# Quantum enhanced localization microscopy with a single photon avalanche diode array

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Conventional single molecule localization microscopy infers the locations of putatively isolated fluorescent emitters to produce super-resolved images. The number of active fluorescent emitters is therefore a critical piece of information during localization; however, this quantity is generally under-determined in widefield fluorescent microscopy. Nevertheless, many fluorophores used in bioimaging are intrinsically single photon sources which exhibit fluorescence antibunching and can emit only one photon following a very short excitation pulse. This general property may provide the necessary physical information for constrained localization in non-sparse scenes by yielding estimates of the number active fluorescent emitters. Here, we leverage recent advancements in single photon avalanche diode (SPAD) arrays to accurately count the number active emitters in a fluorescent sample, while concurrently performing intensity-based multi-emitter localization. We find that SPAD cameras, with their high temporal resolution and single photon sensitivity are capable of precise single molecule localization microscopy, while maintaining the relative simplicity of widefield imaging.

#### Introduction

Far-field optical microscopy is fundamentally limited by diffraction, with the maximum attainable resolution being limited to approximately half the wavelength of light. Several schemes to beat the diffraction limit have been developed in recent years. Many of these schemes utilize the concept of precise localization of isolated fluorescent emitters which blink over a time series of frames [1, 8]. An inherent problem with such methods is the requirement that fluorescent emitters be isolated, slowing down the acquisition of super-resolved images. To address this, we leverage the fact that many fluorophores are intrinsically single photon sources and exhibit fluorescence antibunching. This property can constrain the number of active fluorescent emitters in a region of interest (ROI) and can potentially enable localization in non-sparse scenes [6, 10].

Molecular counting with fluorescence antibunching has a fairly simple motivation: coincidence of photons at multiple detector elements during high speed imaging provides evidence for the number of single photon sources present in the imaged region. Combining the ideas of conventional super-resolution approaches with photon statistics may prove to be a powerful set of methods for bioimaging. SPAD cameras achieve orders of magnitude higher temporal resolutions than standard CMOS cameras, single photon sensitivity, and dark count rates less than 25cps. Furthermore, the reduced readout noise and large fill-factor of recently commercialized SPAD arrays suggests their use for single molecule localization with reduced localization uncertainty. Localization uncertainty, typically the root mean square error (RMSE) of a maximum likelihood or similar statistical estimator, is bounded from below by the inverse of the Fisher information matrix, known as the Cramer-Rao lower bound [2]. Managing the increase in localization uncertainty at high labeling density remains a major bottleneck to localization microscopy. For example, static uncertainty due to molecular crowding can be partially amelioriated by using pairwise or higher-order temporal correlations within a pixel neighborhood [3]. However, the number of fluorescent active emitters in a region of interest remains critical prerequisite information in single molecule localization.

In this study, we present a method for widefield single photon counting in order to rigorously count fluorophores in the sample and subsequently constrain single molecule localization. We investigate the theoretical properties of the zero-lag second-order coherence function  $g^{(2)}(0)$  for widefield photon counting and its spatial properties. Using Bayesian analysis, we derive a posterior distribution on the number of active fluorescent emitters in a region of interest. We then combined this with single molecule localization algorithms and demonstrate resolution of multiple emitters using a multi-emitter fitting algorithm and report localization errors with respect to the Cramer-Rao bound.

## Basic Scheme

We consider a simplified description of widefield photon counting for a a single photon source in the object plane labeled by a continuous-valued coordinate  $\theta = (\theta_u, \theta_v)$ . The spatial profile O of the field in image space is presumed to have a Gaussian shape [4, 7, 11].

$$O(u,v) = \frac{1}{2\pi\sigma^2} e^{-\frac{(x-\theta_u)^2 + (y-\theta_v)^2}{2\sigma^2}}$$
(1)

Therefore the field operator in object space is  $\hat{E} \propto \hat{a}$  and in image space  $\hat{E} \propto O(x,y)\hat{a}$ . Since our SPAD detectors at the image plane must be discrete, the total field at a detector element k centered in image space at  $s_k = (u_k, v_k)$  is then given by integrating over pixels of width  $\delta$ . Moreover, the Gaussian O is presumed to be isotropic and therefore we have  $\hat{E}(s_k) \propto \Gamma_u(u_k, \theta_u) \Gamma_v(v_k, \theta_v)$ . For example,

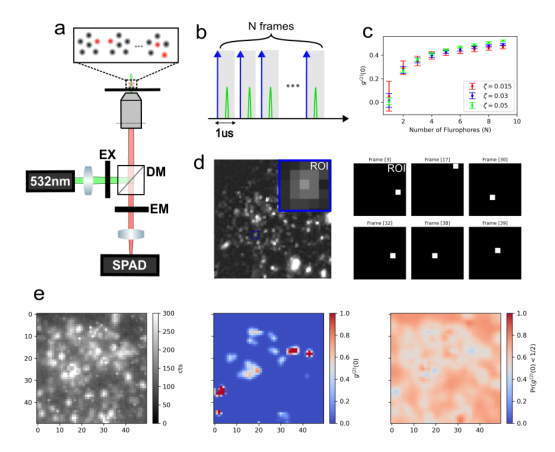


FIG. 1: Single photon counting with a SPAD array (a) Conventional widefield microscopy with integrated SPAD array (b) Single photon imaging scheme using 1us exposures containing a picosecond laser pulse (c) Sum of photon counts over a 5x5 region of interest (ROI), taken with  $N_{\text{frames}} = 5 \times 10^5$ 

$$\Gamma_u(u_k, \theta_u) = \frac{1}{\sqrt{2}} \left( \operatorname{erf} \left( \frac{u_k + \frac{1}{2} - \theta_u}{\sqrt{2}\sigma} \right) - \operatorname{erf} \left( \frac{u_k - \frac{1}{2} - \theta_u}{\sqrt{2}\sigma} \right) \right)$$

We now consider the case of pulsed excitation where the interval between pulses much longer than the fluorescence lifetime. Upon excitation of an isolated fluorescent emitter, a photon is detected at a particular detector element k with probability  $\zeta_k \propto \langle \hat{E}^\dagger(s_k) \hat{E}(s_k) \rangle = \frac{1}{2} \Gamma_v^2 \Gamma_v^2 \text{Tr}(\rho a^\dagger a)$  where  $\rho$  is the density matrix for a two-level system. Similarly, the probability of detection in a region of interest collecting all photons emitted is  $\zeta \propto \text{Tr}(\rho a^\dagger a)$ . Here, we are primarily concerned with the latter quantity, and its application in counting fluorescent emitters.

By temporarily ignoring the spatial profile described by (1), we derive a likelihood on the number of fluorophores in a small ROI with a lateral dimension d=5 pixels. For N fluorophores emitting photons which can be detected within a ROI of the SPAD array, the number of signal photons measured  $n_{\text{signal}}$  following a single excitation pulse will have Binomial statistics  $n_{\text{signal}} \sim \text{Binom}(N, \zeta)$ . Photon pile-up at a single detector ele-

ment can be safely neglected in this model due to its relatively low likelihood. We then model the background signal within the region of interest as a coherent state, which must follow Poissonian statistics  $n_{\text{background}} \sim \text{Poisson}(\lambda)$  for an expected number  $\lambda$  of background counts in the ROI per frame. The total number of counts  $n = n_{\text{signal}} + n_{\text{background}}$  detected in the region of interest following a single pulse is then distributed by the likelihood

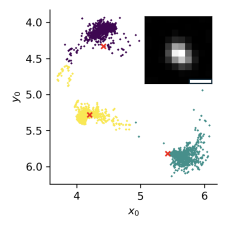
$$p(n = n' \mid N, \zeta) = \sum_{i=0}^{\infty} {N \choose i} \zeta^{i} (1 - \zeta)^{N-i} \frac{\lambda^{n'-i}}{(n'-i)!} e^{-\lambda}$$
(2)

The expression in (2) represents a convolution of Poisson and Binomial probability mass functions. This result is the primary means of inference of the number of active emitters N in a ROI.

In order to begin to perform localization in non-sparse ROIs, we write a posterior distribution on the Binomial parameters used in the likelihood (2) using Bayes rule

$$p(N,\zeta|x) \propto p(x|N,\zeta)p(\zeta)$$
 (3)

We use a Gaussian prior on  $\zeta$  i.e.,  $p(\zeta) = \mathcal{N}(\mu_{\zeta}, \sigma_{\zeta})$ 



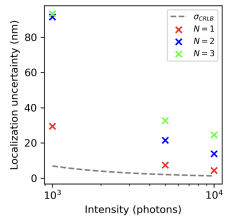


FIG. 2: Single and multi-emitter localization error on sums of photon counts. (left) Localization uncertainty for simulated data for different values of N, plotted with respect to the Cramer-Rao lower bound, shown in dashed gray. (right) Multi-emitter localization by MCMC sampling for N=3, colors indicate a cluster of samples i.e., a single localization. All data was generated with a background rate  $\langle \mathbf{n}_{\text{background}} \rangle = \lambda N_{\text{frames}}/d^2$  per pixel. Scalebar 360nm

with  $\mu_{\zeta}=0.01$  and  $\sigma_{\zeta}=0.005$ . Prior uncertainty in the value of  $\zeta$  stems from fluorophores with potentially heterogeneous photophysical properties as well as varying laser power throughout the excited region. This posterior can be integrated over  $\zeta$  to produce a posterior distribution on the fluor ophore number  ${\cal N}$  i.e.,  $p(N=N'|n) \propto \int_0^1 \prod_j p(n_j|N',\zeta) p(\zeta) d\zeta$  which can be estimated using Monte Carlo methods. The final posterior is then estimated by minibatching the data into batches of  $10^3$  frames and averaging the posterior p(N|n)over minibatches. The fluorophore number N within each ROI is then estimated by the maximum aposteriori (MAP) estimate  $N^*$  given by this distribution.

For localization, we note that the period between excitation pulses is orders of magnitude longer than the fluorescence lifetime, and excitation events are then independent. Total photon counts over  $N_{\text{frames}}$  will therefore follow Poisson statistics, making the localization procedure similar to conventional methods [9]. Denoting the fluorophore coordinates by  $\theta$  and vector of total counts in the region of interest n, we have the following loglikelihood

$$\ell(\mathbf{n}|\theta) = -\log \prod_{k} \frac{e^{-(\mu_{k})} (\mu_{k})^{\mathbf{n}_{k}}}{\mathbf{n}_{k}!}$$

$$= \sum_{k} \log \mathbf{n}_{k}! + \mu_{k} - \mathbf{n}_{k} \log (\mu_{k})$$
(5)

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where, in the multi-emitter regime the expected photon count at a pixel is  $\mu_k = \langle \mathbf{n}_k \rangle = \sum_{m=1}^{N^*} \mu_{k,m}$  given  $\mu_{k,m} = \zeta N_{\text{frames}} \Gamma_u(u_k, x_{0,m}) \Gamma_v(v_k, y_{0,m}) + \lambda N_{\text{frames}} / d^2$ . In the multi-emitter regime, optimization of (4) by sampling is a suitable choice (see Results).

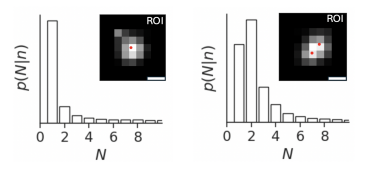


FIG. 3: Posteriors on the number of fluorescent emitters Nand localization for  $N^* = 1$  (left) and  $N^* = 2$  (right) quantum dots. Scalebars 360nm

### Results

Quantum dots coated on a glass coverslip were excited using a picosecond 532nm pulsed laser triggered at 500kHz. Emission light was collected using an oilimmersion 100× objective with numerical aperture (NA) 1.4 (Nikon). The emission signal was then filtered to exclude the laser line (Semrock) and projected onto the SPAD512 sensor (Pi Imaging Technologies) using a tube lens. A simplified diagram of the complete system is depicted in (Figure 1a). Each acquisition consists of  $N = 5 \times 10^5$  frames (500ms), synchronized with each laser pulse, using a 1us exposure per frame (Figure 1b,d). To confirm the presence of single photon sources in the sample, we investigated properties of the zero-lag second order coherence function  $g^{(2)}(0)$ . The following empirical estimate of  $g^{(2)}(0)$  is used [6]

$$g^{(2)}(0) = \frac{G^{(2)}(0) - B}{\langle G^{(2)}(m \neq 0) \rangle - B}$$
 (6)

where  $B=N_{\rm frames}\lambda\zeta$  is the expected number of background-signal coincidences in the region of interest. The quantity  $G^{(2)}(m)$  represents the number of signal-signal coincidences in the region of interest at a lag time m. The quantity  $\langle G^{(2)}(m\neq 0)\rangle$  is the average number of coincidences in pairs of frames at nonzero lag  $m\in[1,100]$ , in units of frames. As expected, simulation from the likelihood (2) shows saturation of  $g^{(2)}(0)$  with increasing values of N (Figure 1c). Moreover, bright clusters of quantum dots exhibit elevated  $g^{(2)}(0)$  values as can be seen in maps of the  $g^{(2)}(0)$  computing using sliding window over the array (Figure 1e).

For localization by optimization of (4), we use Goodman and Weare's Markov Chain Monte Carlo (MCMC) algorithm [5] to sample from the posterior on fluorophore locations. In all simulations we assume a uniform prior on coordinates over the ROI and  $\zeta$  is known and identical over fluorophores. Fluorophore locations can then be estimated from the posterior samples by K-means clustering of the (x,y) coordinates of the first particle and identification of cluster centers (Figure 2a). To validate our estimator, we compare its RMSE to the single emitter Cramer-Rao lower bound, which bounds the variance of a statistical estimator of  $\theta$ , from below (Chao 2016). For an isolated emitter, the Poisson log-likelihood (5) is convenient for computing the Fisher information matrix for  $\theta$  and thus the The Fisher information is [9]

$$I_{ij}(\theta) = \sum_{k} \frac{1}{\mu_k} \frac{\partial \mu_k}{\partial \theta_i} \frac{\partial \mu_k}{\partial \theta_j}$$
 (7)

The Cramer-Rao bound is then found by  $\operatorname{var}(\theta) \geq I^{-1}(\theta)$ . We find that for thousands of expected signal photon counts, localization uncertainty lies in an acceptable range for localization microscopy (Figure 2b). Example posteriors and multi-emitter fitting on experimental quantum dot data are found in (Figure 3). The  $\zeta$  value is treated as unknown but homogeneous across fluorophores in the ROI. A major criticism is likely that we have not measured  $\zeta$  precisely and that we can only provide self-consistency for its simultaenous value with N.

#### Discussion

Many fluorescent emitters exhibit random variations of brightness known as blinking. Blinking increases the observed photon-number fluctuations and could be expected to affect the value of  $g^{(2)}(0)$  or the posterior on the number of active fluorescent emitters. However, the signal photon number per frame will follow Binomial statistics even in the presence of blinking, the only consequence of which is an effective reduction of the detection probability  $\zeta$ . If the effect of censoring photons by blinking

and lowering the quantum yield can be accounted for, the technique used here may be compatible with common super-resolution techniques such as stochastic optical reconstruction microscopy (STORM).

The acquisition times necessary to obtain sufficient photon counts for computing the necessary statistics can potentially be very short. Most fluorophores have relaxation times in the nanosecond range and thus photons can be collected at a rate of at tens of millions of excitation pulses per second. These rates are currently difficult to obtain, however, due to limitations in detector throughput. The SPAD camera used in this study has a minimum exposure time in the microsecond range. Furthermore, the data volume can quickly become intractable due to the need for several thousands of frames for a millisecond-scale exposure time. This is currently a complication for techniques like STORM and advancements in the automation for data acquisitions are necessary. The speed of MCMC based localization remains a limitation for post-processing, and optimization of the processing time for localization is left for future work.

In conclusion, we propose a single molecule imaging technique that allows for simultaneous counting of localization of fluorescent molecules by modeling the quantum properties of fluorescence emission. The technique does not require a nonclassical light source and is designed to supplement standard single molecule localization microscopy techniques. The proposed method can be implemented with a standard widefield fluorescence microscope.

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