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Published in: The Annals of Applied Statistics

DOI (link to publication from Publisher): [10.1214/17-AOAS1054](https://doi.org/10.1214/17-AOAS1054)

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Publication date: 2017

Document Version Publisher's PDF, also known as Version of record

[Link to publication from Aalborg University](https://vbn.aau.dk/en/publications/2b8669b9-221e-432c-a912-aaa8a8ac763c)

Citation for published version (APA): Hooghoudt, J. O., Barroso, M., & Waagepetersen, R. P. (2017). Toward Bayesian inference of the spatial distribution of proteins from three-cube Förster resonance energy transfer data. The Annals of Applied Statistics, 11(3), 1711-1737.<https://doi.org/10.1214/17-AOAS1054>

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TOWARD BAYESIAN INFERENCE OF THE SPATIAL DISTRIBUTION OF PROTEINS FROM THREE-CUBE FÖRSTER RESONANCE ENERGY TRANSFER DATA¹

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Förster resonance energy transfer (FRET) is a quantum-physical phenomenon where energy may be transferred from one molecule to a neighbor molecule if the molecules are close enough. Using fluorophore molecule marking of proteins in a cell, it is possible to measure in microscopic images to what extent FRET takes place between the fluorophores. This provides indirect information of the spatial distribution of the proteins. Questions of particular interest are whether (and if so to which extent) proteins of possibly different types interact or whether they appear independently of each other. In this paper we propose a new likelihood-based approach to statistical inference for FRET microscopic data. The likelihood function is obtained from a detailed modeling of the FRET data-generating mechanism conditional on a protein configuration. We next follow a Bayesian approach and introduce a spatial point process prior model for the protein configurations depending on hyperparameters quantifying the intensity of the point process. Posterior distributions are evaluated using Markov chain Monte Carlo. We propose to infer microscope-related parameters in an initial step from reference data without interaction between the proteins. The new methodology is applied to simulated and real datasets.

1. Introduction. In the biology community there is a vast interest in studying the biomolecular structure and dynamics of macromolecular assemblies in order to understand their functions [\[Alber et al.](#page-24-0) [\(2017\)](#page-24-0), [Polo and Jackson](#page-26-0) [\(2011\)](#page-26-0), [Krissinel](#page-25-0) [and Henrick](#page-25-0) [\(2007\)](#page-25-0), [Puglisi](#page-26-0) [\(2005\)](#page-26-0)]. Because the interactions between proteins and the typical size of proteins (1–100 nm) is at the nanoscale level [\[Erickson](#page-25-0) [\(2009\)](#page-25-0)], no information can be obtained from conventional optical microscopic techniques, which at best can resolve distances down to ∼200 nm [\[van Putten et al.](#page-26-0) [\(2011\)](#page-26-0)]. Instead, Förster resonance energy transfer—also referred to as fluorescence resonance energy transfer—microscopy is widely used to obtain such information. Förster resonance energy transfer (FRET) provides information about distances of

Received August 2016; revised March 2017.

¹Supported in part by the Center for Stochastic Geometry and Advanced Bioimaging which is funded by the Villum Foundation and by the Danish Council for Independent Research—Natural Sciences, Grant 7014-00074B, "Statistics for point processes in space and beyond."

Key words and phrases. Bayesian inference, Markov chain Monte Carlo, Förster resonance energy transfer, spatial point process, spatial distribution, proteins, fluorophores.

the order of 2 to 10 nm within or between molecular structures and is the preferred tool for investigating spatial relationships in biochemistry [\[Wu and Brand](#page-27-0) [\(1994\)](#page-27-0), [Gryczynski, Gryczynski and Lakowicz](#page-25-0) [\(2005\)](#page-25-0), [Clegg](#page-25-0) [\(1995,](#page-25-0) [2006\)](#page-25-0)].

FRET is the nonradiative transfer of the surplus of energy from an excited *donor* fluorophore (fluorescent molecule) to a sufficiently nearby *acceptor* fluorophore by dipole–dipole interaction [\[Heitler](#page-25-0) [\(1954\)](#page-25-0), [Rohatgi-Mukherjee](#page-26-0) [\(1978\)](#page-26-0)]. The widespread use of FRET in biological research is based on the possibility to label, in vivo or in vitro, proteins with fluorophores that are spectrally matched [\[Miyawaki, Sawano and Kogure](#page-26-0) [\(2003\)](#page-26-0), [Bunt and Wouters](#page-25-0) [\(2004\)](#page-25-0)]. The energy transfer due to the FRET mechanism is a stochastic process, and the probability that energy transfer occurs between a donor and an acceptor fluorophore is heavily dependent on the distance between them. The probability that energy transfer occurs is commonly referred to as the efficiency of the energy transfer. The usefulness of FRET lies in the fact that various techniques exist by which the fraction of donor excitations that result in energy transfer—that is, the efficiency—can be quantified.

Two main methods for determining the FRET efficiency are as follows: fluorescence lifetime measurements [\[Wallrabe and Periasamy](#page-26-0) [\(2005\)](#page-26-0), [Lakowicz](#page-26-0) [\(2009\)](#page-26-0), [Chen et al.](#page-25-0) [\(2013\)](#page-25-0)] and spectral methods [\[Sun et al.](#page-26-0) [\(2011\)](#page-26-0), [Zimmermann, Riet](#page-27-0)[dorf and Pepperkok](#page-27-0) [\(2003\)](#page-27-0)]. We focus in this paper on the most commonly applied spectral method called three-cube FRET. Due to the FRET mechanism, a certain fraction of the de-excitations of a donor result in energy transfer to an acceptor instead of donor photon emission, thereby the rate by which photons are emitted from the donors decreases—a phenomenon referred to as quenching of the donor—while instead photons are emitted by the acceptors. Spectral methods now rely on determining the decrease in the donor emission due to FRET. For three-cube FRET, intensity measurements are carried out using three different filter sets—often referred to as cubes—each comprising an excitation filter, a dichroic mirror, and an emission filter. This results in three digital intensity images [\[Zal and](#page-27-0) [Gascoigne](#page-27-0) [\(2004\)](#page-27-0), [Wallrabe et al.](#page-26-0) [\(2006\)](#page-26-0), [Periasamy et al.](#page-26-0) [\(2008\)](#page-26-0), [Periasamy and](#page-26-0) [Day](#page-26-0) [\(2011\)](#page-26-0)]. Two images are obtained by exposing the sample to light in the donor absorption spectrum and recording emitted intensities both in the donor and the acceptor excitation spectrum. The third image is obtained by exposing the sample to light in the acceptor spectrum and also recording light in the acceptor spectrum.

Given FRET image data, the task is to obtain information concerning the spatial configuration of the donors and acceptors in the sample. For example, [Wallrabe](#page-26-0) [et al.](#page-26-0) [\(2003\)](#page-26-0) study the clustering of ligand–receptor complexes in endocytic membranes using confocal FRET microscopy. They differentiate between a clustered or a random distribution of proteins by considering the dependence of the FRET efficiency on donor and acceptor concentrations. In particular, independence of the efficiency on acceptor concentration or a decrease in the efficiency for a higher unquenched donor signal for a fixed acceptor concentration are both indicators for clustering [\[Kenworthy and Edidin](#page-25-0) [\(1998\)](#page-25-0), [Kenworthy](#page-25-0) [\(2001\)](#page-25-0)]. [Goswami et al.](#page-25-0)

[\(2008\)](#page-25-0) instead compare observed distributions of fluorescence intensity and fluorescence anisotropy with values expected from a Poisson distribution of nanoclusters.

Other, computational, approaches mainly rely on the construction of a configuration of donors and acceptors and computing the FRET efficiency related to this configuration by numerical computation of the energy transfer probabilities for each of the donors [\[Wolber and Hudson](#page-27-0) [\(1979\)](#page-27-0), [Corry, Jayatilaka and Rigby](#page-25-0) [\(2005\)](#page-25-0)]. This simple approach has been extended by various authors by simulating FRET events explicitly using Monte Carlo techniques. The extended approach gives the possibility to include additional physical complexity into the model to account for possible photobleaching of donors and acceptors during a FRET measurement or the effect that temporarily unavailable acceptors can have on the FRET efficiency [\[Frederix et al.](#page-25-0) [\(2002\)](#page-25-0), [Berney and Danuser](#page-24-0) [\(2003\)](#page-24-0), [Corry, Jayatilaka](#page-25-0) [and Rigby](#page-25-0) [\(2005\)](#page-25-0)]. [Corry, Jayatilaka and Rigby](#page-25-0) [\(2005\)](#page-25-0) further carefully studied the FRET efficiency in relation to various fixed donor and acceptor configurations (e.g., pentamers) and give a concise overview of the development of the numerical Monte Carlo approaches.

[Loura and Prieto](#page-26-0) [\(2011\)](#page-26-0), [Loura, Fernandes and Prieto](#page-26-0) [\(2010\)](#page-26-0) and [Lakowicz](#page-26-0) [\(2009\)](#page-26-0) give excellent reviews of methods to extract spatial information in membrane biophysics from FRET data. Methods determining the complex structures of a protein or the spatial distribution of protein complexes in living cells are given in e.g. [Raicu et al.](#page-26-0) [\(2009\)](#page-26-0) and [Bonomi et al.](#page-24-0) [\(2014\)](#page-24-0).

The previous mentioned contributions are based on a detailed understanding of the FRET data generating mechanism. This knowledge, however, so far has not been applied to obtain a complete statistical model of FRET data allowing for a principled statistical analysis. In this paper we present a first attempt to conduct a full likelihood-based Bayesian analysis of three-cube FRET image data. The potential advantages of such an approach are that the posterior distribution gives detailed quantitative information regarding model parameters and donor–acceptor interactions as well as measures of uncertainty regarding this information. To obtain the likelihood function, we derive, based on physical considerations, an accurate statistical model for the distribution of the image intensities, conditional on a point pattern consisting of donors and acceptors. We further impose a spatial point process prior [\[Møller and Waagepetersen](#page-26-0) [\(2004\)](#page-26-0)] for the unknown configuration of donors and acceptors. Since our resulting posterior distribution is of a complicated form, we use Markov chain Monte Carlo (MCMC) to sample from the posterior distribution [\[Gamerman and Lopes](#page-25-0) [\(2006\)](#page-25-0), [Gilks, Richardson and Spiegelhalter](#page-25-0) [\(1996\)](#page-25-0)]. It is difficult to infer simultaneously microscope-related parameters and possible interactions between donors and acceptors. We therefore propose to infer microscope-related parameters in an initial step based on reference data without interactions between donors and acceptors. We asses the Bayesian inference procedure by a simulation study and by applying it to an empirical in-vitro reference dataset.

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2. Observation model for three-cube FRET image data. A three-cube FRET dataset consists of three images each corresponding to a rectangular region *W* which is a union of rectangular pixels, $W = \bigcup_{i \in \mathcal{G}} C_i$, indexed by a grid \mathcal{G} . Each pixel *Ci* records a light intensity due to emission from donors or acceptors. The images are created by (1) excitation of donors and measurement of donor emission, (2) excitation of acceptors and measurement of acceptor emission or (3) excitation of donors and measurement of acceptor emission (due to FRET). We represent the images by vectors $Y_{DD} = (Y_{DD}^i)_{i \in \mathcal{G}}$, $Y_{AA} = (Y_{AA}^i)_{i \in \mathcal{G}}$ and $Y_{DA} = (Y_{DA}^i)_{i \in \mathcal{G}}$. The first letter in the subscripts denotes whether donors (D) or acceptors (A) were excited, and the second letter denotes in which channel emission was measured. We assume that a pixel value Y_k^i , $k = DD$, AA, DA, $i \in \mathcal{G}$, is subject to additive normal noise; that is,

$$
(2.1) \t Y_k^i = I_k^i + \varepsilon_i,
$$

where I_k^i denotes light intensity due to emission and the noise terms ε_i are independent and $N(0, \sigma^2)$ distributed.

We now specify models for the I_k^i given configurations of donor and acceptor proteins in *W* whose positions form point patterns, respectively, X_D and X_A .

2.1. *Some fluorescence resonance energy transfer theory*. An excited donor $d \in X_D$ surrounded by a configuration X_A of acceptors can de-excite in three ways: either by emission with a rate k_{DE} , by nonradiative decay (e.g., internal heat conversion) with a rate k_{DN} , or by FRET to an acceptor $a \in \mathbf{X}_A$ with a rate $k_{F,da}$. We will refer to the sum of the first two mechanisms as the *intrinsic* de-excitation rate k_D , that is, $k_D = k_{DE} + k_{DN}$. According to [Förster](#page-25-0) [\(1948\)](#page-25-0), $k_{F,da}$ is given by

$$
k_{F,da} = k_D \left(\frac{R_0}{\|d - a\|}\right)^6,
$$

where R_0 is the so-called Förster distance, defined as the distance between the donor and acceptor at which the de-excitation rate due to FRET equals the intrinsic de-excitation rate; that is, $k_{F,da} = k_D$ if $\|d - a\| = R_0$. The probability that *d* deexcites due to FRET to a specific donor a in X_A thus becomes

$$
P_{da} = \frac{k_{F,da}}{k_D + \sum_{\widetilde{a} \in \mathbf{X}_A} k_{F,d\widetilde{a}}} = \frac{(R_0/\|d-a\|)^6}{1 + \sum_{\widetilde{a} \in \mathbf{X}_A} (R_0/\|d-\widetilde{a}\|)^6}.
$$

Figure [1](#page-5-0) shows an example of the computation of P_{da} for a specific configuration of acceptors *a* around a donor *d*. The total probability that *d* de-excitates due to FRET is $P_{dA} = \sum_{a \in \mathbf{X}_A} P_{da}$. The probability that *d* de-excites by emission or by nonradiative decay is $P_{dD} = 1 - P_{dA}$.

FIG. 1. (a) *Donor fluorophore surrounded by three acceptors at distances* R_0 , $\frac{1}{2}R_0$ *and* $2R_0$ *.* (b) The table shows the de-excitation "path widths" $(R_0/||d - a||)^6$ for energy transfer from the *donor to each of the acceptors and the corresponding energy transfer probabilities Pda*.

2.2. *Model for intensities given protein configurations*. Our model for the intensities given the configurations X_D and X_A is inspired by the model for simulation of FRET data in [Corry, Jayatilaka and Rigby](#page-25-0) [\(2005\)](#page-25-0). However, in contrast to [Corry, Jayatilaka and Rigby](#page-25-0) [\(2005\)](#page-25-0), we introduce the simplifying assumption that a donor or acceptor is always available for excitation [see also [Wolber and Hudson](#page-27-0) [\(1979\)](#page-27-0), [Berney and Danuser](#page-24-0) [\(2003\)](#page-24-0)]. This is a reasonable assumption if the intensity of the laser is moderate so that the inter arrival times of photons at a donor are large compared with the de-excitation times. We can then regard the times of excitations of donors and acceptors as Poisson processes and use standard results for Poisson processes to obtain closed-form distributional results for the I_k^i .

In the [Appendix](#page-23-0) we show that $I_{DD}^i = G_D N_{DD}^i$ and $I_{DA}^i = G_A N_{DA}^i$, where N_{DD}^i and N_{DA}^i are the number of photons detected by the detector in, respectively, the DD-channel and the DA-channel, and N_{DD}^i and N_{DA}^i are both Poisson distributed. Further, G_D and G_A are unknown positive parameters related to the sensitivity of the detector in, respectively, the donor and acceptor emission spectrum. The means of I_{DD}^i and I_{DA}^i are

$$
\mu_{DD}^i = M_D \sum_{d \in \mathbf{X}_D \cap C_i} (1 - P_{dA})
$$

and

$$
\mu_{DA}^i = GM_D \sum_{a \in \mathbf{X}_A \cap C_i} \sum_{d \in \mathbf{X}_D} P_{da},
$$

where M_D and G are unknown positive parameters. We assume that the means μ_{DD}^i/G_D and μ_{DA}^i/G_A of, respectively, N_{DD}^i and N_{DA}^i are sufficiently large so that the Poisson distributions of N_{DD}^i and N_{DA}^i can be well approximated by normal distributions. Then

(2.2)
$$
I_{DD}^i \sim N(\mu_{DD}^i, G_D \mu_{DD}^i)
$$
 and $I_{DA}^i \sim N(\mu_{DA}^i, G_A \mu_{DA}^i)$.

By a similar line of arguments, we also obtain

$$
(2.3) \tI_{AA}^i \sim N(\mu_{AA}^i, G_A \mu_{AA}^i),
$$

where

$$
\mu_{AA}^i = (M_D/K)n(\mathbf{X}_A \cap C_i),
$$

K is an unknown positive parameter and $n(X_A \cap C_i)$ denotes the number of acceptors within pixel C_i . In cases of intensity data with a large proportion of zeros, we instead use truncated normal distributions with point masses at zero for I_{DD}^i , I_{DA}^i and I_{AA}^i ; see Section 2 of Supplement B [\[Hooghoudt and Waagepetersen](#page-25-0) [\(2017a\)](#page-25-0)].

Equations (2.1) , (2.2) and (2.3) specify the distribution of the FRET data conditional on the protein configurations X_D and X_A . The distribution is parameterized by $\psi = (M_D, G, K, G_D, G_A, \sigma^2)$. We refer to the components of ψ as microscope-related parameters. The parameters *G* and *K* are known as the socalled *G*- and *K*-factors [\[Zal, Zal and Gascoigne](#page-27-0) [\(2002\)](#page-27-0), [Chen, Puhl and Ikeda](#page-25-0) [\(2007\)](#page-25-0)]. The parameter M_D can be interpreted as the mean donor emission detector read-out intensity due to one donor excitation.

3. Bayesian inference of spatial characteristics of protein configurations.

We adopt a Bayesian approach to infer the microscope-related parameters ψ and spatial characteristics of the configurations X_D and X_A of proteins. A spatial point process prior (specified in Section [3.1\)](#page-7-0) is used for $X = (X_D, X_A)$, where this prior again depends on a parameter vector θ . We also assign a prior to θ , thus including also this parameter in the posterior inference. As detailed later in Sections [4.4](#page-13-0) and [5,](#page-15-0) we recommend to infer the microscope-related parameters in an initial step using reference data without interactions between donors and acceptors. In a second step, investigating interactions in a dataset of biological scientific interest, *ψ* can then be fixed at estimates obtained from the first step. Letting *y* denote an observation of $Y = (Y_{DD}, Y_{DA}, Y_{AA})$ and $(\mathbf{x}_D, \mathbf{x}_A)$ a realization of **X**, the joint posterior distribution is

(3.1)
$$
p(\mathbf{x}_D, \mathbf{x}_A, \theta | y, \psi) \propto p(y | \mathbf{x}_D, \mathbf{x}_A, \psi) p(\mathbf{x}_D, \mathbf{x}_A | \theta) p(\theta).
$$

Here $p(z)$ and $p(z|u)$ are generic notation for a probability density of a random quantity *Z* and the conditional density of *Z* given another random quantity $U = u$.

3.1. *Priors*. We model a priori **X***^D* and **X***^A* as independent Poisson processes on *W* with intensities θ_D and θ_A ; that is, the prior density of $(\mathbf{X}_A, \mathbf{X}_D)$ with respect to independent unit rate Poisson processes is

(3.2)
$$
p(\mathbf{x}_D, \mathbf{x}_A | \theta) = \theta_A^{n(\mathbf{x}_A)} \theta_D^{n(\mathbf{x}_D)} \exp[-|W|(\theta_A + \theta_D - 2)],
$$

where $|W|$ denotes the area of *W* and $n(\mathbf{x})$ denotes the number of points in a point configuration **x** [see, for instance, equation (6.2) in [Møller and Waagepetersen](#page-26-0) [\(2004\)](#page-26-0)]. We further impose independent conjugate Gamma hyperpriors for θ_D and *θA*. The Poisson prior can be viewed as a null model for the case of no interaction between donors and acceptors or within donors, respectively, acceptors. Compared with other more complex point process models like Markov point processes (like the Strauss hard core model considered in Section [4\)](#page-9-0), the Poisson prior is advantageous in having a known density function. A potential problem is that the Poisson prior is in some sense a strong prior which assigns little probability to point configurations with strong clustering or regularity. This can lead to biased results as demonstrated in Section [4.](#page-9-0) Densities for more flexible Markov point process prior models, on the other hand, contain intractable normalizing constants that depend on the unknown parameters in the point process model. This then precludes the use of standard Markov chain Monte Carlo algorithms (Section 3.2) for evaluation of the posterior distribution.

The Gamma distributions for θ_D and θ_A are defined through shape parameters *α* and rate parameters *β*. As the mean of the Gamma distribution is *α/β* and its variance α/β^2 , the signal-to-noise ratio related to the distribution is defined by the square root of the shape parameter, that is,

$$
\frac{S}{N} = \frac{\alpha/\beta}{\sqrt{\alpha/\beta^2}} = \sqrt{\alpha}.
$$

In a typical FRET experiment there is quite some uncertainty concerning the true values of the numbers of proteins within the sample, and so we have defined not too confined priors for the intensities θ_D and θ_A . We have chosen to set the signalto-noise ratio always equal to 2, resulting in the value of 4 for the shape parameter. In our applications we further specify the prior mean *m* of each of the parameters so that the rate parameter β follows from $\beta = \alpha/m = 4/m$. We also use Gamma priors for the components of ψ ; see the discussion of prior elicitation for ψ in Section [6.](#page-16-0)

3.2. *Markov chain Monte Carlo*. To evaluate the posterior distribution, we use a Markov chain Monte Carlo algorithm [\[Gamerman and Lopes](#page-25-0) [\(2006\)](#page-25-0)] where the components (X_D, X_A) , θ_D, θ_A and (if applicable) the components of ψ are updated in turn. Gibbs updates are used for the full conditional Gamma distributions of θ_D and θ_A , while random walk Metropolis updates on the log scale are used for the components of ψ . For the point configurations (X_A, X_D) we first randomly choose

to either update X_A or X_D (with probability 1/2 for each choice). We then use birthdeath updates as outlined in Sections 7.1.2–7.1.3 in [Møller and Waagepetersen](#page-26-0) [\(2004\)](#page-26-0). If, for example, \mathbf{X}_A is chosen to be updated, then with probability $1/2$ it is proposed to remove a point chosen from the uniform distribution on **X***A*. Otherwise it is proposed to insert a new acceptor point at a location chosen from the uniform distribution on *W*. In case it is proposed to remove a point $u \in \mathbf{X}_A$, the Metropolis– Hastings ratio becomes

$$
\frac{p(\mathbf{x}_D, \mathbf{x}_A \setminus \{u\}, \theta | y, \psi) n(\mathbf{x}_A)}{p(\mathbf{x}_D, \mathbf{x}_A, \theta | y, \psi) |W|} = \frac{p(y | \mathbf{x}_D, \mathbf{x}_A \setminus \{u\}, \psi) n(\mathbf{x}_A)}{p(y | \mathbf{x}_D, \mathbf{x}_A, \psi) \theta_A |W|}.
$$

If it is proposed to insert a new acceptor point $v \in W$, then the Metropolis–Hastings ratio is

$$
\frac{p(y|\mathbf{x}_D,\mathbf{x}_A\cup\{v\},\psi)\theta_A|W|}{p(y|\mathbf{x}_D,\mathbf{x}_A,\psi)(n(\mathbf{x}_A)+1)}.
$$

The expressions for updating X_D are similar. The described birth-death updates are repeated a large fixed number of times between the updates of the parameters θ_D , θ_A and ψ .

To keep the MCMC updates for donor and acceptor points numerically feasible, only those acceptors that reside within $4R_0$ of a donor are taken into account as a possible path for energy transfer for the donor. This important simplification will not lead to any significant difference in posterior results, as the transfer probability *P_{da}* for a donor *d* and an acceptor *a* is very small when $\|d - a\| > 4R_0$, thereby adding or removing a point in pixel *i* can only affect the values of likelihood factors $p(y_{DD}^l, y_{DA}^l, y_{AA}^l | \mathbf{x}_D, \mathbf{x}_A, \psi)$ for pixels *l* in a neighborhood of *i* [note that the likelihood factors as $\prod_{i \in \mathcal{G}} p(y_{DD}^i, y_{DA}^i, y_{AA}^i | \mathbf{x}_D, \mathbf{x}_A, \psi)$. Exploiting this simplification, we have implemented an ingenious algorithm that recomputes the transfer probabilities *Pda* only for donors and acceptors which are influenced by the adding/removing of a point. A detailed description of the MCMC sampler is provided in Supplement B [\[Hooghoudt and Waagepetersen](#page-25-0) [\(2017a\)](#page-25-0)].

3.3. *Inferring spatial characteristics*. In statistics for spatial point processes, the *K*-function is a common tool for inferring interactions from a spatial point pattern. We adapt this approach and use the cross *K*-function [e.g., [Møller and](#page-26-0) [Waagepetersen](#page-26-0) [\(2004\)](#page-26-0)] to measure interactions between donors and acceptors given point configurations x_A and x_D . In general, for point processes X_1 and X_2 of intensities ρ_1 and ρ_2 , $\rho_2 K_{12}(t)$ is the expected number of \mathbf{X}_2 points within distance *t* from a typical point of X_1 . In case of no interaction between X_1 and X_2 , $K_{12}(t) = \pi t^2$. Values of $K_{12}(t)$ greater (smaller) than πt^2 signifies positive (negative) interaction between \mathbf{X}_1 and \mathbf{X}_2 . It is common to consider the cross *L*-function $L_{12}(t) = \sqrt{K_{12}(t)/\pi}$ which is equal to *t* in the case of no cross interaction, while $L_{12}(t) > t$ [$L_{12}(t) < t$] means positive (negative) cross interactions. We will refer to $L_{12}(t) - t$ as the "centered" cross *L*-function. The centered cross *L*-function is

equal to zero in the case of no cross interaction, while larger (smaller) than zero for positive (negative) cross interactions.

Given configurations \mathbf{x}_D and \mathbf{x}_A of donors and acceptors, we estimate the cross *K*-function by

$$
\widehat{K}_{\mathbf{x}}(t) = \sum_{u \in \mathbf{x}_A, v \in \mathbf{x}_D} \frac{1[\|u - v\| \le t]}{n(\mathbf{x}_A)n(\mathbf{x}_D)|W \cap W_{u-v}||W|^{-2}},
$$

where W_{u-v} is *W* translated by $u - v$ [e.g., Section 4.4.3 in [Møller and](#page-26-0) [Waagepetersen](#page-26-0) [\(2004\)](#page-26-0)]. The cross *L*-function is estimated by $\widehat{L}_{\mathbf{x}}(t) = \sqrt{\widehat{K}_{\mathbf{x}}(t)/\pi}$. To infer cross spatial interactions between donors and acceptors given FRET data, we consider the posterior distribution of $\widehat{L}_{\mathbf{X}}(t) = \sqrt{\widehat{K}_{\mathbf{X}}(t)/\pi}$ or its centered version. We also considered so-called cross *G*- and *J* -functions [e.g., [Møller and](#page-26-0) [Waagepetersen](#page-26-0) [\(2004\)](#page-26-0)], but in our simulation studies the cross *L*-function gave a more clear impression of the nature of donor–acceptor interactions.

4. Simulation studies. Our primary target of inference is the cross *L*function, \hat{L}_X for the configuration $\mathbf{X} = (\mathbf{X}_A, \mathbf{X}_D)$ of donors and acceptors which is unknown in practice. However, we also need to infer the microscope-related parameters ψ . From a Bayesian perspective, if the right prior distribution is chosen, the posterior distribution by definition provides the correct inference given the data *Y* and prior information. However, in our case, the Poisson prior [\(3.2\)](#page-7-0) is partly chosen for convenience in order to yield tractable MCMC computations, and is not necessarily the best possible representation of prior information. Thus, from a pragmatic point of view, it makes sense to assess possible bias of our Bayesian inference procedure.

In particular, we focus in Section [4.3.1](#page-11-0) on the posterior mean $L_{|Y}$ of $L_{\mathbf{X}}$ as a predictor of $\widehat{L}_{\mathbf{X}}$. The posterior mean $L_{|Y}$ is further an estimate of $L^E = E[\widehat{L}_{\mathbf{X}}]$, which is the expected value of $\overline{L}_{\bf X}$ over replicated data **X**. Note in this connection that had we used the true distribution of \overline{X} as the prior, then $EL|Y|$ and L^E would be exactly equal—that is, $L_{|Y}$ would be an unbiased predictor/estimate both of L_X and L^E . In Section [4.3.3](#page-13-0) we assess the performance of the full posterior distribution of L_X given *Y* for inference regarding L_X . In Section [4.3](#page-10-0) we consider ψ to be a fixed known parameter. Section [4.4](#page-13-0) is concerned with inference regarding *ψ*.

4.1. *Simulation of synthetic data*. To generate synthetic data for the simulation study, the point configuration $\mathbf{X} = (\mathbf{X}_A, \mathbf{X}_D)$ is generated on a 1000 nm by 1000 nm square region as a realization of a bivariate Strauss hard core process. This point process has density (with respect to a bivariate process of independent unit rate Poisson processes) of the form

(4.1)
$$
f(\mathbf{x}_A, \mathbf{x}_D) \propto \beta_D^{n(\mathbf{x}_D)} \beta_A^{n(\mathbf{x}_A)} \gamma^{s_R(\mathbf{x}_A, \mathbf{x}_D)} \text{HC}(\mathbf{x}_A, \mathbf{x}_D, \mathfrak{R}_A, \mathfrak{R}_D, \mathfrak{R}_{DA}),
$$

where $s_R(\mathbf{x}_A, \mathbf{x}_D)$ is the number of unordered pairs of points $\{u, v\}$ with $u \in \mathbf{x}_A$, $v \in x_D$, and interpoint distance $||u - v||$ less than *R*. Values of γ less than one lead to repulsion between donors and acceptors, while values of *γ* greater than one lead to attraction. The term $HC(\mathbf{x}_A, \mathbf{x}_D, \mathcal{R}_A, \mathcal{R}_D, \mathcal{R}_{DA})$ is one if the following hard core condition is satisfied: all donors have an interpoint distance greater than \Re_b , all acceptors have an interpoint distance greater than \Re_A , and all pairs of points where one is a donor and the other an acceptor have an interpoint distance greater than \Re_{DA} . Otherwise the hard core term is zero, whereby it serves to model that donors and acceptors have a physical extent that prevents them from getting arbitrarily close to each other. Different settings of the Strauss hard core process parameters are used to create different point pattern types described in Section 4.2 below.

Next, conditional on the configuration **X** and the various microscope-related parameters ψ , the intensity data *Y* is generated from the model specified in Section [2.](#page-4-0) Regarding the observation model, we fix the measurement variance σ^2 at 25, let each of *G, K, G_D, G_A* equal to 1, and consider values 1, 5, 20 of M_D in order to generate data of varying signal to noise ratios defined by $\mathbf{E}[\mu_{DD}^i/(G_D\mu_{DD}^i+\sigma^2)^{1/2}]$ and $\mathbf{E}[\mu_k^i/(G_A\mu_k^i+\sigma^2)^{1/2}], k = DA, AA$. For each point pattern type we generate 100 independent synthetic point patterns $\mathbf{X}^{\text{synth},i}$ and associated synthetic image data $Y^{\text{synth},i}$, $i = 1, \ldots, 100$.

4.2. *Point pattern types*. The basic point pattern types considered are *dimer*, *clustered, Poisson hard core* and *repulsive*. For all types, \Re_D , \Re_A and \Re_{DA} are at least 2 nm. The parameters β_D and β_A are further adjusted to have on average 1000 donors and 1000 acceptors.

In case of *dimer*, we specify large values of $\Re_D = \Re_A$, which essentially means that only proteins of different types can appear close to each other. Thus the only clusters possible are mini-clusters consisting of one donor and one acceptor, that is, dimer clusters. For the *clustered* case, $\Re p = \Re A$ are reduced, which enables formation of a wider range of clusters containing several donors and acceptors. In case of *dimer* and *clustered*, values of $\gamma_{DA} = 2$, 8 correspond to respectively moderate and strong interactions. For *Poisson hard core*, $\gamma_{DA} = 1$, while all hard core distances are 2. In case of *repulsive*, varying values of \Re_{DA} , *r* and γ_{DA} < 1 generate different strengths of repulsive interaction between donors and acceptors. Table [1](#page-11-0) gives an overview of the different parameter settings considered.

4.3. *Inference regarding spatial characteristics*. We estimate *L^E* by the empirical average of L-functions $\widehat{L}_{\mathbf{X}^{\text{synth},i}}$ obtained from the $\mathbf{X}^{\text{synth},i}$. From each synthetic dataset $Y^{\text{synth},i}$ we further obtain an MCMC estimate $\overline{L}_{|Y^{\text{synth},i}}$ of the posterior mean $L_{|Y}$ synth,*i* of \widehat{L}_X given Y ^{synth,*i*}. The mean posterior *L*-function $EL_{|Y}$ is estimated by the mean of the $\bar{L}_{|Y}$ synth,*i*. The sampling variability of $L_{|Y}$ is further represented by the variation of the $\bar{L}_{|Y^{synth},i}$. When considering inference for the cross *L*-function in the following Sections [4.3.1](#page-11-0)[–4.3.3,](#page-13-0) ψ is fixed at the value used for generating the synthetic datasets.

Parameter settings used in the Strauss hard core model [\(4.1\)](#page-9-0) *to create the various point pattern types. The values for the homo* (\Re_{DD} , \Re_{AA}) *and hetero* (\Re_{DA}) *hard core distances and the hetero interaction radius* (*R*) *presented in the table follow from the considerations stated in the text and by the choice of the Förster distance* $R_0 = 6$

4.3.1. *Bias of posterior mean*. We assess the bias of *L*|*^Y* by considering the mean (over replicated data **X**, Y) of L_X and its prediction $L_{|Y}$, thereby we also assess how $L_{|Y}$ performs as an estimate of L^E (the expected cross L -function for X).

In Figures [2](#page-12-0) and [3](#page-13-0) the estimates of $L^{E}(t) - t$ and $EL_{1Y}(t) - t$ are shown for each of the point pattern types in Table 1 for the three values of M_D (1, 5, 20). In Figure [2\(](#page-12-0)a), for example, L^E for dimer-type 1.2 shows that the underlying point patterns are clustered for distances $r < 16$ (as $L^E > 0$) and are slightly repulsive for distances $16 < r < 24$. Further, for distances $24 < r < 40$, there seems to be some slight clustering again, while, for $r > 40$, the pattern displays complete spatial randomness. The negative values that occur for $r < 5$ are due to the minimum imposed hardcore distance of 2. The negative part for small r is also visible in the plots (b)–(h), where L^E otherwise indicates clustering among donors and acceptors and in plot (i), where $L^{E}(t) - t$ is close to zero otherwise.

The general impression from the plots is that *L*|*^Y* is biased downward when the true point patterns are of dimer or clustered types (a)–(h) and biased upward in the cases of the repulsive types (i) – (m) (including the Poisson hard core case). For $M_D = 1$, where the signal-to-noise ratio is very low, the mean of L_V is very close to zero, and it does not seem possible to infer in this case cross interactions between donors and acceptors. However, for $M_D = 5, 20$, there is always a pro-

FIG. 2. *In each plot the solid line is the centered LE-function*. *The other lines show the centered EL*_|*Y for varying* M_D : *dashed line*: $M_D = 1$ *, dotted line*: $M_D = 5$ *, dashed-dotted line*: $M_D = 20$. *The plots are for the dimer and clustered point pattern types*.

nounced peak (positive or negative) of the mean $L_{|Y}$ -function where the peak is of the right sign and located in the right place of the peak of the L^E -function. Moreover, the bias consistently decreases when M_D and hence the signal-to-noise ratio increases. Despite the bias, the results suggest that qualitatively correct statements can be made regarding independence, clustering or repulsion between donors and acceptors.

4.3.2. *Variability of posterior mean L-function*. In addition to bias, the extent to which valid qualitative conclusions can be made from the posterior *L*|*^Y* -function of course also depend on its variability. Figure [4](#page-14-0) shows for each basic point pattern type and M_D either 5 or 20, 98% envelopes for $L_Y(t)$ based for each *t* on the minimal and maximal values of $L_{|Y}(t)$ over the 100 replications. These envelopes are fairly narrow for distances up to 100 and show that, in the setting of the simulation study, qualitative conclusions regarding the nature of interaction between donors and acceptors will be consistently correct over replicates.

FIG. 3. *Continuation of Figure* [2.](#page-12-0) *Plots are for the Poisson hard core and repulsive types*.

4.3.3. *Inference based on full posterior distribution*. The results in Sec-tion [4.3.1](#page-11-0) showed that the posterior mean *L*-function $L_{|Y}$ can exhibit substantial bias as an estimate of L^E and hence also as a predictor of \widehat{L}_X . This can invalidate the use of the full posterior distribution for inferring the uncertainty regarding the estimation of L^E or the prediction of \widehat{L}_X . As an example Figure [5](#page-15-0) shows $L_{|Y^{\text{synth}},1}(t) - t$ and the 98% central posterior interval for $\widehat{L}_{\textbf{X}}(t) - t$ given $Y = Y_{synth,1}$ for the same point pattern types as in Figure [4](#page-14-0) and $M_D = 5$, 20. Also, the true $\widehat{L}_{\mathbf{X}^{\text{synth},1}}(t) - t$ are shown in each plot.

For the lower signal-to-noise ratio with $M_D = 5$, $\widehat{L}_{\mathbf{X}^{\text{synth},1}}(t)$ falls outside the 98% posterior interval for several point pattern types. Thus the posterior intervals do not always give a useful quantification of the uncertainty regarding the knowledge of L_X . However, for the higher signal-to-noise ratio with $M_D = 20$, the envelopes do include or almost include the $\hat{L}_{\mathbf{X}^{\text{synth},1}}$ function.

4.4. *Simulation studies for microscope parameters*. So far the vector *ψ* of microscope-related parameters has been assumed to be known, which is rarely the case. We have investigated Bayesian inference for ψ in a simulation study for which the full details are given in Supplement C [\[Hooghoudt and Waagepetersen](#page-25-0) [\(2017b\)](#page-25-0)]. We here just comment on results obtained for simulations with $G_D =$ $G_A = K = G = 1$, $M_D = 20$ and $\sigma^2 = 25$. Figure [6](#page-16-0) shows boxplots of the posterior mean of each microscope parameter over 40 replicated datasets for each point pattern type. The main features are as follows:

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FIG. 4. *Distribution of the centered* $\bar{L}_{|Y}$ *-function for the various point pattern types, summarized by* 98% envelopes based on $\overline{L}_{|Y^{\text{synth},i}}$, $i = 1, ..., 100$, together with the mean value $\widehat{E}L_{|Y}$ (middle *solid line*). *The value of M_D is* 5 *or* 20. *In each plot twenty of the* $\overline{L}_{|Y^{\text{synth},i}}$ *'s are shown with solid gray curves*, *the dashed line is the centered L E-function*.

1. for the Poisson hard core patterns the posterior means of all the parameters coincide with or are very close to their respective synthetic values [see plots (a) – (f) for type number 9].

2. for all the clustered patterns (type number $1-8$) inference for M_D , G and K is biased. The posterior means \overline{M}_D are significantly below their synthetic value, while \bar{G} and \bar{K} are above their target values. Further, the bias increases for the patterns generated with $\gamma_{DA} = 8$ (type number 2, 4, 6, 8) compared to the corresponding patterns generated with $\gamma_{DA} = 2$ (type number 1, 3, 5, 7).

3. for repulsive patterns (type number $11-13$) M_D and *K* are on or close to target, while \overline{G} is negatively biased.

4. the posterior means of σ , G_A and G_D are on target for almost all point pattern types [plots (d)–(f)].

As explained in detail in Supplement C, the biased results are due to the mismatch between the Poisson point process prior and the actual point processes used for the

FIG. 5. Posterior distribution of $\widehat{L}_{\bf X}$ given the first synthetic dataset $Y^{\text{synth},1}$ for each point pat*tern type*. *Dashed*: *posterior mean L*|*^Y* synth*,*¹ , *dashed-dotted*: 98*% envelopes and solid gray*: *twenty posterior realizations of L* **X**. *Solid black shows the true L* **X**synth*,*¹ .

simulations. The microscope-related parameters thus a posteriori take on values to "soothe" this mismatch, resulting in biased results. Further, in this setting with joint inference of ψ and spatial characteristics, the posterior mean *L*-functions are strongly biased as well, being close to zero for all distances for all point pattern types (not shown).

5. Two-step approach to likelihood-based inference. In the previous Section [4.4](#page-13-0) we observed that applying the Bayesian inference methodology using the Poisson process prior on Poisson hard core patterns gave reliable estimates for all microscope parameters. This suggests an approach where microscoperelated parameters are inferred from reference datasets constructed with absence of donor–acceptor interactions. Therefore, we propose a two-step approach where the microscope-related parameters are inferred in a first step using reference data. In the second step the values of the microscope parameters are fixed at the posterior estimates from the first step in order to make inference on the spatial configura-

FIG. 6. *Boxplots of the posterior mean of all the six microscope parameters for the forty replicated runs for each of the point pattern types (referred to by their type number, see Table [1\)](#page-11-0) for* $M_D^s = 20$ *. The horizontal lines are drawn at the corresponding synthetic values*.

tion of donors and acceptors of a three-cube FRET sample of biological scientific interest. To illustrate the approach, we have carried out the first step on empirical reference three-cube FRET data, as discussed in the following section.

6. Data example. In this section we apply our Bayesian methodology to empirical in vitro three-cube FRET data obtained from donor or acceptor fluorophore labeled transferrin proteins [\[Welch](#page-27-0) [\(1992\)](#page-27-0)] attached to polylysine slides [\[Shima](#page-26-0) [and Sakai](#page-26-0) [\(1977\)](#page-26-0)]. Transferrin bound to polylysine is known to be approximately randomly (i.e., Poisson hardcore) distributed [\[Wallrabe et al.](#page-26-0) [\(2007\)](#page-26-0)]. The objective is to infer the microscope parameters related to the experimental setup.

We initially conducted an exploratory analysis [described in Supplement A, [Hooghoudt, Barroso and Waagepetersen](#page-25-0) [\(2017\)](#page-25-0)] where we quantified the amount of photobleaching and compared empirical mean–variance relationships of the image data with the ones implied by our model. From these mean–variance relationships, as well as other non-Bayesian methods discussed in Supplement A, we obtained rough estimates for the microscope parameters that were used to set the prior means in the Bayesian analysis. We thus use a pragmatic Bayesian approach where the rough non-Bayesian estimates entering in the priors are refined by introducing information obtained through the likelihood derived from our observation model.

Due to certain computational issues discussed in Section 6.2 and Section [6.3.1,](#page-19-0) we are at this stage only able to use a small subset of the full data in the Bayesian inference. Improving the computational methodology is an important topic of further research.

6.1. *The image dataset*. Three cube FRET measurements have been carried out on three samples, to which we refer as samples 1, 2 and 3. Sample 1 is prepared to consist of twice as many donors (D) as acceptors (A), that is, $D : A \approx 1 : \frac{1}{2}$, while samples 2 and 3 are prepared such that, respectively, $D : A \approx 1 : 1$ and \overline{D} : $A \approx \frac{1}{2}$: 1. Three-cube FRET data is obtained on each sample on a square grid containing 512×512 square pixels. The pixel side length is 0.279 μ m and the focal volume depth is approximately 5 pixels $(1.4 \mu m)$ [\[Wallrabe et al.](#page-26-0) (2007)]. The image data are shown in Figure [7.](#page-18-0)

The emission in the DD-channel (Y_{DD}) and AA-channel (Y_{AA}) are corrected for background emission, while the DA-channel data (Y_{DA}) is also corrected for spectral bleedthrough by the methods described in [Elangovan et al.](#page-25-0) [\(2003\)](#page-25-0).

We noticed that around the edges of the 512×512 images, often very low or zero intensity regions occurred due to improper sample preparation. Therefore, the exploratory statistical analysis has been based solely on the central rectangular section of the images consisting of 100×100 pixels (see also Supplement A, Section 1).

In order to obtain sufficient photon count statistics—that is, sufficiently high signal-to-noise ratio for each pixel—each sample has been remeasured ten times. We then create an aggregated dataset by summing pixelwise over the ten measured intensities for each channel. We note that by remeasuring the sample instead of increasing the measurement time, we obtain information concerning the amount of photobleaching occurring for remeasurements (Supplement A, Section 3) and the pixel intensity variance in the three channels. The latter information gives the possibility to deduce the empirical mean–variance relationship of the image data and to obtain estimates for G_D and G_A (Supplement A, Section 6).

6.2. *Inference procedure setup*. The prior distributions for the microscope and Poisson point process parameters were specified as Gamma distributions. For each

FIG. 7. *Channel intensity images of the aggregated channel dataset of sample* 2. (a) *DD-channel*, (b) *AA-channel*, (c) *DA-channel*. *Plots* (a)*–*(c) *consist each of* 512 × 512 *pixels*. *Plots* (d)*–*(f) *show enlargements of the square subregions of the plots* (a)–(c), *each consisting of* 100×100 *pixels*. *Above each plot is stated the mean* (*me*), *maximum* (*ma*) *and minimum* (*mi*) *pixel intensity value in the image*. *In each image the gray levels are constructed by using ten equally spaced intervals between zero and the maximum value of the image*. *Black/white refers to the lowest/highest intensity interval*.

parameter the shape parameter is set to 4 based on the reasoning in Section [3.1,](#page-7-0) and we use a pragmatic Bayesian approach where the prior means of five of the microscope parameters are set by aid of the rough estimates $M_D \approx 2.6$, $G \approx 0.7$, $K \approx$ 0.7, $G_D \approx 7.4$ and $G_A \approx 5.5$ obtained from the preliminary statistical analysis in Supplement A, Sections 3–6 and 8. The prior mean of the measurement noise σ^2 we have set, rather ad hoc, to 50.

By the statistical analysis in Section 8 of Supplement A, it was further found that the point process intensities θ_A and θ_D of the samples can be roughly related to the donor and acceptor solution concentrations applied for the sample preparation, thereby we found that, for sample 1, $\theta_D \approx 2e^2 \mu m^{-2}$ and $\theta_A \approx 1e^2 \mu m^{-2}$; while, for sample 2, $\theta_D \approx 2e3 \ \mu m^{-2}$ and $\theta_A \approx 2e3 \ \mu m^{-2}$; and, for sample 3, $\theta_D \approx$ 1e3 μ m⁻² and $\theta_A \approx 2e^3 \mu$ m⁻². We use these values as prior means for θ_A and θ_D for each of the samples. The applied prior means as used in the Bayesian analyses are summarized in Table [2.](#page-19-0)

In the MCMC computations we used random walk Metropolis–Hastings updates for the log microscope-related parameters. The values of the random walk update standard deviations τ are also shown in Table [2.](#page-19-0) We tuned the τ values

Prior mean 2*.*5 1.0 1.0 5.0 5.0 50*.*0 2e3 1e3 2e3 2e3 1e3 2e3 *τ* 0*.*05 0.1 0.1 0.5 0.5 1*.*0– – – – – –

to get an approximately 30% proposal acceptance rate for each of the microscope parameters. The total number of MCMC updates is 5e9 for each run. Metropolis– Hastings updates for the microscope parameters and Gibbs updates for the point process parameters are made after every 1e4 birth/death updates of donor or acceptor points.

The MCMC chains converge slowly due to bad mixing as discussed in Section 6.3. This means that we need many rounds of birth-death updates for the donor/acceptor points followed by updates of microscope and point process parameters. In each round we need to update a large fraction of the donors and acceptors. Thus, for a fixed fraction, each round takes more computing time the higher posterior expected number of donors and acceptors. The a posteriori expected number of donors and acceptors in each pixel is fairly high (of the order 300). To keep the computation time at an acceptable level, we therefore perform the inference on a small 10×10 subset of pixels which contain a posteriori of the order of 3e4 donors and acceptors.

6.3. *Results of the inference*.

6.3.1. *Assessment of MCMC samples*. In Figure [8](#page-20-0) the traceplots of the microscope parameters and the Poisson point process intensities for sample 1 are shown. Posterior mean values are displayed in the upper left corner of each of the plots. The traceplots indicate poor mixing of the MCMC samples except for G_A , G_D and σ .

The poor mixing is due to high posterior correlation between certain parameters as visualized by the scatterplots in Figure [9](#page-21-0) in which the posterior realizations of M_D , G , K , θ_D , θ_A are plotted against each other. Especially M_D and θ_D and G and θ_A are highly correlated, but fairly strong correlations are also evident between *M_D* and *K* and between *K* and θ_D . Similar scatterplots of G_D , G_A and σ^2 versus each of the other parameters (not shown) do not show any clear correlation with any of the parameters.

6.3.2. *Posterior results*. In Table [3](#page-21-0) the 95% posterior intervals and posterior means for each of the microscope parameters and Poisson intensities are stated

FIG. 8. *Traceplots of the microscope and the Poisson point process parameters for sample* 1. *Posterior mean values are displayed in the upper left corner of each of the plots*. *For plotting a subsampling of* 10 *has been applied*.

for each of the three samples. The microscope parameters M_D , G , G_A , G_D and *σ* should be equal for all samples, and this may seem contradicted by their posterior means that vary across samples. There is, on the other hand, considerable overlap between almost all 95% posterior intervals so that the Bayesian inference does not contradict that the microscope parameters are equal across samples.

Figure [10](#page-22-0) shows the posterior distributions of the centered *L*-function for the three samples. As expected, there is no indication of clustering nor repulsion since the centered posterior *L*-functions are close to zero and the posterior means are approximately zero.

7. Discussion. This paper presents a first attempt to implement likelihoodbased inference for FRET data. We thus, based on physical considerations, devel-

FIG. 9. *Scatterplots of the posterior realizations of MD,G,K,θD,θA versus each other for sample* 1. *For plotting a subsampling of* 100 *has been applied*.

oped a realistic observational model for FRET data given the underlying configurations of donors and acceptors. Based on this model, we proposed to implement Bayesian inference using MCMC.

	Sample		
		2	3
M_D	$2.3 - 7.0$ (4.2)	$3.1 - 8.6(5.5)$	$2.8 - 6.8(4.7)$
G	$0.46 - 1.21(0.78)$	$0.50 - 1.36(0.89)$	$0.19 - 0.52(0.32)$
K	$0.61 - 1.93(1.08)$	$0.31 - 0.98(0.60)$	$0.51 - 1.49(0.92)$
G_A	$1.9 - 6.2(4.0)$	$5.3 - 12.7(8.9)$	$8.0 - 13.4(10.6)$
G_D	$1.2 - 5.7(3.3)$	$3.0 - 10.0$ (6.3)	$1.7 - 6.5(3.9)$
σ	$4.4 - 12.0(8.3)$	$3.8 - 10.7(7.2)$	$4.2 - 11.6(7.9)$
θ_D	$6.1e2 - 19.3e2(11.1e2)$	$7.8e2 - 2282(13.4e2)$	$9.0e2 - 23.5e2(14.7e2)$
θ_A	$6.1e2 - 15.0e2(9.7e2)$	$8.3e2 - 20.4e2$ (12.8e2)	15.0e2-34.1e2 (23.5e2)

TABLE 3 *Posterior results for each of the three samples*: 95*% posterior intervals and posterior means* (*in brackets*) *for each of the parameters*. *Prior means of the parameters are given in Table* [2](#page-19-0)

FIG. 10. *Posterior distribution of the L-function for samples* 1*–*3. *For each sample the posterior distribution is summarized by the posterior mean* (*solid line*) *of the L-function and* 95*% envelopes based on minimal* (*lower dashed-dotted line*) *and maximal* (*upper dashed-dotted line*) *values of* 39 *posterior realizations of the L-function*.

We quantify spatial dependence by considering the posterior mean of the cross *L*-function for the donors and acceptors. Our simulation results show that the posterior mean of the *L*-function can be used to distinguish between clustering, absence of interaction and repulsion between donors and acceptors. Due to bias, one needs to be careful when making quantitative statements regarding strength of interaction based on the posterior means of the *L*-functions. However, we believe that it is meaningful to make relative comparisons of strength of interactions between samples observed under the same experimental conditions and thus with the same signal-to-noise ratios.

Partly due to poor mixing of the proposed MCMC procedure, we were forced to consider only a small subset of the full data. A key objective for further research is therefore to obtain a more efficient MCMC scheme so that efficient use of the full data becomes feasible. [Haario, Saksman and Tamminen](#page-25-0) [\(2001\)](#page-25-0) suggest to use joint updates, but they consider posterior distributions of fixed dimensional random vectors. However, preliminary experiments with this approach indicate that we need joint updates involving both the microscope parameters and the donor-acceptor point patterns. It is not clear how to do this. Our data example illustrated the use of reference data with no donor–acceptor interactions to infer the microscope-related parameters. In future work it would interesting to apply an improved MCMC algorithm to conduct Bayesian inference for an experimental sample with possible interactions.

The Poisson point process prior for protein configurations was chosen partly for computational reasons. To implement Bayesian inference with more flexible Markov point process priors allowing for both repulsive and attractive interactions requires more advanced Markov chain Monte Carlo methods developed in [Møller](#page-26-0) [et al.](#page-26-0) [\(2006\)](#page-26-0) and [Murray, Ghahramani and MacKay](#page-26-0) [\(2006\)](#page-26-0). However, these methods are highly computationally demanding since they involve so-called perfect simulation from the point process prior, which can lead to unacceptable computing times in the case of protein configurations of high cardinality which are frequently encountered for FRET data.

APPENDIX: DERIVATION OF OBSERVATION MODEL

In this section we refer to notation introduced in Section [2.](#page-4-0) Let L_d denote the set of times in the observation time span $[0, T]$ where a donor *d* in X_D is excited by a photon from the laser. We assume that L_d is a homogeneous Poisson process on [0, T] with intensity $\lambda_D > 0$. The process L_d can be decomposed as

$$
L_d = L_{dE} \cup L_{dN} \bigcup_{a \in \mathbf{X}_A} L_{da},
$$

where L_{dE} denotes the times of excitations of *d* which resulted in emission in the D channel, L_{dN} is the times of excitations resulting in nonradiative de-excitation and *Lda* denotes the times of excitations that resulted in FRET to acceptor *a* and subsequent emission in the *A* channel. The so-called quantum yield 0 *<* q_D < 1 is the probability of emission for donors conditional on that de-excitation is by emission or nonradiatively, that is, $q_D = k_{DE}/k_D$. Similarly, q_A denotes the quantum yield for acceptors. Invoking the random labeling theorem for Poisson processes [e.g., Proposition 3.7 in [Møller and Waagepetersen](#page-26-0) [\(2004\)](#page-26-0)], L_{dE} , L_{da} and L_{dN} are independent Poisson processes with intensities $\lambda_D q_D P_{dD}$, $\lambda_D q_A P_{da}$ and $\lambda_D[(1 - q_D)P_{dD} + \sum_{a \in \mathbf{X}_A} (1 - q_A)P_{da}]$.

For each $i \in \mathcal{G}$ we let

$$
L_{DD}^i = \bigcup_{d \in \mathbf{X}_D \cap C_i} L_{dE}
$$

and

$$
L_{DA}^i = \bigcup_{d \in \mathbf{X}_D, a \in \mathbf{X}_A \cap C_i} L_{da}
$$

be the Poisson processes of donor excitation times which result in photon emissions for respectively donors and acceptors in the pixel *Ci*.

The emitted photons fall on the detector independently of each other with a probability $0 < h < 1$ of detection. In the point process literature the detected photons are called an independent thinning with retention probability *h*. Further, of the detected photons only independent thinnings with retention probabilities $0 < Q_D < 1$ ($0 < Q_A < 1$) are registrated in the donor (acceptor) channel of the detector. The probabilities Q_D and Q_A are respectively the detector quantum yields in the donor and acceptor channel [\[Pawley](#page-26-0) [\(2006\)](#page-26-0)]. The detected photon counts N_{DD}^i and N_{DA}^i of emissions in the pixel C_i are thus Poisson distributed with means $Q_D h q_D \lambda_D T \sum_{d \in \mathbf{X}_D \cap C_i} P_{dD}$ and $Q_D h q_A \lambda_D T \sum_{d \in \mathbf{X}_D, a \in \mathbf{X}_A \cap C_i} P_{d a}$. Finally, $I_{DD}^i = G_D N_{DD}^i$ and $I_{DA}^i = G_A N_{DA}^i$, where G_D and G_A are amplification factors depending on the detector and channel. Defining $M_D = G_D Q_D h q_D \lambda_D T$

and $G = (G_A Q_A q_A)/(G_D Q_D q_D)$, we arrive at the specified means of I_{DD}^i and I_{DA}^i .

The mean of I_{AA}^i is found in a similar fashion. In the AA-channel acceptors are directly excited by the laser which is now broadcasting in the acceptor excitation spectrum with an intensity $\lambda_A > 0$. An excited acceptor can only de-excite due to emission or nonradiatively, and the detected photon counts N_{AA}^i of emissions in the pixel C_i are Poisson distributed with means $Q_A h q_A \lambda_A T \sum_{a \in X_A \cap C_i}^{\infty} 1$ and $I_{AA}^i = G_A N_{AA}^i$. Defining $M_A = G_A Q_A h q_A \lambda_A T$ and $K = M_D / M_A$, we arrive at the specified mean of I_{AA}^i .

Acknowledgments. The authors sincerely thank the Editor, Associate Editor and referees for their constructive suggestions and comments, which contributed substantially to the improvement of the work.

SUPPLEMENTARY MATERIAL

Supplement A: Preliminary statistical analysis of the in vitro three-cube FRET dataset (DOI: [10.1214/17-AOAS1054SUPPA](http://dx.doi.org/10.1214/17-AOAS1054SUPPA); .pdf). The supplementary material contains a preliminary statistical analysis of the in vitro three cube FRET dataset. Sample preparation and experimental setup are discussed as well the channel dataset extracted from the samples. The amount of photobleaching for the remeasurements in the three channels is studied and the mean–variance relations of the dataset are compared with the ones given by our statistical model. Further, estimates for the microscope parameters are obtained through various non-Bayesian methods.

Supplement B: The MCMC sampler (DOI: [10.1214/17-AOAS1054SUPPB;](http://dx.doi.org/10.1214/17-AOAS1054SUPPB) .pdf). The supplementary material contains a detailed description of the steps performed in the MCMC sampler.

Supplement C: Inference of the microscope parameters (DOI: [10.1214/17-](http://dx.doi.org/10.1214/17-AOAS1054SUPPC) [AOAS1054SUPPC;](http://dx.doi.org/10.1214/17-AOAS1054SUPPC) .pdf). The supplementary material contains a detailed description of the inference of the microscope parameters.

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