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1	Spatial targeting of infectious disease control: identifying multiple, unknown sources
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19 Summary

21	1. Geographic profiling (GP) was originally developed as an analytical tool in criminology, where it uses the spatial					
22	locations of linked crimes (for example murder, rape or arson) to identify areas that are most likely to include the					
23	offender's residence. The technique has been extremely successful in this field, and is now widely used by police					
24	forces and investigative agencies around the world. More recently, the same method has been applied to biological					
25	data, notably in spatial epidemiology, where it uses the locations of disease cases to identify infection sources: the					
26	identification of these sources is critical to control efforts of diseases such as malaria, since targeted intervention is					
27	more efficient and cost effective than untargeted intervention.					
28	2. Here we solve the problem of identifying multiple sources, even when the number of sources is unknown – a					
29	requirement for many biological studies. We present a new, rigorous mathematical and computational method, and					
30	show why previous Bayesian methods were often outperformed by the empirically-developed Criminal Geographic					
31	Targeting (CGT) algorithm used in criminology.					
32	3. We use simulations and real-world examples to compare our model to both the CGT algorithm and to an existing					
33	Bayesian model. We demonstrate that our method combines the advantages of both previous methods, particularly					
34	in cases featuring large data sets and multiple sources.					
35	4. Our approach provides an increase in search efficiency over other methods and is likely to lead to improved					
36	targeting of interventions and more efficient use of resources. We suggest that the Dirichlet process mixture (DPM)					
37	model provides a useful and practical tool for conservation biologists and epidemiologists that can be used to inform					
38	management decisions and public health policy.					
39						
40	Kevwords					
41	Bayesian statistics, criminology, Dirichlet process mixture, epidemiology, geographic profiling					
42						
43	Abbreviations					
44	GP, geographic profiling; DPM, Dirichlet process mixture; MCMC, Markov Chain Monte Carlo					
45						

46 Introduction

47 In many areas of biology (for example invasion biology and epidemiology), models describing the ways in which 48 animals, plants or pathogens spread outwards from a central source are of considerable importance. Such models are 49 routinely used to generate risk maps in epidemiology, or to predict the effect of global climate change on the spread 50 of invasive species (Kolar & Lodge 2001). Surprisingly, very few models exist which run backwards in time, using 51 current spatial patterns to identify sources of infections or biological invasions, despite the fact that the identification 52 of these sources can be used to target control efforts, dramatically improving the efficiency of interventions. 53 Recently, geographic profiling (GP) – a technique originally developed in criminology to help prioritise large lists of 54 suspects in cases of serial crime (Rossmo, 2000) – has been successfully applied to biological data, providing a way 55 of doing exactly this (Le Comber & Stevenson 2012).

56

57 Investigations of serial crime typically involve too many, rather than too few, suspects; for example, the 58 investigation into the Yorkshire Ripper murders in the UK between 1975 and 1980 generated 268,000 names 59 (Doney 1990). In criminology, GP techniques use spatial data concerning the locations of connected crime sites to 60 create a surface of search priority that is overlaid on a map of the study area to produce a geoprofile, which in turn 61 allows the police to prioritise investigations by systematically checking suspects associated with locations in 62 descending order of the height on the geoprofile (Rossmo 2000). There are a number of different geographic 63 profiling software programs available, including Rigel (Miller 2003), developed by Environmental Criminology 64 Research Inc. (ECRI), CrimeStat (Levine 1996), funded by the U.S. National Institute of Justice, and Dragnet (Canter 2000), developed at the University of Liverpool. Other authors (for example Snook et al. (2002, 2005)) have 65 66 made a case for the use of human judges. Of different programmes available, the most widely used is the criminal geographic targeting (CGT) algorithm of Rossmo (Rossmo 1993), which forms the basis of Rigel (Miller 2003), in 67 68 which information from multiple crime sites is combined by means of summing over independent distributions. The 69 CGT is used by organisations including the Royal Canadian Mounted Police, the Bureau of Alcohol, Tobacco, 70 Firearms and Explosives, the Los Angeles Police Department, the National Crime Agency in the UK and the United 71 States Marine Corps and has also been used to identify source populations during biological invasions and sources 72 of infection during disease outbreaks (Le Comber et al. 2006; Raine et al. 2009; Le Comber et al. 2011; Stevenson et 73 al. 2012).

74						
75	The development of geographic profiling has – understandably – been driven by the need for practical solutions to					
76	the problems encountered by law enforcement agencies. O'Leary (O'Leary 2009; O'Leary 2010; O'Leary 2012)					
77	placed GP in a Bayesian framework, mathematically formalising the problem. However, the model put forward by					
78	O'Leary makes the simplifying assumption that all observed data points originate from a single source, and hence					
79	performs extremely badly in cases where there are actually multiple sources (see Methods and Results). Thus,					
80	despite the mathematical appeal of O'Leary's approach, the CGT algorithm continues to be widely used as a result of					
81	its proven track record (Rossmo 2000).					
82						
83	Here, we present a well-defined mathematical approach that unifies existing methods in a single framework.					
84	Crucially, our method explicitly deals with the issue of multiple sources – a situation typical of biological data sets,					
85	but less common in criminology. Under these circumstances, our model outperforms both the CGT algorithm and a					
86	simple Bayesian model based on the work of O'Leary (O'Leary 2010). Further, we develop a computational					
87	approach using Markov Chain Monte Carlo (MCMC) methods that extends the technique to large data problems.					
88	Finally, we demonstrate the effectiveness of our model using a real-life example of malaria cases in Egypt.					
89						
90	Specifically, we assert that (1) one of the reasons for the CGT algorithm's improved performance relative to the					
91	simple Bayesian model lies in its ability to deal with multiple sources; and hence by constructing a Bayesian model					
92	that incorporates the ability of the CGT algorithm to deal with multiple sources while maintaining the mathematical					
93	rigour of the simple Bayesian model, we can outperform both of the existing methods; (2) this method can be					
94	extended to large data problems using MCMC; (3) this method can be used to provide practical solutions to real-life					
95	problems, such as those found in epidemiology.					
96						

97 Geographic Profiling Models

98 The traditional (CGT) and Bayesian approaches to geographic profiling differ in both their construction and

99 implementation. In the following sections we specify each in common terms.

101 CGT algorithm

The traditional method begins by considering a distance-decay function around each individual data point. The height of the surface is a measure of how confident we are that the source location lies at this point. The decay function can take a number of forms, but in criminological applications it is typical to use a two-part distribution that increases to a maximum at a distance *B* from the data point, and then declines beyond this:

$$f(d) = \begin{cases} \frac{1}{d^h}, & \text{if } d > B\\ \frac{kB^{g-h}}{(2B-d)^g}, & \text{if } d \le B \end{cases}$$
^[1]

106

107 where d is the distance (either Euclidian or Manhattan) from the observation. This distribution was originally 108 proposed by Rossmo (2000), but here we have used the notation of O'Leary (O'Leary 2009; O'Leary 2010) 109 (correcting for a mistake in the direction of the inequalities). In this paper we use the Euclidean distance throughout. 110 Although this decay function is often referred to as a probability distribution, this is not technically true as there is 111 no requirement for the surface to integrate to unity (nor, in criminology, any need for it to do so, since the analysis is 112 used to produce ranked scores rather than probabilities). Thus, in the traditional method the decay function is better 113 described as a surface of search priority, subject to the more general constraint that points high up on the surface 114 represent areas of high priority. This measure of priority is modelled as an additive quantity, meaning that the 115 information from several observations can be combined by summing together the independent surfaces. The end 116 result of this process of summation is a single surface that represents our integrated knowledge of the source 117 location, which is referred to as a jeopardy surface (Rossmo, 2000).

118

The search efficiency of the model can be calculated using the hit score percentage; the proportion of the area that we must search before the true source location is found. The smaller the hit score percentage, the more accurate the geoprofile, with a hit score percentage of 50% representing what we would expect from a non-prioritised random or uniform search (see Rossmo 2000).

123

124 Simple Bayesian model

125 We compare the CGT algorithm against a simple Bayesian model based on the initial approach described by

126 O'Leary (O'Leary 2010; O'Leary 2012), and ignoring subsequent extensions relating to the choice of priors. This

127 approach differs from the CGT in that distributions are defined and manipulated according to the laws of 128 probability. The starting point is to write down the probability of the data, given the known location of the source. 129 This is achieved through the use of a probability distribution, which we will refer to as the migration profile, in 130 which the probability of finding an observation at any point in the domain is expressed relative to the location of the 131 source. Assuming independence between observations, the probability of the sample is simply the product over the 132 probabilities of the individual data points (in fact, Rossmo (1995) considered a similar formulation in which the 133 CGT algorithm is applied in log space). By placing a suitable prior on the source location and applying Bayes' rule it 134 is possible to derive the posterior distribution of the source location, given the observations.

135

136 Unsurprisingly, the choice of method makes a big difference to the results. While the CGT algorithm tends to create 137 a patchy distribution of peaks and troughs, entertaining the possibility of a number of different source locations, the 138 simple Bayesian method tends to place the majority of the posterior probability mass around the spatial mean of the 139 data points (at least for many choices of prior and likelihood, including those considered here). Another important 140 difference between the methods is in the rate of convergence. In the Bayesian approach the variance of the posterior 141 distribution tends to decrease rapidly as more data is added, whereas in the CGT method the variance of the 142 geoprofile can never be less than the variance of the decay function. Generally, when there is in fact a single source 143 location the Bayesian method is predicted to outperform the traditional method. However, if there is the potential for 144 multiple source locations then the Bayesian method is predicted to converge quickly on the wrong answer, while the 145 traditional method will still perform well. In this study, we test this prediction using a variety of simulations (see 146 Results 1 and 2, below).

147

148 The Dirichlet process mixture model

149 Our primary objective is to address the issue of multiple sources within a well-defined Bayesian framework. The

150 tool that allows us to do this is the Dirichlet Process Mixture (DPM) model, which has a strong mathematical

151 foundation (Ferguson 1983; Green & Richardson 2001) and is finding increasing application within biology (e.g.

152 Huelsenbeck et al. 2006; Huelsenbeck & Andolfatto 2007; Dorazio et al. 2008). Unlike many clustering approaches,

153 DPM models do not require the user to specify the number of clusters beforehand, making them extremely useful in

154 situations where there is no strong prior information about the exact number of clusters. In place of a fixed number

155 of clusters, the DPM model describes the process of cluster formation using a single 'concentration parameter', α . 156 Specifically, if we have already seen n observations, of which n_A came from cluster A, then the (prior) probability of 157 the next observation also belonging to cluster A is given by $n_A/(n + \alpha)$. It follows that, no matter how many observations we have seen, there is always a positive probability $\alpha/(n + \alpha)$ of the next observation originating from a 158 159 previously undiscovered cluster. While we may not believe there to be a truly unlimited number of clusters, by 160 allowing for the possibility of an expanding number of clusters we can ensure that our model is always appropriate 161 for the quantity of data at hand. Obviously the choice of the concentration parameter α has a strong influence on the 162 model. Although an appropriate value of α could be fitted from training data, here we chose instead to integrate over 163 our uncertainty by placing a diffuse hyper-prior over α (of the form $h(\alpha)=1/(1+\alpha)^2$, see Appendix 2 for details). 164 Where stronger prior information is available, the model can easily be adapted to include this.

165

The second part of the DPM model is the calculation of the posterior distribution of source locations, conditional on 166 167 a particular partition of the data into clusters. This part is mathematically very similar to the simple Bayesian model, 168 with the only difference being that a different posterior distribution is produced for each cluster. The likelihood of 169 all observations in the same cluster is equal to the product of the migration profile over each of the observations. By 170 incorporating an appropriate prior on the source location and applying Bayes' rule we arrive at the posterior 171 distribution of the source location from which this particular subset of observations derived. Carrying out this step 172 for each cluster independently we obtain a set of posterior distributions – one for each of the (potentially) multiple 173 source locations.

174

175 Finally, in order to obtain an analytical solution to the DPM model described above we would be required to sum 176 over all possible partitions of the *n* data points into up to *n* clusters, weighted by the posterior probability of the partition in each case. The number of such partitions is given by the n^{th} Bell number (B_n) which becomes 177 178 prohibitively large for values as low as n=10 (B₁₀=115,975). Thus, for any reasonably sized data set we must turn to 179 MCMC methods for a practical solution. Fortunately, a detailed exposition of MCMC algorithms for DPM models is 180 provided by Neal (2000), and we need only to adapt these algorithms to our specific application. A more detailed 181 description of the DPM model, including expressions relating to posterior inference under the analytical and MCMC 182 forms of the solution, is provided in Appendices 1 to 3.

8

184 It is important to emphasise that the DPM model can be adapted to use any migration profile that satisfies the laws 185 of probability (i.e. integrates to unity). The essence of the DPM model lies in the way that information is combined between clusters, and not in the specific details of the migration profile used. This can be seen in the logic of our 186 187 study, which has four parts. (i) First, when comparing directly the CGT, simple Bayesian, and DPM models, we use 188 the distribution from the CGT (described in equation [1]) as our migration profile in all three approaches. This 189 ensures that the only difference between methods lies in the way that information is being combined, and not in any 190 other assumptions relating to migration. (ii) Next, we validate the MCMC version of our proposed solution using 191 this same migration profile, thereby ensuring that our MCMC results are directly comparable with our analytical 192 results. (iii) From this, we move on to consider simulated data generated from a distribution more typical of those 193 assumed in biology – the normal distribution – and explicitly compare the full form of the DPM model with the 194 CGT under this assumption. (iv) Finally, we examine a real-world data set – an outbreak of malaria in Cairo – using 195 all three models.

196

197

198 Methods(i) Comparing the simple Bayesian, CGT and DPM models

199 As mentioned above, our first task is to compare the simple Bayesian, CGT and DPM models purely in terms of the 200 way that information is combined in each case, and controlling for any differences between models, such as the 201 migration profile. We simulated 6, 7, 8 or 9 data points from the distribution given in equation [1] (B=0.5, f=4, g=4), 202 emanating from either 1, 2 or 3 sources, truncated them to fit the available grid. For the purposes of simulation we 203 split the domain into a 100*100 grid, and replicated each combination of the number of data points and sources 1000 204 times. Sources were chosen to fall within the central 50*50 cells in a random, uniform manner. For each simulated 205 data set we then used each of the three methods described above to search for the 'unknown' source locations, with 206 search efficiency being measured in terms of the hit score percentage. The same distribution (distribution [1] with B=0.5, f=4, g=4) was used as the search distribution in each of the three methods. By designing simulations in this 207 208 way we can capture an idealised situation in which all three methods make the same assumptions about the true 209 dispersal distribution, and furthermore these assumptions are exactly correct (thereby removing another possible 210 source of model error).

9

212 (ii) MCMC validation

- 213 For the reasons described previously, the analytical form of the DPM model can deal with only small data sets, and
- 214 for larger data sets an MCMC implementation of the solution is required. For each of the 12000 simulations
- described above (1000 replicates of each combination of 1, 2 and 3 sources and 6, 7, 8 or 9 data points), we also
- 216 used an MCMC implementation of the model, and calculated the correlation between the surface produced by the
- analytical form of the model and the MCMC form (see Appendix 3 for details of the MCMC algorithm). We also
- repeated the comparison of the DPM model with the CGT for larger data sets (1, 2 and 5 source locations; 20, 40,
- 219 60, 80 and 100 spread points), using just the MCMC implementation of the model.
- 220

When running the MCMC, multiple chains were run simultaneously, with convergence being assessed using the Gelman-Rubin (GR) diagnostic statistic (Gelman et al. 2003) evaluated on the concentration parameter α (using a value of GR=1.1 as a threshold for convergence). After the burn-in period, samples were obtained until the largest standard error of any point on the estimated surface was less than 0.01. Samples were not thinned, as it has previously been shown that this does not increase statistical power in situations such as this (Link & Eaton 2012).

227 (iii) Further comparison of the CGT and DPM models

The migration profile used above (distribution [1]) was designed for criminological applications. In some cases, including many biological applications, it may be more appropriate to assume alternative migration profiles. Here, we assume a bivariate normal migration profile, centred on the unknown source location(s), and with variance σ^2 . In some cases, there will be biological data on dispersal patterns that can be used to inform the choice of σ ; for example, studies have shown that most malaria transmission occurs close to the larval breeding sites – usually between a few hundred meters and a kilometer– and rarely exceeds 2-3 km (Carter et al. 2000).

234

We are also required, as part of the DPM model, to choose a prior on the source location(s). For the sake of simplicity we use an empirical Bayes approach, assuming a bivariate normal prior, centred on the spatial mean of the observed data, and with variance τ^2 , where τ was set to the maximum distance in either latitude or longitude between the crime sites. τ equals one standard deviation of the normal prior; hence, we expect our source to lie within this distance of the centre around two-thirds of the time, and the model allows for sources well outside the area bounding the crimes. Hence, there is a diffuse, non-informative prior over and beyond the normal search area.

We simulated 6, 7, 8 or 9 data points from a bivariate normal distribution with standard deviation sigma = 1 and emanating from either 1, 2 or 3 sources. For the purposes of simulation we split the domain into a 100*100 grid, and replicated each combination of the number of data points and sources 1000 times. For each simulated data set we then used the two best performing methods described above (CGT and DPM) to search for the 'unknown' source locations, with search efficiency being measured in terms of the hit score percentage. The CGT uses the distribution describe in equation [1] with parameters fitted from the data as described by Rossmo (2000), while the DPM uses the spatial mean to fit phi, with sigma fixed at 1.

249

250 (iv) Case study

251 We tested the performance of our model in a real world example by using the MCMC implementation of the DPM 252 model to reanalyse data from Le Comber et al. (2011). In this study, spatial data relating to 139 recorded 253 Plasmodium vivax malaria cases were collected, and buffer zones of 2 km were created around the locations of these malaria cases and merged to form a polygon of 296.5 km² (Hassan 2006). All accessible aquatic habitats within this 254 255 study area (surface/cryptic; temporary/semipermanent/permanent) were located and characterised between April and 256 September 2005. These included water tanks, water pools created through pipelines or drainage system breakage, 257 seepage from slum housing, natural springs, pools and ditches filled with ground water. Water sources included in 258 this analysis were identified as bodies of water harbouring at least one mosquito larva over the study period (n = 59). 259 A total of 11 mosquito species were identified, including the malaria vectors An. sergentii and An. pharoensis, as 260 well as other, non-vector, species. Of these 59 sites, seven tested positive for one or both of the malaria vectors An. 261 sergentii and An. pharoensis (An. sergentii is well established as the most dangerous malaria vector in Egypt (Said et al. 1986)). 262

264	A dispersal distance of sigma = 0.018, roughly corresponding to 1km, was used in the DPM model in
265	correspondence with values in the literature (e.g. Carter et al. 2000) and a value of tau = 0.328 was fitted from the
266	observed data (see above).
267	
268	The model is written in R (R core team 2012) and integrates with Google Maps via the R package RgoogleMaps
269	(Loecher 2012). The model used in this paper is available from the authors on request as an R package called
270	'Rgeoprofile'.
271	
272	
273	Results
274	(i) Comparing the simple Bayesian, CGT and DPM models
275	
275	Starting with the first set of simulations (1000 replicates of each combination of 1, 2 and 3 sources and 6, 7, 8 or 9
276	data points), we used a fully factorial ANOVA to test the effect on the hit score percentage (or average hit score
277	percentage when the number of sources was > 1) of model type, number of sources and number of spread points.
278	Three model types were examined; the analytical form of the DPM model, the classical CGT algorithm and the
279	simple Bayesian model.
280	
281	Model type, number of points and number of sources all significantly affected the relative performance of the three
282	models (ANOVA: model type: F _{2,35964} =4787.05,p< 2e-16; sources: F _{2,35964} =13099.30,p<2e-16; points: F ₃ ,
283	$_{35964}$ =106.23, p<2e-16). All interactions were highly significant, with the <i>F</i> value for model type*sources interaction
284	having the largest effect size ($F_{4,35964}$ =2840.12, p<2e-16); none of the other F values exceeded 52. Tukey post-hoc
285	tests at α =0.05 showed that (1) the CGT significantly outperformed the simple Bayesian model, by an average of
286	1.81% (95% CI: 1.75-1.86%); (2) the DPM model showed a statistically significant improvement over both the CGT
287	algorithm, albeit only by 0.3% (95% CI: 0.25-0.36%) and the simple Bayesian model, again by about 2% (95% CI:
288	2.1-2.2%). Across all 12,000 runs, the DPM model performed better than the CGT in 68.2% of trials, and as well or
289	better in 74.9%, and better than the simple Bayesian model in 64.6% of trials, and as well or better in 91.5%.

- However, although the DPM model outperformed the simple Bayesian model overall, the simple Bayesian modelhad a small advantage when there was a single source (Figure 1).
- 292

293 (ii) MCMC validation

- 294 For the same simulated data sets described above we calculated the correlation between the surface produced by the
- analytical form of the DPM model and the MCMC form. The two surfaces tended to extremely highly correlated (r
- 296 $(\text{mean} \pm \text{sd}) = 0.9998 \pm 0.0010$, demonstrating that the MCMC algorithm does indeed find the same or at least

297 extremely similar – posterior distributions as the analytical form of the model.

- 298
- For the second set of simulations (1000 replicates of each combination of 1, 2 and 5 sources and 20, 40, 60, 80 or
- 300 100 data points) we performed the same analysis as in Results part 1, with extremely similar results (ANOVA:

301 model type: $F_{1,29992}=167.7$, p<2e-16; sources: $F_{2,29992}=10603.1$, p<2e-16; points: $F_{4,29992}=1986.2$, p<2e-16; model

302 type*sources: $F_{2,29992}$ =463.5, p<2e-16; model type*points: $F_{4,29992}$ =17.4, p<2e-16; sources*points: $F_{8,29992}$ =2916.7,

303 p<2e-16; model type*sources*points: $F_{8, 29992}=0.9$, p=0.87). Tukey post-hoc tests at $\alpha = 0.05$ showed that the DPM

304 model outperformed the CGT algorithm in a statistically significant way; again, this improvement was most marked 305 when the number of sources was > 1 (Figure 2).

306

307 (iii) Further comparison of the CGT and DPM models

308 In the next set of simulations, in which a normal migration profile was assumed, we used ANOVA to test the effect 309 on the hit score percentage (or average hit score percentage when the number of sources was > 1) of model type,

310 number of sources and number of spread points. The two best performing model types from previous simulations

311 were examined; the CGT and the DPM.

- 313 The best performing ANOVA was selected by AIC to include a single significant interaction term. Model type,
- 314 number of points and number of sources all significantly affected the relative performance of the two models
- 315 (ANOVA: model type: $F_{19991}=3693.6, p \le 2e-16$; sources: $F_{2,19991}=2038, p \le 2e-16$; points: $F_{3,19991}=39.1, p \le 2e-16$).
- 316 Model type*sources interaction was also significant ($F_{4,19991}$ =222.1, p<2e-16). Tukey post-hoc tests at α =0.05

showed that the DPM model showed a statistically significant improvement over the CGT algorithm with an effect
size of 4.1% (95% CI: 3.9-4.2%). The MCMC implementation of the DPM outperforms the CGT 67.1% of the time,

and performs as well or better 67.2% of the time. In our simulations this equates to searching on average 410 fewer

320 cells (95% CI: 394-421) before finding all of the sources.

321

322 (iv) Case study

The median hit score percentages for the seven vector breeding sites identified in Hassan (2006) were 0.34% for the DPM model, compared to 0.43% for the CGT and 1.2% for the simple Bayesian model. Note that the hit scores reported here differ from those in Le Comber et al. (2011), although the surface produced is the same in both cases. The difference arises because the DPM model uses RgoogleMaps (Loecher 2012), and thus the exact dimensions of the search area (which affects the hit score) are set by the available zoom levels in the Google Maps data. To allow direct comparison, we used the same search area for the CGT and the DPM mode.

329

330 For five of the seven sites, hit score percentages for the DPM were less than half a per cent. An additional output of 331 our model is that it can provide a barplot of the posterior probability of the number of realised sources (Figure 3). In 332 this case our model indicated the highest probability for seven sources, with a likely range of 6-10. Interestingly, 333 some of these correspond to areas where no vector species were found by Hassan (2006) (Figure 4). One possibility, 334 of course, is that these are false-positive results. Alternatively, it is possible that some sources were missed in the 335 original survey, especially given the often considerable difficulty of locating small, transient breeding populations of 336 mosquitoes (Carter et al. 2000) and since searches were carried out in a single year (2005), whereas the malaria 337 cases spanned four (2001-2004) (Hassan 2006; Le Comber et al. 2011).

338

339 Discussion

Overall the DPM model is an improvement on the existing methods. When the number of sources is greater than one it outperforms them (Results (i)), it does not require that the number of sources is known *a priori* and, in addition, it generates estimates of their number. Even in conditions specifically designed to maximise the performance of the CGT algorithm, the DPM model still obtains a small advantage, reflecting the way in which it appropriately combines information from observations, rather than taking a simple sum (as in the CGT) or product (as in the simple Bayesian model). The DPM model's analytical method cannot be extended to very large numbers of observations, but the approach can be implemented in an MCMC algorithm which accurately constructs the posterior distribution, as demonstrated in Results (ii).

348

With these facts established we move on to consider cases in which the DPM model may have a practical advantage over other approaches. The later set of simulations (Methods (iii) and Results (iii)) demonstrate that there are biologically plausible settings in which the use of the DPM model can result in an appreciable increase in search efficiency compared with other methods. Finally, and perhaps most encouragingly, we find that the DPM model leads to an increase in search efficiency when applied to a real-world data set describing malaria transmission in Cairo. The improvement over the CGT algorithm is small, but justifies further investigation of this model on a range of data sets.

356

357 In its construction, the DPM model forms a bridge between the seemingly disparate methodologies of the CGT and 358 the simple Bayesian approach to geographic profiling. From a practical point of view the major difference between 359 the two existing approaches lies in whether distributions should be summed (CGT) or multiplied (simple Bayesian). 360 The DPM model works by splitting the data into groups, with each group corresponding to a different source 361 location. The laws of probability then dictate that distributions should be multiplied within groups, but summed 362 between groups. Thus, if all points are assigned to a single source we arrive back at the simple Bayesian model, 363 while if all points are assigned to different sources we arrive at something more akin to the CGT algorithm. In this 364 context, our concentration parameter α can be understood as a prior over the complete spectrum of models, which 365 allows us to transition between a single-source model and a multiple-source model. When α is set to zero, the DPM 366 model becomes mathematically equivalent to the simple Bayesian model; conversely, as α tends to infinity, we 367 converge on the CGT algorithm. In the majority of cases – particularly those dealing with biological data – the most 368 likely explanation for the data will often lie between these two extremes. For example, in the malaria analysis, the 369 DPM model assigned the highest probability to seven sources from 139 disease case locations (Figure 3). 370

In our simulations, the DPM model outperformed both other approaches when there were multiple sources. In cases
with a single source – a common scenario in criminology – the improvement over the CGT, although statistically

373 significant, was minimal when the dispersal distribution was drawn from Equation [1] (when this assumption was 374 relaxed, the improvement was more marked). The comparison between the DPM model and the simple Bayesian 375 model shows that latter has a small advantage when there is a single source. However, when there is more than one 376 source, the DPM shows a large improvement (this is perhaps unsurprising, since the simple Bayesian model assumes 377 that there is a single source). In real-world applications of GP models it will often (perhaps even always) be the case 378 that the true number of sources is unknown, therefore the principal advantage of the DPM model lies in its ability to 379 rigorously handle the problem of multiple sources. In fact, since the difference between the simple Bayesian model and the DPM model is small when there is a single source, and the advantage offered by the DPM model when there 380 381 are multiple sources is larger, we would argue that the DPM model is preferable in real-world applications of GP. In 382 our simulations, the DPM model outperformed both other approaches in cases with multiple sources. In cases with a 383 single source – a common scenario in criminology – the improvement over the CGT, although statistically 384 significant, was minimal when the dispersal distribution was drawn from Equation [1] (when this assumption was

385 relaxed, the improvement was more marked).

386

387 However, formulating the problem in a rigorous Bayesian framework also allows for a number of useful extensions. 388 First, our model produces a true probability surface, allowing us to calculate the marginal probability of different 389 numbers of sources, as in Figure 3. Second, we can produce a probability surface conditional on a particular number 390 of sources, thereby allowing us to break the overall picture down into different scenarios (we can imagine a different 391 search strategy, conditional on there being one source, two sources etc.). Third, the DPM model explicitly calculates 392 the posterior probability under the model that a particular observation is derived from a particular source. This may 393 be of interest in criminology, where crime linkage is an important problem (Rossmo 2000), and may also be useful 394 in biological data sets, where the spatial linkage can be validated against other forms of information (for example 395 genetic data).

396

So far, the DPM model is constructed with flexibility in mind, rather than statistical power. For particular cases it may be possible to increase the power of the model by incorporation of stronger prior information – for example, by inferring the concentration parameter from training data. Similarly, where empirical evidence has shown that nonnormal dispersal profiles are appropriate (for example, Cauchy distributions in some bird species (Winkler et al. 2005; VanHoutan et al. 2007) or bivariate Student's t-distributions in seeds (Nathan & Muller-Landau 2000)), these
can be used within the same general framework.

403

As well as producing a range of new outputs, the DPM model could also be extended to incorporate new inputs. For example, one useful possible extension of our approach is the utilisation of the outputs produced by niche models to generate priors in the DPM model. Niche modelling is a well-developed field that has recently been placed on a Bayesian footing (Elith & Leatherwick 2009), making its incorporation into the DPM model relatively straightforward. A Bayesian niche model produces a probabilistic estimate of the suitability of habitat for the organism being studied that can be used as a prior in the DPM model. Combining these two approaches would go some way towards producing a spatially explicit niche model approach, as called for by Peterson et al (2003).

411

412 In epidemiology and invasion biology, much more attention is paid to models that run forwards in time to generate 413 risk maps or forecasts of future incidence than those that run backwards to locate sources. GP, on the other hand, is 414 radically different, running backwards in time to use current locations to infer sources (Le Comber & Stevenson 415 2012). The DPM model structure described above also differs from many spatially explicit epidemiological models, 416 such as the shot noise Cox process (Møller 2003), in assuming a distribution of point sources, rather than a smoothly 417 varying hazard function over space. This feature also distinguishes the DPM approach from many existing methods 418 that are routinely used to detect clusters in ecological and epidemiological data (see Pullan et al. 2012 for a review). 419 The impact that these different modeling assumptions may have on our conclusions should be explored in further 420 work. In fact, as O'Leary (O'Leary 2010; O'Leary 2012) has shown, a fully Bayesian implementation of GP can 421 easily be extended to run forwards in time. Despite the difficulties faced by all predictive models, this could potentially be important in areas of biology including epidemiology, invasion biology and in conservation biology 422 423 (e.g. planning reintroductions of animals or plants).

424

The DPM model we present here is a general method that can be applied to data describing spread from common source. Evidence-based targeting of interventions is a crucial component in the fight against infectious disease, and targeted interventions are more efficient and more cost-effective than untargeted interventions; for example, malaria is strongly dependent on the location of vector breeding sites, and most transmission only occurs within short

429	distances of these sites (Carter et al. 2000). Because of this clustering, untargeted intervention is highly inefficient.					
430	In the Cairo study, the DPM model identified five of the seven breeding sites in less than half a percent of the total					
431	search area, representing a dramatic improvement over a non-targeted search.					
432						
433	Although our implementation of the DPM model can deal with large data sets (>1000 data points), GP methods also					
434	work well with very small data sets (Rossmo 2000; Stevenson et al. 2012), allowing their use in the early stages of					

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likely to lead to improved targeting of interventions, and more efficient use of resources.

an outbreak or invasion, when control efforts are most likely to be successful. The DPM model provides a useful

practical tool for conservation biologists and epidemiologists, offering improvements over other methods that are

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Figure 1 Comparison of the analytical form of the DPM model against (A) the simple Bayesian model, and (B) the CGT algorithm, expressed as the hit score percentage of the simple Bayesian model minus the hit score percentage of the DPM model, and the hit score percentage of the CGT algorithm minus the hit score percentage of the DPM model, respectively. Thus, points above the red line indicate cases in which the DPM model outperformed the other models. In both cases, the DPM model has a statistically significant advantage, although this is more pronounced for the comparison with the simple Bayesian model. In both comparisons, the relative performance of the DPM model improves as number of sources increases.





Figure 2 Comparison of the MCMC implementation of the DPM model against the CGT algorithm, expressed as
the hit score percentage of the CGT algorithm minus the hit score percentage of the DPM model. Again, points
above the red line indicate cases in which the DPM model outperformed the other model. The DPM model
outperformed the CGT algorithm, especially as number of sources increases.



578 Figure 3 Marginal likelihood of different numbers of realised infection sources for the Cairo data. The DPM model

579 estimates that there are 6-10 sources, and assigns the highest likelihood to seven sources.





Figure 4 Geoprofile from 139 *Plasmodium vivax* cases in Cairo, Egypt, using (A) the simple Bayesian model; (B)
the CGT algorithm; (C) the DPM model. (D) shows a close-up of the DPM surface. In all cases the observed data



points are shown as black circles, while the empirically identified sources are shown as blue squares.