Log-Gaussian Cox processes and sampling paths:

towards optimal design

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Abstract

Goal of this paper (placeholder abstract). Evaluate a wide variety of path designs

in terms design-based heuristics and model-based criteria for spatial prediction

using Bayesian LGCP models. Identify promising initial designs for later op-

timization and sequential design (not actually optimizing yet). Illuminate any

relationships among design characteristica and predictive criteria that will be

helpful for constrained optimization. Discuss sequential contruction of paths as

a precursor to online sequential design.

Keywords: log-Gaussian Cox process, optimal sampling, model-based design,

spatial sampling design

1. Introduction

Spatial point process models have long been considered generally infeasible

because of their computational demands, but recent advances in Bayesian com-

puting have made the Log-Gaussian Cox process an attainable model in prac-

tice (Rue et al., 2009; Lindgren et al., 2011; Illian et al., 2012; Simpson et al.,

2016). Variable sampling effort leads to a degraded point pattern Chakraborty

et al. (2011) and it is relatively simple to accomodate variable sampling effort

in these models using modern computing tools (Yuan et al., 2017). However,

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the literature on optimal sampling for spatial point process models is in its infancy (Liu and Vanhatalo, 2020). In this article, we present a variety of sampling path designs and assess their optimality for LGCP models.

Point pattern data are routinely collected in species distribution studies and ordnance response projects. These applications may use quadrat sampling or line-transect sampling, with transect sampling being more common. When the objective is mapping where events occur in space, various spatial mapping procedures have been used. Traditionally these have involved aggregating the data to grid cell counts or computing moving averages. Aggregation has the downside of introducing arbitrary structure into the data by the choice of gridding scheme or averaging window, and requires uneccessary computation effort (Simpson et al., 2016). Software is now available to fit spatial point process models to data acquired via distance sampling and simultaneously estimate the detection function (Johnson et al., 2014; R Core Team, 2019).

In ecological settings, sampling plans are often designed around the goal of estimating total abundance. Ordnance response surveys are typically designed with the objective of detecting (but not necessarily mapping) intensity hotspots (USACE, 2015; Flagg et al., 2020). However, to our knowledge, there has been very little work done in deciding where to collect data when the goal is to map the intensity using a spatial point process model. In this paper, we adapt nearest-neighbor criteria to the path design setting and introduce a sequential path construction scheme as a starting point for future optimization. We then compare a variety of path design schemes with respect to a suite of model-based and design-based criteria for simulated point pattern data.

1.1. Spatial design

Design-based sampling. Most classical sampling work has been done for points or small quadrats approximated as points. Space-filling criteria may be good starting points (Borkowski and Piepel, 2009). Latin hypercube sampling has space-filling properties McKay et al. (1979); Husslage et al. (2011).

Space-filling curves. Used in design of dense or stretchable circuits (Ogorzałek, 2009; Ma and Zhang, 2016) and high-dimensional data visualization in bioinformatics (Anders, 2009). Peano curve is very flexible for filling irregular shapes (Fan et al., 2014).

Space-filling curves are one-dimensional paths constructed iteratively; as the number of iterations goes to infinity, the limiting path has nonzero area and actually fills the space (Sagan, 1994). For applications we stop after a finite number of iterations. The Hilbert curve is fast and simple to construct.

Model-based spatial design. Regularity is optimal for spatial prediction but randomness and a variety of interpoint distances are best for parameter estimation (Diggle and Lophaven, 2006). Inhibitory plus close pairs is a good compromise (Chipeta et al., 2017).

50 1.2. Paths as sampling designs

While some ideas about the characteristics of a good point design apply to paths, creating an optimal path design is not as simple as connecting the points of a point design with line segments. There are many ways to connect points into a path, so optimal design criteria must apply to the whole path and not only to the waypoints.

Pollard et al. (2002) adaptively zigzagged their line transects in a species abundance survey.

The Visual Sample Plan software includes features to create systamatic transect plans and augment plans with additional transects in regions lacking spatial coverage (Matzke et al., 2014). It helps the user choose the transect spacing to maximize the probability of detecting the presence of a hotspot of specified size and intensity. However, it does not employ criteria to optimize spatial prediction.

Liu and Vanhatalo (2020) used narrow quadrats (swaths along line-transects) as their sampling units. The transects were short relative to the size of the study region and not connected into a path.

2. Materials and methods

Heuristics of a good path design:

- Should start with a sparse design with regular spacing, then refine with infill
 - Provides good spatial coverage even if aborted early
 - Imagine downloading a high-resolution intensity jpeg over 56k
 - Path should avoid sharp turns but is allowed to cross itself
- One option is to generate two segments at a time, first a short-to-medium length segment to get to the start of the next transect, then a mediumto-long segment for the transect
 - Could have new segment length be negatively correlated with the previous segment length

2.1. Design-based criteria

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We give attention to some design-based criteria that tie directly to practical considerations of data collection.

Path length. The total distance that traveled is often a constraint. Minimize it.

Corners. Data collection equipment (e.g. metal detectors) may have limited mobility, requiring minimizing the nubmer or angle of turns.

- Nearest neighbor distance. A common criterion for space-filling designs, we adapt it to be meaningfully calculated for any point on a path. Define the kth-order nearest neighbor distance as $\operatorname{nnd}_k(u) = \min |u v|$ where v is any point in the set of path segments at least k steps away from the segment containing u. If k = 0, this includes v in the same segment as u so trivially $\operatorname{nnd}_0(u) = 0$
- for all u in the path. $\operatorname{nnd}_1(u)$ includes all segments except the one containing u. $\operatorname{nnd}_2(u)$ excludes the segment containing u and segments with which it shares

vertices. Segments not accessible by a connected path starting at u are always included. Maximize $\min[\operatorname{nnd}_2(u)]$, $\operatorname{avg}[\operatorname{nnd}_2(u)]$, and $\operatorname{avg}[\operatorname{nnd}_1(u)]$.

(Move details to appendix.)

5 2.2. Model-based criteria

Mean-squared prediction error. Minimize MSPE for the GP.

Posterior prediction variance. Minimize maximum prediction variance and average prediction variance for GP.

2.3. Sampling schemes

These are the focus, move them earlier?

Parallel transects. Parallel transects running the length of the site in the vertical axis. Three ways of choosing the horizontal coordinate: simple random sample (SRS), systematic with a random starting point and even spacing, inhibitory plus close pairs.

Latin hypercube sampling. Random Latin hypercube design connected by shortest path. Waypoints generated by the 1hs R package Carnell (2020). Connected into a the shortest path by the TSP package (Hahsler and Hornik, 2020).

Space-filling curves. Hilbert curve generated by HilbertVis package (Anders, 2009). This is a deterministic design, so a random offset is added.

Particle movement model. Models the way data are actually collected. Waypoints generated sequentially by generating a jump distance and a direction.

The jump distance is generated from a scaled beta distribution, and negatively
correlated with previous jump distance. This behavior should approximately
alternate between a short "transition" and a long transect. The negative correlation was achieved by applying a 1-x transformation to a beta autoregressive
process (McKenzie, 1985). The direction angle is drawn from a bimodal distribution that is symmetric around 0 (a normal distribution reflected about 0).

explain the Strauss part

set up to think about adaptive sampling (adding a transect at a time or stopping early but don't actually do it here)

2.4. Model fitting

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INLA (Rue et al., 2009), SPDE (Lindgren et al., 2011), off-grid (Simpson et al., 2016)

2.5. Simulation procedure

We consider a fictitious site \mathcal{R} with the simple shape of a 1500 unit by 700 unit rectangle. In this site, we will simulate two data generating models meant to produce random intensity functions with with hotspots. First, a LGCP with latent GP mean $\mu = \log(250/|\mathcal{R}|)$ and a Matérn covariance with $\nu = 1$, $\sigma = 2$, and range = 200. This model produces relatively unstructured hotspots due to large variability in the GP.

Second, the superposition of a two-stage cluster process superposed and a LGCP. The cluster process (a Neyman-Scott or, more specifically, a Thomas process) is constructed as follows. The number of clusters is Poisson with mean 3. The number of events per cluster is Poisson with mean 200. The cluster centers distributed uniformly over \mathcal{R} . Events come from a bivariate normal distribution with mean equal to the cluster center and variance $\Sigma = \tau^2 \mathbf{I}$, $\tau = 50$. The LGCP has $\mu = \log(250/|\mathcal{R}|)$ and Matérn covariance with $\nu = 1$, $\sigma = 1$, and range = 200. This model is based upon the typical conceptual model of a firing range, with a background process (represented by the LGCP) and a small number of higher-intensity foreground clusters containing the events of interest.

Path design schemes:

- Simple random sample of north-south line transects
 - Number of transects = 10, 25, 50, 70
 - Expect high variance, large prediction error in big gaps.
- Systematic sample of north-south line transects

- Number of transects = 10, 25, 50, 70
- Uniformly distributed starting point
- Constant spacing

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- Expect low bias and ok variance, can miss structures at certain sizes,
 may not have best space-filling properties.
- Systematic sample of north-south serpentine transects
 - Number of transects = 7, 22, 47, 67
 - Uniformly distributed starting point
 - Constant spacing
 - Number of zigzags = 5.8
 - Horizontal zigzag distance set so that the total horizontal distance traveled equals 2100 units (the length of 3 non-zigzag line transects)
 - Expect better space-filling properties than line-transect designs, lower bias/variance farther from path, would be better at estimating anisotropic covariance than line-transects.
- Inhibitory plus close pairs sample of north-south line transects
 - Total number of transects (including pairs) = 10, 25, 50, 70
 - Number of pairs = 0.1, 0.2 times the total number of transects (rounded up or down to nearest whole number)
 - Pairs uniformly distributed within radius of primaries, max pair radius =
 1500/total number of transects
 - Position of primaries generated from a 1-dimensional Strauss process with $\gamma = 0.05$
 - A compromise between SRS and systematic in every way.
- Latin Hypercube Sampling waypoints
 - Number of bins = 50,300,1200,2400

Expect low bias/variance per unit distance traveled, many sharp corners, some big open areas.

• Hilbert curve

- Order = 3, 4, 5, 6
 - Created in square and then scaled to fit in \mathcal{R}
 - A uniform random offset added equal to spacing between segments
 - Expect good space filling, good bias and variance, lots of short segments.

• Random particle movement

- Distance cutoff = 6700, 17200, 34700, 49700
- Segment lengths uniform 50 to 500 units
- Adjacent segments uncorrelated or $\rho = -0.8$
- Turn angle $N(\mu = \pi/3, \sigma = \pi/6)$ or $N(\mu = \pi/2, \sigma = \pi/12)$
- Angle multiplied by discrete uniform over $\{-1,1\}$
- Strauss-esque thinning, antirepulsion = 0.8, pair radius = 80
- All combinations of the above, plus pair distance of 300 for 6700 distance cutoff
- Expect variation in all characteristics due to extreme randomness, but some near-optimality that could be harnessed by search algorithms, should see exploration followed by infill, negative ρ with turns centered tightly on $\pi/2$ should mimic zigzagging among parallel transects.

(Probably should move explanations of schemes to appendix.)

100 designs from each scheme. All events within a 2 unit radius of the path are observed. Whole experiment repeated for 5 realizations from each data generating model.

Model:

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Mesh with dual colored by node weight

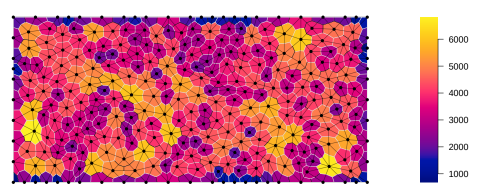


Figure 1: Illustration of the mesh and associated numerical integration scheme used to approximate the latent GP.

- **X** is a Poisson process on \mathcal{R} with intensity $\lambda(u)$
- $\log[\lambda(u)] = \mu + \mathbf{e}(u)$
 - $\mu \sim \text{Unif}(-\infty, \infty)$

 - PC prior on σ and ρ with $\Pr(\sigma > 3) = 0.1$ and $\Pr(\rho < 100) = 0.1$ (Fuglstad et al., 2019; Simpson et al., 2017)
 - SPDE approach of Lindgren et al. (2011) using mesh in Figure 1
 - Likelihood factorization of Simpson et al. (2016)

3. Results

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look at examples of designs that minimize each criterion look at examples of designs along the Pareto front

4. Discussion

discuss starting points for optimization and sequential design

5. Conclusions

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Appendix A. Notation and Terminology

- process defined on $\mathcal{D} \subset \mathbb{R}^d$, domain of the intensity function, in this manuscript d=2
 - observation window $S \subset \mathcal{D}$
 - define three regions:
 - the domain \mathcal{D} over which the process mathematically operates
 - the study region \mathcal{R} over which inferences are desired
 - the observed/sampled observation window \mathcal{S}
 - general relationship is $S \subset \mathcal{R} \subset \mathcal{D} \subset \mathbb{R}^d$ where all of the subset symbols taken to mean "subset or equal"
 - \mathcal{D} can be bounded or unbounded (often equal to \mathbb{R}^d), \mathcal{S} practically always bounded, \mathcal{R} bounded or unbounded depending on application and inferential goals
 - the "fully surveyed" (censused) situation is $\mathcal{S} = \mathcal{R}$
 - survey path \mathcal{P} is a one-dimensional subset of \mathcal{R}
 - set of one or more sequences of waypoints connected by line segments
 - ${\mathcal S}$ is the set of all points within a fixed (and assumed known) radius of ${\mathcal P}$
 - **X** point process on \mathcal{R} , $\mathbf{x} = \{x_1, \dots, x_n\}$ realized point pattern
 - $-\mathbf{X}_{\mathcal{S}} = \mathbf{X} \cap \mathcal{S}$ the restriction of \mathbf{X} to \mathcal{S} , $\mathbf{x} = \mathbf{X} \cap \mathcal{S}$ the realized observeable point pattern

• point $x \in \mathbf{x}$ called an event

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- intensity function $\lambda(u)$
- types of "points" in space:
 - -x event in the point pattern
 - s numerical integration node
 - -u arbitrary location in \mathcal{D} used to index intensity function and predictors
- z(u) a column vector of covariates/predictors at u (not used in this manuscript)
- \bullet "point" refers to a u unless clearly stated otherwise
- bold for sets and spatial processes, normal italics for spatial vectors
- y and variations will be used for objects derived from the point pattern, e.g. marks, pseudodata
 - distance sampling fits into the framework with expansion of notation to include a (nontrivial) detection function and differentiate between the observed and observable point patterns

Appendix B. Extension of Nearest Neighbor Distance to Paths

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