Theoretical comparison of real-time feedback-driven single-particle tracking techniques

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Real-time feedback-driven single-particle tracking is a technique that uses feedback control to enable single-molecule spectroscopy of freely diffusing particles in native or near-native environments. A number of different RT-FD-SPT approaches exist, and comparisons between methods based on experimental results are of limited use due to differences in samples and setups. In this study, we used statistical calculations and dynamical simulations to directly compare the performance of different methods. The methods considered were the orbital method, the Knight’s Tour (grid scan) method and MINFLUX, and we considered both fluorescence-based and interferometric scattering (iSCAT) approaches. There is a fundamental trade-off between precision and speed, with the Knight’s Tour method being able to track the fastest diffusion but with low precision, and MINFLUX being the most precise but only tracking slow diffusion. To compare iSCAT and fluorescence, different biological samples were considered, including labeled and intrinsically fluorescent samples. The success of iSCAT as compared to fluorescence is strongly dependent on the particle size and the density and photophysical properties of the fluorescent particles. Using a wavelength for iSCAT that is negligibly absorbed by the tracked particle allows an increased illumination intensity, which results in iSCAT providing better tracking for most samples. This work highlights the fundamental aspects of performance in RT-FD-SPT and should assist with the selection of an appropriate method for a particular application. The approach used can easily be extended to other RT-FD-SPT methods.
I. INTRODUCTION

Single-molecule spectroscopy (SMS) gives access to the properties and dynamics of biomolecules and other molecular or nanoscaled systems that are normally averaged out in bulk experiments. It allows the real-time detection of dynamics that are unsynchronized across separate particles, thus circumventing ensemble averaging and unveiling many hidden details in molecular processes. SMS typically measures the fluorescence emitted by a fluorescent probe or the autofluorescence of a molecule or nanoparticle and allows the spectrum, intensity, lifetime, and polarization of the fluorescence to be detected simultaneously. The use of a confocal pinhole, total internal reflection, or two-photon excitation allows a smaller observation volume to be probed, thus reducing the background noise and enabling the measurement of faint single emitters. Example applications include protein conformational dynamics, transcription in single DNA molecules, enzyme reactions, and light-harvesting complexes switching between different functional states.

During the past three decades, unprecedented advances have been made in the application of SMS to numerous types of biomolecules but its full potential remains to be developed. This is mainly due to its limitations not having been well addressed yet. One major limitation of standard approaches to SMS is its requirement to remove a biomolecule from its natural environment. Performing SMS in vivo is a highly challenging endeavor, considering the extremely high packing density of macromolecules in their native environment, which puts a significant constraint on the ultrahigh selectivity and sensitivity required for a successful single-molecule experiment. For this reason, SMS is commonly performed on a highly purified sample of molecules in an in vitro environment. Furthermore, collection of a statistically significant number of fluorescence photons requires a sufficiently long measurement time. Therefore, to suppress the diffusive motion of biomolecules, they need to be immobilized on a substrate, in an enclosing matrix, or in solution, or confined to a small subspace like a liposome or vesicle. Immobilization is commonly done by attaching a highly diluted concentration of molecules to a microscope coverslip through the use of molecular linkers, or by spin-coating the molecules in a polymer host matrix. Molecules can also be trapped in solution using physical, optical, electrokinetic, or thermodynamic principles. Immobilization or confinement of biomolecules allows the use of a tightly focused light source to probe it for extended periods of time. However, this is a highly artificial environment and
Theoretical comparison of real-time feedback-driven single-particle tracking techniques therefore strongly limits the biological relevance of these experiments. Immobilization of biomolecules may also introduce non-physiological interactions and structural distortions.

One key step closer to in vivo SMS is a technique that allows the real-time detection of freely diffusing single molecules for extended periods of time. This can be done by real-time feedback-driven single-particle tracking (RT-FD-SPT), which uses feedback control to keep a particle in the observation volume, without disturbing the particle’s local environment. RT-FD-SPT is not to be confused with conventional, image-based single particle tracking, which is also able to measure a particle’s motion but uses a wide field of view that does not allow for concurrent spectroscopy. RT-FD-SPT has, for example, been used to measure fluorescence lifetimes, spectra, and antibunching of freely diffusing molecules in live cells. RT-FD-SPT also has better time resolution and a much longer tracking range than conventional image-based SPT.

Despite a rather slow development of RT-FD-SPT methods during the past ca. two decades compared to other single-molecule techniques, there are already many different RT-FD-SPT methods in use, and accurately evaluating the performance of different methods is key to choosing the right method for a specific application. Comparisons can be made using experimental results, but this is not ideal due to considerable variation in experimental setups and samples. Some methods have also been investigated theoretically, using either statistical analysis or dynamical simulations but this has so far only been done for a limited number of methods and using approaches or criteria that do not allow a direct comparison of all RT-FD-SPT methods. We started to address this shortcoming in a previous study, wherein a comparison of different RT-FD-SPT methods was made using statistical analysis. In this work, we expand that study to more thoroughly investigate three commonly used RT-FD-SPT methods and extend the comparison using a dynamical simulation. We consider both fluorescence emission and scattering as photon sources. Interferometric scattering (iSCAT) is considered for the latter. Due to the limited theoretical development for iSCAT-based tracking methods we first derived an expression for its Cramér-Rao bound before evaluating its performance. Overall, our study provides a direct comparison of the performance of different RT-FD-SPT methods. It also illustrates some fundamental aspects of RT-FD-SPT performance that are often overlooked in experimental literature or studies purely focused on localization (e.g., Ref. 27), such as an important trade-off in RT-FD-SPT between tracking precision and tracking speed, and the
Theoretical comparison of real-time feedback-driven single-particle tracking techniques relationship between localization precision and realized tracking error, specifically in the context of Kalman filtering.

In Section II we describe the different position sensing methods that were investigated and derive an expression for the iSCAT Cramér-Rao bound. Section III shows the results of the statistical analysis and dynamical simulations, and Section IV provides a discussion of the results as well as concluding remarks.

II. POSITION SENSING

RT-FD-SPT is based around a feedback system that uses a measured particle position to continually re-center the particle in the observation volume. Such a feedback system consists of a “position sensor”, a control system, and an output actuator. Approaches differ mainly in how the particle position is measured. In this study, we focus on single-detector methods that can be used in two dimensions. This is done for simplicity and due to the fact that, while iSCAT is capable of 3D tracking when the motion is sufficiently slow, large axial motion would not easily be tracked in an RT-FD-SPT context. The three methods considered are also well-suited to illustrate the fundamental aspects of performance that this study aims to highlight. The analysis can, however, easily be extended to three dimensions or other position sensing methods. The 2D results are immediately applicable to scenarios such as diffusion in flat lipid bilayers.

Three position-sensing methods commonly used in two dimensions are the orbital, Knight’s Tour, and MINFLUX methods. The former two involve a beam scanning pattern over a semi-continuous area, while MINFLUX involves a “constellation” of a small number of points. All three methods have been extended to three dimensions. These methods have thus far been applied using fluorescence measurements, but we will also consider them in the context of the technique called interferometric scattering (iSCAT), which uses scattered light.

A. Scanning patterns

In the orbital method, a Gaussian laser spot is scanned in a circle around the particle of interest (Fig. 1a). If the particle is not in the center and the orbital frequency is sufficiently
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**FIG. 1:** (a) Orbital scanning pattern with beam waist $w$ and scanning length $L$. (b) Knight’s Tour scanning pattern showing the scanning length $L$ and beam waist $w$. (c) MINFLUX scanning pattern, showing scanning length $L$ and beam size parameter fwhm. The particle is kept in the center of the scanning pattern in each case.

The detected intensity varies sinusoidally, enabling one to use the amplitude and phase of this variation to determine the particle position in the $x$–$y$ plane. For sufficiently large values of the orbital radius $R = L/2$, where $L$ denotes the orbital diameter or “scanning length” (see Fig. 1), the average detected intensity has a local minimum when the particle is in the center of the orbital. For small $R$, there is a maximum average intensity in the center. A value of $R = w/\sqrt{2} \approx 0.7w$, where $w$ is the beam waist, is the largest value of $R$ for which there is a maximum average intensity in the center, and this has been shown to be the optimal geometry to maximize position information.

The Knight’s Tour method also uses a Gaussian laser spot, but instead of scanning in a circle, the spot is scanned across a grid of points by means of a “knight’s tour” (Fig. 1b). This is a heuristic to cycle through the points of the grid in a way that continually samples the entire area, rather than from one side to the other as in a raster scan. This allows feedback to be applied on every photon detection without the need to wait for a scan to complete, which would improve the tracking of fast-moving particles that might move across the scan area faster than a raster scan does. The Knight’s Tour method was originally developed as a way of increasing the size of the scan area and thus increasing the speed at which particles can be tracked. Before its implementation, a larger scan area was obtained by increasing the beam waist with the orbital method, which decreases the tracking precision. In this work, we use the knight’s tour pattern from the first work by Wang and Moerner. Different scan patterns have been used, and it has also been extended to three dimensions by the
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Welsher group in a method called 3D-DyPLoT\textsuperscript{10,17,40}.

MINFLUX utilizes an illumination minimum – typically using a doughnut-shaped laser beam spot – to measure the particle position (Fig. [1]). Keeping the particle in the center of the illumination minimum minimizes the total number of detected photons, enabling more position information to be encoded, in principle, in each detected photon\textsuperscript{23,45}. This concept is in contrast to the other two methods, where the number of detected photons is maximized. The standard two-dimensional MINFLUX approach utilizes four scan points and the original implementation involved fast scanning of the beam. A recently developed version, called p-MINFLUX\textsuperscript{46}, uses interleaved laser pulses, which eliminates the need for fast beam scanning but, instead, requires a fast pulsed laser and time-gating electronics.

B. iSCAT

RT-FD-SPT has traditionally been based on fluorescence, but it is also possible to make use of scattered light by using the technique called interferometric scattering (iSCAT). This technique uses the interference of scattered and reflected light, which, for small particles, provides a significantly stronger signal than the pure scattering signal. The principle is as follows\textsuperscript{41}, an incident light field with electric field amplitude $E_i$ on a particle close to an interface (for example the glass-water interface of a coverslip close to the particle) produces a scattered field with amplitude $E_s$ from the particle as well as a reference field $E_r$ from the reflection at the interface. These two fields interfere, giving rise to a detected intensity $I_{det}$ given by

$$I_{det} \propto |rE_i + sE_i|^2 = |E_i|^2 (r^2 + 2|r||s| \cos \theta + |s|^2),$$

where $r$ is the reflectivity of the interface, $s$ is the complex scattering coefficient, and $\theta$ is the difference in phase between the reflected and scattered fields. If the illumination originates from a laser beam with cross-sectional area $A$, $s$ can be determined from the scattering cross-section $\sigma_{scat}$ through the relation $|s|^2 = \sigma_{scat}/A$.

Due to the square on $|s|$, for a small particle the pure scattering term, $|E_i|^2|s|^2$, is negligible compared to the interference term, $|E_i|^2r||s| \cos \theta$. Even though the detected signal is dominated by the reference term, $|E_i|^2r^2$, the latter should remain constant and can be estimated by measuring the background $N_{bkg}$ when there is no particle in the observation.
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volume. The actual iSCAT measurement is then made in terms of the contrast $C_f$ between the interference and the background, defined as

$$C_f = \left| \frac{N_{\text{det}} - N_{\text{bkg}}}{N_{\text{bkg}}} \right|,$$

where $N_{\text{det}}$ is the number of photons detected during the measurement.

iSCAT essentially provides only a different source of photons and the actual position measurement can thus still be done using a variety of methods. Traditionally, the method has used a camera but it has recently also been implemented using a knight’s tour scan and real-time feedback. The combination of iSCAT with RT-FD-SPT is a promising but as yet untapped area of development.

C. Static localization

A particle that moves slowly in comparison with the feedback bandwidth can be tracked well enough to be considered approximately static relative to the tracking beam. For such a tracking situation it is useful to determine the accuracy with which a static particle’s position can be determined. The main limitations on the accuracy of any particular method are the number of detected photons and the signal-to-background ratio (SBR). These two factors are usually strongly related but do not necessarily show a one-to-one relationship. We therefore develop a framework to compare the precision of the different position sensing methods given a certain number of detected photons and a certain SBR. To this end, we use the Fisher information, which provides a measure of the amount of information that an observed random variable carries about an unknown parameter. In our case, the observed variable is the photon counts and the unknown parameter is the particle position. We are ultimately interested in the Cramér-Rao bound (CRB), which can be calculated from the Fisher information. The CRB is a lower bound on the variance of any position estimator, and thus a direct measure of precision. For fluorescence-based localization, the CRB calculation is described in Ref. 24.

For iSCAT, no CRB has as yet been determined to the best of our knowledge. The major difference between the photon statistics of iSCAT and fluorescence is that for iSCAT, the statistics are determined by the background. This is because the interference counts are not measured directly, but measurements are made in terms of the contrast $C_f$ between the interference and background counts. This contrast measurement is dominated by the reference
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(background) counts, and the uncertainty in the estimate of the interference counts is therefore mostly determined by the photon noise of the reference counts. Since the reference term is produced purely by the laser, we can assume that the reference counts \( N_r \approx N_{bkg} \) are Poisson distributed. We further assume that \( N_r \) is large, and it can thus be well approximated by a Gaussian distribution with variance \( N_r \). This leads to the estimate of the interference counts \( N_{int} \approx N_{det} - N_{bkg} \) having a variance of \( \sigma_N^2 = \sigma_{det}^2 + \sigma_{bkg}^2 \approx 2 \sigma_{bkg}^2 \approx 2N_r = 2N_{int}/C_f \). Note that this variance is different from the Poisson variance \( \sigma_{N,Poisson}^2 = N_{int} \) that would be applicable if it was possible to measure the interference counts directly. As as \( C_f \) is typically less than 5%, the variance \( \sigma_N^2 \) is large compared to the mean. We derive an expression for the iSCAT CRB using this variance \( \sigma_N^2 \).

We follow a modified approach of Ref. [24] and consider a particle at a position \( \vec{r}_m \in \mathbb{R}^d \) probed with an illumination pattern comprising \( K \) distinct intensities \( \{I_0(\vec{r}), \ldots, I_{K-1}(\vec{r})\} \), with a resulting collection \( \vec{n} = \{n_0, n_1, \ldots, n_{K-1}\} \) interference photons per pattern point and a total number of interference photons \( N_{int} = n_0 + n_1 + \ldots + n_{K-1} \).

Assuming \( n_i \) to be sufficiently large, \( \vec{n} \) is approximately Gaussian with mean \( \hat{N}_{int} \) and standard deviation \( \sigma_N \). We can approximate \( n_i \) as also being Gaussian with expected values \( p_i \hat{N}_{int} \) and standard deviations \( p_i \sigma_N \), where

\[
p_i = \frac{I_i}{\sum_{j=0}^{K-1} I_j},
\]

with \( 0 \leq i \leq K - 1 \).

Thus the probabilities for measuring \( \vec{n} \) conditioned to \( \hat{N}_{int} \) are given by

\[
P(\vec{n}|\hat{N}_{int}) = \prod_{i=0}^{K-1} \frac{\exp -\frac{1}{2} \left( \frac{n_i - \hat{N}_{int} p_i}{\sigma_N p_i} \right)^2}{\sigma_N p_i \sqrt{2\pi}},
\]

where \( p_{K-1} = 1 - \sum_{j=0}^{K-2} p_j \). The Fisher information is therefore given by

\[
\{F_{\vec{n}}\}_{ij} = E \left( -\frac{\partial^2}{\partial p_i \partial p_j} \ln P(\vec{n}|\hat{N}_{int}) \right) \text{ with } i, j \in [0, \ldots, K - 2]
\]

\[
= \frac{\hat{N}_{int}^2}{\sigma_N^2} \left( \frac{1}{p_{K-1}^2} + \delta_{ij} \frac{1}{p_i^2} \right) \text{ with } i, j \in [0, \ldots, K - 2],
\]

which differs from the expression for fluorescence in the \( N \)-dependent factor as well as the square on \( p_i \). The result in terms of position, \( F_{\vec{r}_m} \), is given by

\[
F_{\vec{r}_m} = \mathcal{J}^\dagger F_{\vec{n}} \mathcal{J}^*,
\]
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with $\mathbf{J}^* \in \mathbb{R}^{(K-1) \times d}$ the Jacobian matrix of the transformation from $\bar{r}$-space to the reduced $\bar{p}$-space, which is a $(K-1)$-dimensional space since $p_{K-1} = 1 - \sum_{j=0}^{K-2} p_j$. There are, therefore, $K - 1$ independent parameters. We finally get

$$F_{\bar{r}_m} = \frac{N_{int}^2}{\sigma_N^2} \sum_{i=0}^{K-1} \frac{1}{p_i^2} \left[ \left( \frac{\partial p_i}{\partial r_{m1}} \right)^2 \ldots \left( \frac{\partial p_i}{\partial r_{md}} \right)^2 \right], \quad (8)$$

The Cramér-Rao lower bound on the covariance matrix $\Sigma(\bar{r}_m)$ is given by $\Sigma_{CRB} = F_{\bar{r}_m}^{-1}$. In our results, we use the arithmetic mean of the eigenvalues of this matrix (recall that $d$ is the number of dimensions):

$$\tilde{\sigma}_{CRB} = \sqrt{\frac{1}{d} \text{tr}(\Sigma_{CRB})}. \quad (9)$$

III. RESULTS

A. Static localization

1. Fluorescence

We first compare the three different scan patterns based on fluorescence using $N = 100$ photons at two different signal-to-background ratio (SBR) values and different values of the scanning length $L$. For the orbital and Knight’s Tour methods, the beam waist was scaled relative to $L$. For the orbital method, the relation $L = 2w/\sqrt{2}$ was used, while for the Knight’s Tour method, the grid spacing was chosen such that $L = 3.75w$, consistent with Ref. [32]. Therefore, for these two methods, $L$ is bounded by the diffraction limit, whereas for MINFLUX it is not. The scaling of the beam size with scanning length for the orbital and Knight’s Tour methods also means that the SBR does not depend on the scanning length for these methods (if the beam size is kept constant, an increase in the scanning length would lead to a decrease in the SBR). For MINFLUX, however, there is a dependence of SBR on scanning length; specifically, the SBR scales with the average illumination intensity and thus increases with increased $L$. In this case, the assumed SBR of 10 was for $L = 100$ nm, and the adjusted SBR for other scanning lengths ranged from 2.5 for $L = 50$ nm to 300 for $L = 700$ nm. In practice, the SBR would be bounded by saturation, but since MINFLUX does not show good performance at large $L$, these results are not of practical relevance.
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Furthermore, at these large $L$ values, the results for different values of the SBR $\geq 10$ are very similar to each other and the results in Fig. 3 are therefore representative of the performance. The results are shown in Fig. 2.

All three methods show improved precision with decreased $L$, with the lower SBR showing a general decrease in precision. The differences between the three methods are also clear. The orbital method (Figs. 2a and 2b) shows a lower CRB in the center of the scan range, along with a moderate decrease in CRB with decreased $L$. The Knight’s Tour method (Figs. 2c and 2d) has a very even CRB across the scan range due to even illumination. It too has a moderate improvement in precision with decreased $L$. MINFLUX (Figs. 2e and 2f) shows a dramatically lower CRB in the center of the scan range compared to the edges, along with a large decrease in the central CRB with decreased $L$. Here we also see the greatest impact of the SBR: in the case of SBR $= \infty$ (Fig. 2e), the central CRB decreases without bound as $L$ tends to 0, whereas with a finite SBR (Fig. 2f) there is a limit to how low the central CRB can be made by decreasing $L$. The three methods are compared directly in Figs. 2g and 2h where the beam waist for the orbital and Knight’s Tour methods is set to $w = 400$ nm, and two different MINFLUX $L$-values are shown. It is clear that MINFLUX indeed provides more information per photon, but the trade-off is a smaller scanning range. With the same value of $L$, the performance of MINFLUX is somewhat worse than that of the orbital method. This fact is important, as a large scanning range is required for tracking fast-moving particles.

2. iSCAT

We next compare the different scan patterns using iSCAT-based localization, considering $N_{int} = 100$ and two contrast values: $C_f = 5\%$ and $C_f = 1\%$ (Fig. 3). The shape of the CRB curve for the orbital method (Figs. 3a and 3b) is quite different than with fluorescence. As we have previously shown, this is due to the uncertainty in the parameter $p_i$ (Eq. 3) having a maximum in the center, both for iSCAT and fluorescence. For fluorescence, this uncertainty is counteracted by the dependence of the position uncertainty on the derivative of $p_i$, which has a minimum in the center. For iSCAT, the dependence on the parameter uncertainty itself is quadratic and therefore weighs more strongly. The CRB curve for the Knight’s Tour method (Figs. 3c and 3d) is again very flat. MINFLUX (Figs. 3e and 3f) shows
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FIG. 2: CRB along the x-axis for different $L$-values (indicated in nm next to each curve) for (a, b) orbital method, (c, d) Knight’s Tour method, (e, f) MINFLUX, (g, h) different methods (MF - MINFLUX, KT - Knight’s Tour, Orb - orbital method). $N = 100$ photons for all curves. Left column: $SBR = \infty$; right column: $SBR = 10$. For MINFLUX, $SBR = 10$ corresponds with $L = 100$ nm, while SBR was scaled for other $L$-values (see text for details).
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a similar trend as with fluorescence, with the central precision again depending strongly on the scanning length, resulting in a good precision only in the center.

\[ C_f = 5\% \quad \text{Orbital method} \quad C_f = 1\% \]

\[ \text{Knight's Tour method} \]

\[ \text{MINFLUX} \]

\[ \text{FIG. 3: CRB along the x-axis for different } L \text{-values (indicated in nm next to each curve) for (a, b) orbital method, (c, d) Knight’s Tour method and (e, f) MINFLUX with iSCAT localization. } N_{int} = 100 \text{ photons, left column: } C_f = 5\%, \text{ right column: } C_f = 1\%. \]

An interesting feature of the iSCAT CRB expression is its dependence on the number of scan points (Fig. 3). The improvement in precision when using more scan points is due to a more accurate measurement of the background when it is measured at every point.
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For all curves, $L = 500$ nm, $N_{int} = 100$ photons, $C_f = 1\%$.

In order to compare iSCAT and fluorescence, we consider the relative number of fluorescence and scattering counts at the same illumination intensity, as well as the average iSCAT contrast, for a specific sample. The average contrast is proportional to $2\sqrt{\sigma_{scat}/r}$, where $\sigma_{scat}$ is the scattering cross-section and $r$ is the reflectivity of the interface, while the relative fraction of iSCAT counts to fluorescence counts, $N_{scat}/N$, is proportional to $2\sqrt{\sigma_{scat}/\sigma_{abs}\Phi_f}$, where $\sigma_{abs}$ and $\Phi_f$ are the absorption cross-section and fluorescence quantum yield, respectively. We consider four example samples that may be tracked using both methods: an HIV virion labeled with the quantum dot QD525, the eGFP mutant of green fluorescent protein (a mutant featuring enhanced fluorescence), the main light-harvesting complex of plants (LHCII), and phycobilisome (PB), the main light-harvesting from the cyanobacterial strain *Synechocystis* PCC6803. These examples include intrinsically fluorescent (eGFP, LHCII and PB) and artificially labeled (QD525-labeled HIV) nanoscaled biological systems displaying a wide range of sizes. GFP is a relatively small protein, while PB is a very large (6.2 MDa) multidomain pigment-protein complex; LHCII is a trimeric pigment-protein complex with a molecular weight of 128 kDa, while HIV is an order of magnitude larger in all dimensions. Furthermore, GFP and PB are both water-soluble, while LHCII is a membrane protein and typically embedded in detergent micelles for *in vitro* studies.

The scattering cross-section of a particle can be estimated using the formula $\sigma_{scat} = \frac{8\pi^3\alpha^2(\lambda/n_m)^{-4}}{3}$, where $\alpha = 3V(n_s^2 - n_m^2)/(n_s^2 + 2n_m^2)$ is the particle’s polarizability, $V$ is its volume, $n_s$ its refractive index, $n_m$ the refractive index of the surrounding medium, and $\lambda$ the wavelength of the incident light. The calculated values for $\sigma_{scat}$, $C_f$, and $N_{scat}/N$ are
displayed in Table S1, along with other relevant parameters derived from literature. $C_f$ is calculated from Eq. (2). We assume HIV to be spherical with a radius of 73 nm. PB is hemidiscoidal with a radius of 30 nm and a thickness of 10 nm. LHCII is cylindrical in shape with a radius of 3.65 nm and a height of 4.5 nm. GFP is also cylindrical with a radius of 1.2 nm and a height of 4.2 nm. We also consider the case of LHCII in a detergent micelle, as this is typically how it is measured in solution. In this case, the length of an n-dodecyl-β-D-maltoside molecule (2.3 nm) was added to the effective radius. In each case, the effective radius is given in Table S1, equal to the radius of a sphere with the same volume as the particle. The peak absorption wavelength (except for QD525 for which the absorbance continues to increase at shorter wavelengths) of each sample is also shown in Table S1. The reflectivity of a glass-water interface at normal incidence (0.4%) for a typical microscope coverslip was used for the calculations.

The CRB as a function of the detected number of photons is shown in Fig. 5 for iSCAT and fluorescence. The iSCAT performance follows the size of the particle. For a large particle such as a virus, iSCAT is orders of magnitude more precise than fluorescence, whereas for small particles such as GFP or LHCII, fluorescence-based localization is better. For LHCII in a micelle and PB, the precision of the two methods is similar. The asymptotic shape of the iSCAT curve is due to the factor $\hat{N}^2 - \sigma_N^2 \sigma_H^2$ in the Fisher information expression (Eq. 8). The corresponding factor for fluorescence is simply $\hat{N}^4$. For large values of $\hat{N}$, $\hat{N}^2 - \sigma_N^2 \sigma_H^2$ is approximately equal to $\hat{N}^2 \sigma_N^2$. If we assume $\hat{N}$ to be large and Poisson-distributed, then $\sigma_N^2 \approx \hat{N}$ and therefore $\hat{N}^2 / \sigma_N^2 \approx \hat{N}$. It follows that for both iSCAT and fluorescence, at high count rates, $\sigma_{CRB}^2 \propto \hat{N}^{-1/2}$ (Eq. 9).

The result in Fig. 5a assumes the same illumination intensity for iSCAT and fluorescence. However, iSCAT can be performed at a wavelength where the sample does not strongly absorb, and thus it is possible to use a much higher illumination intensity for iSCAT at a wavelength where phototoxicity is strongly limited, leading to improved precision. To investigate this possibility, we chose a separate wavelength for iSCAT for each sample, and assumed a 1000 times increased illumination intensity. This is based on single-molecule fluorescence experiments typically using an excitation power of around 1 μW, and iSCAT experiments using around 1 mW. The calculation values in this case are shown in Table S2 and the results are shown in Fig. 5b. In this case, both PB and LHCII in a micelle have a markedly improved precision with iSCAT, and for pure LHCII and GFP the precision is
FIG. 5: CRB as a function of the number of photons for both iSCAT and fluorescence for different samples, using the same illumination intensity for a and a 1000 times increased intensity for iSCAT for b. The upper x-axes show fluorescence counts. The iSCAT counts are higher than the fluorescence counts by the ratios given in ?? for a and b, respectively. The average iSCAT counts for all samples except HIV-QD are shown on the lower x-axes. For HIV-QD, the iSCAT counts are $\sim 10^4$ times higher than the average for the other samples, as indicated next to the curve.

roughly the same for the two methods.

The relationship between scattering and absorption cross-sections and the iSCAT and fluorescence CRBs is further illustrated in Fig. 6. Here the logarithm of the ratio of iSCAT CRB to fluorescence CRB is depicted as a function of the scattering cross-section and the product of the absorption cross-section with the fluorescence quantum yield. The examples considered above are indicated, along with a number of other representative samples: a 12-nm diameter CdSe/ZnS quantum dot (QD), a 40-nm fluorescent microsphere, fibrinogen labeled with Alexa Fluor 647, and the bacterial light harvesting complex LH2. These samples
Theoretical comparison of real-time feedback-driven single-particle tracking techniques all show improved precision with iSCAT, to varying degrees. It is a simple task to estimate the position of any other sample, and thus evaluate the possible suitability of iSCAT for that sample, using this figure.

![Graph showing the logarithm of the ratio of iSCAT CRB to fluorescence CRB as a function of the scattering cross-section and the product of absorption cross-section with fluorescence quantum yield.](image)

**FIG. 6**: Logarithm of the ratio of iSCAT CRB to fluorescence CRB as a function of the scattering cross-section and the product of absorption cross-section with fluorescence quantum yield. $N = 1 \times 10^4$ fluorescence photons and the iSCAT illumination is assumed to be 1000 times higher than the fluorescence illumination (each sample has a different assumed illumination that would produce $1 \times 10^4$ fluorescence photons). Positive (blue) values indicate better precision with iSCAT while negative (red) values indicate better precision with fluorescence. The positions of the example samples are indicated. The lower right region is excluded due to undefined values (corresponding to low numbers of photons in Fig. 5).

**B. Dynamic simulation**

1. **Control System Design**

   To compare the methods' dynamic performance, the complete control system was simulated numerically. The output is assumed to be a piezoelectric sample stage, which is assumed to have a linear response, a general feature of piezoelectrics. For simplicity we
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design a second-order system with a step response similar to that of a real stage (with its manufacturer-provided controller) (??). Details can be found in the SI; we show the final result here. With the system state

$$\mathbf{x} = \begin{bmatrix} x_p \\ x_s \\ \dot{x}_s \end{bmatrix},$$

where $x_p$ and $x_s$ are the particle and stage positions, respectively, the dynamical system is given by

$$\dot{\mathbf{x}} = \begin{bmatrix} 0 & 0 & 0 \\ 0 & -2 & 4 \\ 0 & -4 & 0 \end{bmatrix} \mathbf{x} + \begin{bmatrix} 0 \\ 2 \\ 4 \end{bmatrix} u + \begin{bmatrix} \sqrt{2D} \\ 0 \\ 0 \end{bmatrix} \xi(t)$$

(11)

$$y = \begin{bmatrix} 1 & -1 & 0 \end{bmatrix} \mathbf{x} + w_n,$$

(12)

where $u$ is the control input, $D$ is the two-dimensional diffusion coefficient, $\xi$ is a white-noise process with mean 0 and variance 1, $w_n$ is an error term that mostly originates from Poisson noise propagating through the measurement process, and $y$ is the measured position. We also implemented a Kalman filter as this is commonly used in real RT-FD-SPT systems.

For the orbital method, we used the position estimate from Ref. 28, for MINFLUX, we used the online position estimator from Ref. 24, and for the Knight’s Tour method, we used the estimator

$$\hat{\mathbf{r}} = \sum_{i=1}^{40} \hat{p}_i \cdot \bar{r}_{b_i},$$

(13)

where $\hat{\mathbf{r}}$ is the estimated position vector, $\bar{r}_{b_i}$ are the position vectors of the beam positions $b_i$, and $\hat{p}_i$ are the estimated values of the parameters defined in Eq. 3, based on the detected photons at each scan point.

We performed dynamic simulations of each method by numerically integrating Eq. 12 in two dimensions, computing estimates for $\mathbf{y}_k$ using the estimators mentioned above, and subsequently estimating $\mathbf{x}_k$ using the Kalman filter equations ($\mathbf{y}_k$ and $\mathbf{x}_k$ are the discretization of $\mathbf{y} = y$ and $\mathbf{x}$ respectively — see the SI for details). The estimated position $x_p$ was directly applied as feedback into the control input $u$.

The parameters used were as follows: the beam waist for the orbital and Knight’s Tour methods was $w = 400$ nm, the MINFLUX beam size parameter was fwhm = 500 nm, the
MINFLUX scanning length was $L = 50 \text{ nm}$, the detected count rate for a particle in the center of the beam was $\Gamma_0 = 12.5 \ \text{kcounts/s}$, and the fluorescence signal-to-background ratio was 10. For iSCAT-based localization, the same assumptions were made for the average contrast and ratio of iSCAT to fluorescence counts as in the case of static precision. The beam scan was performed at 8.33 kHz, and feedback was applied at the same frequency. The Kalman noise covariance $R$ was adjusted empirically for each scan pattern and iSCAT sample, to achieve the fastest possible tracking.

A typical well-tracked trajectory is shown in the left column of Fig. 7, where the diffusion coefficient is $0.1 \ \mu\text{m}^2\text{s}^{-1}$. The stage closely follows the diffusion and the intensity shows only Poisson noise. An “almost tracked” trajectory is shown in the right column of Fig. 7, where the diffusion coefficient is $20 \ \mu\text{m}^2\text{s}^{-1}$. Here, the system is unable to respond fast enough. There are large jumps in the measured intensity as the particle moves in and out of the illumination beam before eventually being lost completely. Performing many such simulation runs for different values of $D$, we can calculate the average tracking error, and the results are shown in Fig. 8.

We first compare the different scanning patterns using fluorescence as shown in Fig. 8a. For each method, three different regions can be identified. At high values of $D$, there is an “untracked” region, where the tracking error follows free particle statistics, i.e., $e_{\tau_0} = \sqrt{2D\tau}$, indicated by the dashed line. At intermediate values of $D$, there is a “well-tracked” region, where the error is lower than the free particle value. At the lowest values of $D$, the system starts tracking noise and the error is worse than in the case when no tracking takes place. Another way of looking at the latter regime is to say that the feedback frequency is too fast.

The scanning methods give rise to at least two important differences. Firstly, in the well-tracked region, the precision of the tracking error lines up with the static result (Fig. 2), with MINFLUX having the best precision, and the Knight’s Tour method having the worst precision. Specifically, the precision in the center of the tracking region determines the tracking error, since good tracking keeps the particle close to the center. Secondly, each method has a different “cutoff” value where the diffusion becomes too fast to track. This value is determined by the size of the tracking area and also confirms the static result. MINFLUX has the lowest cutoff due to its small tracking range ($L = 50 \text{ nm}$), followed by the orbital ($L = 566 \text{ nm}$) and Knight’s Tour ($L = 1500 \text{ nm}$) methods. For very slow diffusion,
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it is clear that the precision does not follow the same trend as in the faster tracked region. This is due to different values being used for the Kalman covariance $R$ for the different scanning patterns. These values were optimized to achieve the highest possible “cutoff” value for each method, and the Knight’s Tour has the largest value. As a result, the effect of noise is dampened, artificially providing better performance in this “noise-tracking” regime. In a real experiment, one would simply decrease the feedback frequency for tracking slower diffusion.

A comparison between iSCAT and fluorescence for the different example samples is shown in Figs. 8c and 8d. For simplicity we assume a constant fluorescence intensity for all the samples, since we are only interested in the relative performance of iSCAT vs fluorescence-based localization. Hence, there is only one curve for fluorescence. The general trend is the same as in Fig. 5 and the main result of static precision is confirmed: for all samples except GFP, iSCAT performs better than fluorescence when an increased illumination intensity is
FIG. 8: Average tracking error as a function of the diffusion coefficient for (a,b) the three different scanning patterns using fluorescence-based localization (a) with and (b) without a Kalman filter, and (c,d) different samples using iSCAT and fluorescence with (c) the same illumination intensity and (d) a 1000 increased illumination intensity for iSCAT. A constant fluorescence intensity was assumed for all samples, justifying why (c) and (d) show only one curve for fluorescence. For (a), (b), and (d), an average over 5 runs was used for each value of the diffusion coefficient, while for (c) the average was over 100 runs due to the large variance in the average error of the unsuccessfully tracked samples. The dashed lines labeled “untracked” represent the expected error for a freely diffusing particle.

used. Interestingly, the improvement over fluorescence is even better here, with even LHCII showing markedly better tracking using iSCAT.

One noticeable difference between the dynamic tracking precision and static CRB is the smaller difference in precision between different methods and samples in the case of dynamic
tracking. This is mostly due to the Kalman filter successfully suppressing the influence of noise. In Fig. 8b, the tracking error is shown for the different scanning patterns using a simulation without a Kalman filter. To amplify the results, a higher fluorescence intensity (60 kcounts/s) was used. Here, the tracking error matches the CRB more closely. Comparing the result with and without the Kalman filter highlights the power of the Kalman filter to compensate for noise. However, there are still fundamental limits to precision that the Kalman filter is unable to fully overcome.

IV. DISCUSSION AND CONCLUSION

The results for the different scanning patterns highlight the fundamental trade-off between tracking speed and tracking precision. The orbital method was shown to provide optimal position information when using a Gaussian beam\(^{23}\). The success of MINFLUX demonstrates, however, that much better precision can be achieved with a non-Gaussian beam profile — specifically, one with an illumination minimum. The Knight’s Tour method, on the other hand, while having worse precision than the orbital method, is much faster due to a much larger scan area. The Knight’s Tour method (or at least a three-dimensional version of it, 3D-DyPLoT) has also shown better experimental results than the orbital method in terms of RT-FD-SPT. While both methods have been shown experimentally to track particles up to \(D = 20 \mu \text{m}^2 \text{s}^{-1}\) (Ref. 40 and 52), 3D-DyPLoT has achieved this at a 6 times lower count rate (20 kHz vs 120 kHz\(^{40,52}\)).

It has recently been shown theoretically and experimentally that selectively sampling only some locations in the Knight’s Tour pattern results in improved precision\(^{26}\). Sampling only at the most information dense locations does, however, decrease the effective scanning area. MINFLUX provides a marked improvement in precision over Gaussian-beam-based localization, but it has not been proven to provide the best possible precision. Similarly, the Knight’s Tour provides a better way of achieving fast tracking than expanding the beam size and using the orbital method, but is certainly not optimal in terms of precision. Gallatin and Berglund\(^{23}\) answered the question of what the best scan path is for optimal precision given a Gaussian beam of a certain size. In fact, all of the work on this topic can be thought of as trying to answer the same question: What is the optimal beam shape and scan pattern for a certain size scan area? In the case of MINFLUX, a simpler version of this problem is
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considered where the size of the scan area is allowed to be arbitrarily small. Gallatin and Berglund likewise simplified the question by assuming a fixed beam shape. Theirs remain the only conclusive answer to a version of the problem. An alternative approach is not to have a fixed scan area at all but to optimize the scanning pattern in real time. Vickers and Andersson showed that this can be done using a control law that optimizes the Fisher information in real time.

An important difference between the orbital method and the other methods is that the orbital method makes use of continuous scanning. With the other methods, the beam has to move between spatial points on a significantly shorter time scale than the dwell time at each point (in our simulations, this scanning time was assumed to be instantaneous). This requires much faster scanning hardware than for the orbital method, thus adding significantly to the cost of implementation.

iSCAT has thus far only been applied once with real-time feedback, in the form of an ABEL trap. In this work, we have shown that the method should outperform fluorescence-based localization for many samples, even ones that are intrinsically fluorescent. This finding makes it a very promising localization technique for RT-FD-SPT, which we hope will see much use in future.

We have considered here only a subset of RT-FD-SPT methods, and only in two dimensions. The approach used can easily be applied to other methods such as other scanning patterns, multi-detector methods, or extended to three dimensions. It would also be very useful for investigating different control systems — extremum seeking controllers, for example — and different estimation and filtering methods. An important aspect of performance not addressed in this work is that of application-specific performance, notably with regard to spectroscopy. As we discuss in Ref. good performance for spectroscopy generally requires an adequate tracking precision to keep the particle in the observation volume, together with a sufficient tracking duration to collect the necessary spectroscopic data. Yet, it is conceivable that an analysis of different methods could be made based on the CRB of some measured spectroscopic quantity or even a simpler parameter such as the diffusion coefficient to find the truly optimal tracking method for a specific application.

To conclude, we have compared different RT-FD-SPT methods directly, and have found that there is a fundamental trade-off between tracking speed and precision. The Knight’s Tour method is the fastest, MINFLUX is the most precise, and the orbital method is a cost-
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effective option in between. iSCAT shows improved performance over fluorescence for most
of the biological examples considered in this study when using a suitably large illumination
intensity. The results are useful for choosing between methods and as a framework for
comparing other RT-FD-SPT methods.

SUPPLEMENTARY MATERIAL

See supplementary material for calculation values of iSCAT parameters and details on
the dynamical simulations.

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DATA AVAILABILITY

The data and code for all calculations and simulations is available at https://github.
com/bvanheerden/spt_sim

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