
 308 associated with it. In reality of course scattering signals will always have some non-zero
 309 measurement noise. This can be attributed to a variety of factors, from a measurement
 310 bias in the detector, to the Brownian motion of the dimer itself. To explore the impact
 311 of measurement uncertainty on orientation estimation model performance we introduce a
 312 Gaussian noise to the measured signal:

$$I(\hat{\mathbf{s}}) = I(\hat{\mathbf{s}}) \pm \epsilon I(\hat{\mathbf{s}}) \quad (10)$$

313 where ϵ is the percentage error associated with the scattering signal. Figure 10 shows the
 314 performance of the model at a range of ϵ using in-plane detector angles 15° , 55° , 90° and
 315 out-of-plane detector at 75° , with β set to 1:

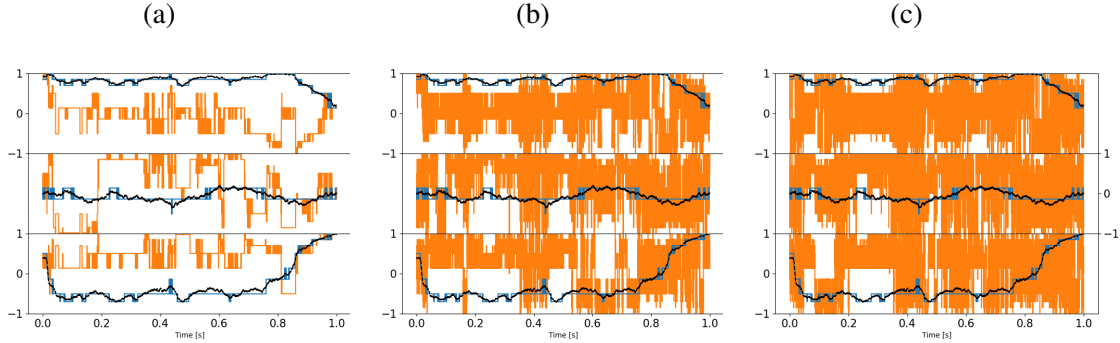


Fig 10: Model prediction for signal error of (a) 1% [$F(K_l) = 7.246$], (b) 15% [$F(K_l) = 0.511$], and (c) 25% [$F(K_l) = 0.536$].

316 As can be seen from Figure 10, the inclusion of signal noise quickly leads to a decrease
 317 in the model's performance. This is due to an inherent feature of the inverse scattering

318 problem: two distinct regions in orientation space can become heavily intertwined and
 319 thus no longer well separated when mapped to intensity space (even though the mapping
 320 remains continuous): so even small uncertainties in the scattering data can lead to large
 321 'mistakes' in the choice of orientation by the neural network. (Indeed if this was not the
 322 case the inverse scattering problem would be quite simple.)

323 To reduce the effects of the signal noise we took the time average of the expected signal
 324 over $0.001s$ and then had our neural network estimate the orientation based on the average
 325 signal.

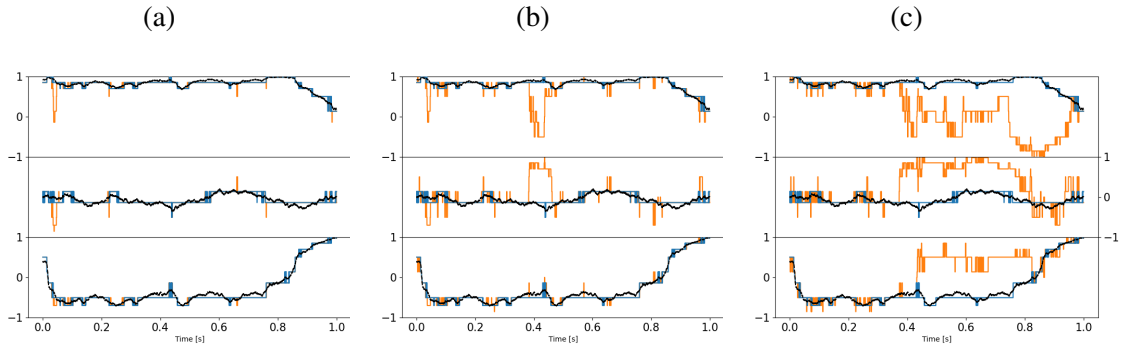


Fig 11: Model prediction for signal error of (a) 1% [$F(K_l) = 4.823$], (b) 15% [$F(K_l) = 1.494$], and (c) 25% [$F(K_l) = 0.882$], time averaged over 1 ms

326 This resulted in a reduction in the overall signal noise and provided a higher degree
 327 of accuracy for our model. There appears to be no clear correlation between the length
 328 over which we time average and the performance of our model. Time averaging over
 329 every $0.05s$ resulted in a drastically worse performance; this is due to the fact that over
 330 longer time periods there is greater uncertainty regarding how the dimer's orientation
 331 has changed, thus tracking the instantaneous orientation becomes harder for the neural
 332 network. Fortunately, time averaging even over $1ms$ seems to provide a satisfactory
 333 estimation of the dimer's angular dynamics within the optical trap.

334 From the above discussion it's clear that estimation of the dimer's orientation is a

335 problem that can be endlessly tuned to fully maximise our end result. Here we simplify the
336 problem somewhat by employing a relatively small finite number of 'reference orientations'
337 to map between scattering and dimer orientation: the precision of estimation could be
338 improved by utilising a greater number of reference orientations, although there remains
339 a balance between the realisable precision of orientation estimate and the noise level of
340 the scattering measurement. Another avenue to further explore would be using the method
341 to optimise the choice of detection angles, essentially to find the region in the mapping
342 between measured scattering and orientation that offers the best degree of confidence
343 through optimal separation of scattering signals for distinct orientations. For sequences
344 of data such as dynamic measurements, a further potential enhancement would be to
345 consider more complex correlations based on prior expectations of the dynamics. Here
346 already we improve the method using a non-uniform prior based on only the immediately
347 previous measurement in time (see Section 2.1): considering a non-uniform grouping of
348 reference orientations might result in a better estimation, if we have information regarding
349 the dimer's preferred axis of rotation.

350 **4 Conclusion**

351 We have developed a method for measuring the dynamics of an optically-trapped colloidal
352 objects based purely on measurements of the object's light scattering at a small number
353 of detection angles. We demonstrate the method using the orientation of an asymmetric
354 dimer as the dynamic variable and object of interest respectively, but in principle the model
355 can be applied to any characteristic that impacts the light scattering pattern produced
356 by a trapped entity such as size and shape. The MSTM package is a flexible tool for
357 calculating the light scattering of complex objects using a representation of the object as
358 a set of micro-particles, enabling training of a neural network to enable categorisation of
359 the mapping between scattering and trapped object characteristics. By taking account of

the physically realistic behaviour of the trapped object and the characteristics of the trap (which impact the dynamics of the object), the Bayesian inference method can be refined to provide a reliable estimation of object characteristics of interest, even in the presence of measurement noise. Fundamentally, the inverse scattering problem is difficult to solve, since the mapping between object characteristics and scattering can be highly complex. We determined the minimum number of detectors required for a reliable estimation in the presence of measurement noise; furthermore, we demonstrated that the arrangement of these detectors is critical for a reliable estimation of an objects orientation. However, Bayesian inference based on neural network estimation of the mapping provides a powerful method for practical applications, extending the use of optical trapping beyond measuring microscopic force response toward detailed structural and dynamic information about complex trapped entities.

Acknowledgement. The authors thank the support for this research from the funding provided by the Leverhulme Trust.

Disclosures. The authors declare no conflict of interest.

References

- 1 C. J. Bustamante, Y. R. Chemla, S. Liu, and M. D. Wang, “Optical tweezers in single-molecule biophysics,” *Nature Reviews Methods Primers*, vol. 1, Mar 2021.
- 2 Z.-q. Yin, T. Li, X. Zhang, and L. Duan, “Large quantum superpositions of a levitated nanodiamond through spin-optomechanical coupling,” *Physical Review A*, vol. 88, no. 3, p. 033614, 2013.
- 3 Y. Arita, S. H. Simpson, P. Zemánek, and K. Dholakia, “Coherent oscillations of a levitated birefringent microsphere in vacuum driven by nonconservative rotation-translation coupling,” *Science advances*, vol. 6, no. 23, p. eaaz9858, 2020.
- 4 M. M. Burns, J.-M. Fournier, and J. A. Golovchenko, “Optical matter: crystallization and binding in intense optical fields,” *Science*, vol. 249, no. 4970, pp. 749–754, 1990.
- 5 P. Gupta and S. Ahlawat, “Raman spectroscopic studies on optically trapped red blood cells,” in *International Conference on Fibre Optics and Photonics*, pp. S3D–2, Optica Publishing Group, 2014.

- 391 6 L. Friedrich and A. Rohrbach, "Tuning the detection sensitivity: a model for axial
392 backfocal plane interferometric tracking," *Optics letters*, vol. 37, no. 11, pp. 2109–
393 2111, 2012.
- 394 7 Y. Yifat, J. Parker, T.-S. Deng, S. K. Gray, S. A. Rice, and N. F. Scherer, "Facile
395 measurement of the rotation of a single optically trapped nanoparticle using the
396 diagonal ratio of a quadrant photodiode," *ACS Photonics*, vol. 8, no. 11, pp. 3162–
397 3172, 2021.
- 398 8 I. A. Carvalho, N. A. Silva, C. C. Rosa, L. C. Coelho, and P. A. Jorge, "Particle
399 classification through the analysis of the forward scattered signal in optical tweezers,"
400 *Sensors*, vol. 21, no. 18, p. 6181, 2021.
- 401 9 D. Watson, N. Hagen, J. Diver, P. Marchand, and M. Chachisvilis, "Elastic light
402 scattering from single cells: orientational dynamics in optical trap," *Biophysical*
403 *journal*, vol. 87, no. 2, pp. 1298–1306, 2004.
- 404 10 R. Bar-Ziv, A. Meller, T. Tlusty, E. Moses, J. Stavans, and S. A. Safran, "Localized
405 dynamic light scattering: Probing single particle dynamics at the nanoscale,"
406 *Physical Review Letters*, vol. 78, p. 154–157, Jan 1997.
- 407 11 N. B. Viana, R. T. S. Freire, and O. N. Mesquita, "Dynamic light scattering from an
408 optically trapped microsphere," *Phys. Rev. E*, vol. 65, p. 041921, Apr 2002.
- 409 12 C. Hu, M. W. Chung, X. C. Shan, H. R. Abbas, and Y. Haw, "Confocal
410 three dimensional tracking of a single nanoparticle with concurrent spectroscopic
411 readouts," *Applied Physics Letters*, vol. 88, 2006.
- 412 13 A. Raudsepp, G. B. Jameson, and M. A. Williams, "Estimating orientation of
413 optically trapped, near vertical, microsphere dimers using central moments and off-
414 focus imaging," *Applied Optics*, vol. 61, no. 2, pp. 607–614, 2022.
- 415 14 D. W. Michael I. Mishchenko, Larry D. Travis, "T-matrix computations of light
416 scattering by nonspherical particles: A review," *Light scattering by Non-Spherical*
417 *Particles*, vol. 55, pp. 535–575, 1996.
- 418 15 G. C. Reythor, *Numerical methods for radiative heattransfer*. Doctoral thesis,
419 Universitat Politècnica de Catalunya, 2006.
- 420 16 W. Vigilante, O. Lopez, and J. Fung, "Brownian dynamics simulations of sphere
421 clusters in optical tweezers," *Optics Express*, vol. 28, p. 36131, Nov 2020.
- 422 17 I. C. D. Lenton, T. A. Nieminen, V. L. Y. Loke, A. B. Stilgoe, Y. Hu, G. Knöner,
423 A. M. Brańczyk, N. R. Heckenberg, and H. Rubinsztein-Dunlop, "Optical tweezers
424 toolbox." <https://github.com/ilent2/ott>, 2020.
- 425 18 A. Nir and A. Acrivos, "On the creeping motion of two arbitrary-sized touching
426 spheres in a linear shear field," *Journal of Fluid Mechanics*, vol. 59, p. 209–223, Jun
427 1973.

Table A1: Reference Orientations vector components, for $n_{refs} = 30$ *

α	$\hat{\mathbf{n}}_{\alpha, x}$	$\hat{\mathbf{n}}_{\alpha, y}$	$\hat{\mathbf{n}}_{\alpha, z}$
1	0.2958759	0.2958759	0.9082483
2	0.9082483	0.2958759	0.2958759
3	0.2958759	0.9082483	0.2958759
4	0.2958759	0.2958759	-0.9082483
5	0.9082483	0.2958759	-0.2958759
6	0.2958759	0.9082483	-0.2958759
7	0.2958759	-0.2958759	0.9082483
8	0.9082483	-0.2958759	0.2958759
9	0.2958759	-0.9082483	0.2958759
10	0.2958759	-0.2958759	-0.9082483
11	0.9082483	-0.2958759	-0.2958759
12	0.2958759	-0.9082483	-0.2958759
13	-0.2958759	0.2958759	0.9082483
14	-0.9082483	0.2958759	0.2958759
15	-0.2958759	0.9082483	0.2958759
16	-0.2958759	0.2958759	-0.9082483
17	-0.9082483	0.2958759	-0.2958759
18	-0.2958759	0.9082483	-0.2958759
19	-0.2958759	-0.2958759	0.9082483
20	-0.9082483	-0.2958759	0.2958759
21	-0.2958759	-0.9082483	0.2958759
22	-0.2958759	-0.2958759	-0.9082483
23	-0.9082483	-0.2958759	-0.2958759
24	-0.2958759	-0.9082483	-0.2958759
25	1.0000	0.0000	0.0000
26	0.0000	1.0000	0.0000
27	0.0000	0.0000	1.0000
28	-1.000	0.0000	0.0000
29	0.0000	-1.000	0.0000
30	0.0000	0.0000	-1.000

*Orientation vector points from centre of sphere 1 to centre of sphere 2.

Table A2: Raw intensities I_k^* and scaled intensities y_k

α	$I(\hat{\mathbf{n}}_\alpha, 15^\circ)$	$I(\hat{\mathbf{n}}_\alpha, 55^\circ)$	$I(\hat{\mathbf{n}}_\alpha, 90^\circ)$	$y(\hat{\mathbf{n}}_\alpha, 15^\circ)$	$y(\hat{\mathbf{n}}_\alpha, 55^\circ)$	$y(\hat{\mathbf{n}}_\alpha, 90^\circ)$
1	5.236437793	0.008879799	0.01023413	-0.566323866	-0.895169311	-0.782655503
2	9.029762808	0.014176754	0.023474524	1.604434643	-0.737872411	1.16224444
3	5.677784222	0.018003042	0.012268563	-0.313760083	-0.624248031	-0.48381477
4	4.054681384	0.008596164	0.007681417	-1.242592777	-0.903592052	-1.157627171
5	5.916429873	0.012267124	0.018806686	-0.17719333	-0.794580264	0.47657917
6	7.154962253	0.040816852	0.007678353	0.531566054	0.053224459	-1.158077234
7	4.857371303	0.057897419	0.009575087	-0.78324749	0.560444156	-0.87946335
8	9.018457316	0.061837715	0.027314068	1.597964991	0.677454091	1.726240514
9	5.001594138	0.007138369	0.009576248	-0.700714897	-0.946882332	-0.879292708
10	4.069312021	0.041444603	0.012267124	-1.234220286	0.071865963	-0.484026249
11	6.542222096	0.050631978	0.023474524	0.180920933	0.34469166	1.16224444
12	7.930714067	0.10058846	0.010228239	0.975495811	1.828185342	-0.78352083
13	5.236437793	0.008879799	0.01023413	-0.566323866	-0.895169311	-0.782655503
14	9.029762808	0.014176754	0.023474524	1.604434643	-0.737872411	1.16224444
15	5.677784222	0.018003042	0.012268563	-0.313760083	-0.624248031	-0.48381477
16	4.054681384	0.008596164	0.007681417	-1.242592777	-0.903592052	-1.157627171
17	5.916429873	0.012267124	0.018806686	-0.17719333	-0.794580264	0.47657917
18	7.154962253	0.040816852	0.007678353	0.531566054	0.053224459	-1.158077234
19	4.857371303	0.057897419	0.009575087	-0.78324749	0.560444156	-0.87946335
20	9.018457316	0.061837715	0.027314068	1.597964991	0.677454091	1.726240514
21	5.001594138	0.007138369	0.009576248	-0.700714897	-0.946882332	-0.879292708
22	4.069312021	0.041444603	0.012267124	-1.234220286	0.071865963	-0.484026249
23	6.542222096	0.050631978	0.023474524	0.180920933	0.34469166	1.16224444
24	7.930714067	0.10058846	0.010228239	0.975495811	1.828185342	-0.78352083
25	8.589197415	0.039227387	0.024433841	1.352317814	0.006024147	1.303159911
26	7.179988381	0.037734463	0.014278111	0.545887442	-0.038309302	-0.188629493
27	4.518783843	0.045969647	0.022162922	-0.977006687	0.206240387	0.9695815
28	8.589197415	0.039227387	0.024433841	1.352317814	0.006024147	1.303159911
29	4.635190763	0.153655662	0.02212562	-0.910391959	3.404054059	0.964102078
30	4.290255329	0.010364637	0.014275348	-1.107783832	-0.851075979	-0.189035408

* I_k values are calculated using MSTM package.