Bayesian analysis of the scatterometer wind retrieval inverse problem: some new approaches

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Summary. The retrieval of wind vectors from satellite scatterometer observations is a non-linear inverse problem. A common approach to solving inverse problems is to adopt a Bayesian framework and infer the posterior distribution of the parameters of interest given the observations using a likelihood model relating the observations to the parameters, and a prior distribution over the parameters. In this paper we show how Gaussian process priors can be used efficiently with a variety of likelihood models, using local forward (observation) models and direct inverse models for the scatterometer. We present an enhanced Markov Chain Monte Carlo method to sample from the resulting multi-modal posterior distribution.

We go on to show how the computational complexity of the inference can be controlled using a sparse, sequential Bayes algorithm for estimation with Gaussian processes. This helps to overcome the most serious barrier to the use of probabilistic, Gaussian process methods in remote sensing inverse problems, which is the prohibitively large size of the data sets. We contrast the sampling results with the approximations found using the sparse, sequential Bayes algorithm.

1. Introduction

Satellite borne scatterometers have been in existence for several decades; retrieving wind vectors from scatterometer observations is a complex inverse problem. In brief, scatterometers emit a pulse of micro-wave radiation which travels largely unmodified through the atmosphere. When it meets the ocean surface it interacts with water waves of a similar wavelength of around 5 cm for the C-band radars used in this study. The strength of the backscattered signal depends on the magnitude and relative orientation of the water waves to the incoming radar beam. The water wave field at centimetre wavelengths is driven by the instantaneous surface wind stress, which in turn is related to the near surface (usually taken at 10 m height) wind vector. The physics of the problem is complex and not fully understood (Janssen et al., 1998). So far, most solutions to this inverse problem have been based on locally inverting an empirical sensor model for the scatterometer observation process (Stoffelen and Anderson, 1997a), often using look up tables (Offiler, 1994). In this paper we propose a statistically based retrieval method for the scatterometer inverse problem, and demonstrate how an innovative sparse, sequential Bayes algorithm can be used to significantly speed up the solution of this inverse problem. We apply our methods to both data assimilation, where we have an informative prior estimate of the wind field from a dynamical model, and, data retrieval, with a zero mean prior.

The paper is organised as follows: in Section 2 the methods for obtaining a local solution to the inverse problem are described. Section 3 shows how the prior vector Gaussian process models are constructed and Section 4 how the priors are combined with the local solutions to provide an a posteriori solution to the inverse problem, using sampling and a variational
approximation (sequential Bayes). The results are shown and discussed in Section 5 while conclusions are drawn in Section 6.

1.1. The aim of scatterometry
Obtaining wind vectors over the ocean is important to numerical weather prediction (NWP) since the ability to produce a forecast of the future state of the atmosphere depends critically on knowing the current state accurately (Haltiner and Williams, 1980). The act of obtaining estimates of the current state of the atmosphere using observations and dynamical models is referred to as data assimilation. However, the observation network over the oceans (particularly in the Southern Hemisphere) is very limited (Daley, 1991). Thus it is hoped that the global coverage of ocean wind vectors provided by satellite-borne scatterometers (Offiler, 1994) will improve the accuracy of weather forecasts by providing better initial conditions for NWP models (Lorenc et al., 1993). The scatterometer data also offer the potential of improved wind climatologies over the oceans (Levy, 1994) and the possibility of studying, at high resolution, interesting meteorological features such as cyclones (Dickinson and Brown, 1996).

1.1.1. Scatterometers
This study uses scatterometer data from the ERS-2 satellite; the on-board vertically polarised microwave radar operates at 5.3 GHz and measures the backscatter from gravity-capillary waves on the ocean surface. Backscatter from the ocean surface is measured by the normalised radar cross section, denoted by $s_o$, and has units of decibels. A 500-km wide swathe is swept by the satellite to the right of the track of its polar orbit, as illustrated in Figure 1. There are 19 cells sampled across the swathe, and each cell has dimensions of roughly 50 by 50 km, which implies that there is some overlap between cells.

Each cell is sampled from three different directions by the ‘fore’, ‘mid’, and ‘aft’ radar beams, giving a triplet, $s^o = (s^o_f, s^o_m, s^o_a)$. This triplet $s^o$, is related to the average wind vector $v$ within the cell (Offiler, 1994) with a dependence on the known incidence and azimuth angles of the beams, which vary across the swathe. We assume that any unmodelled effects are largely related to wind speed and thus their impact is implicitly included in the empirical models which have been developed (Cornford et al., 2001; Bullen et al., 2003). Other geophysical parameters such as rain and sea ice are also believed to have a small effect...
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1.2. Bayesian framework

The inverse problem to be solved is the retrieval of the wind field, \( V = \{ v_i \}_{i=1,N} \), given the scatterometer observations, \( S^o = \{ s^o_i \}_{i=1,N} \), where \( N \) is the number of observations. Since the processes involved include noise, it is necessary to estimate the conditional probability density function, \( p(V | S^o) \). In the usual way for Bayesian approaches to the solution of inverse problems (Tarantola, 1987) the likelihood, \( p(S^o | V) \), factorises as \( \prod_i p(s^o_i | v_i) \), that is the errors on the scatterometer observations are assumed independent, and the posterior distribution is:

\[
p(V | S^o) = \frac{p(S^o | V)p(V)}{p(S^o)} \propto \left( \prod_i p(s^o_i | v_i) \right) p(V)
\]

We show two approaches to the inference of \( p(V | S^o) \) using forward (likelihood) models, based on hybrid Monte Carlo sampling (Duane et al., 1987) and the sequential Bayes algorithm. We also show how it is possible to use direct local inverse models which can improve the speed of the methods.

In the discussion to this point \( p(V) \) has remained undefined. A natural probabilistic representation for wind fields, which links the local models, is to use a vector Gaussian process:

\[
p(V) = \frac{1}{(2\pi)^{N/2} |K_v|^{1/2}} \exp \left( -\frac{1}{2} V' K_v^{-1} V \right)
\]

where \( K_v \) is the joint covariance matrix for \( V \). The parameterisation of the prior wind field model is discussed in Section 3.

2. Non-Bayesian approaches: local solution

The initial problem which must be addressed is the local (that is marginal) inversion of the observation operator. As discussed in Section 1 the forward or likelihood model, which maps the state variables that are to be recovered (\( v \)) to the observables (\( s^o \)) is not perfectly known. Physics based models, such as VIERS1 (Janssen et al., 1998), have been shown to provide reasonable approximations to observed data. Thus empirical approaches have been developed, which exploit some of the prior knowledge from the physics of the observation processes, by using a truncated Fourier model structure (Stoøelen and Anderson, 1997b).

In Section 2.1 we outline the neural networks used to implement the forward model, while Section 2.2 describes the use of mixture density networks to construct local direct inverse models.

A significant issue when dealing with scatterometer data comes from recognising that this data provides information on the wind vector over a 50 by 50 km region of the ocean. There are no other direct measurements of \( v \) at this scale. At best there may be a single buoy measurement somewhere within the region, but there is no guarantee this will be representative. The best approximation to the average wind vector can be derived from NWP models. These ‘pseudo-observations’ are derived from numerical forecasts of the dynamics of the atmosphere using very complex models. These models integrate coupled
Fig. 2. (a) An approximate cross section through the 3D cone \( s_{o_i} = f(v_i) \) which defines the forward model. The lighter part is the upwind segment, the darker the downwind segment. A satellite observation is plotted as \( \times \), and corresponding closest points on the model manifold \( f(v_i) \), are marked \( \ast \). Also shown are samples from the noise distribution (small grey dots), thus giving an idea of instrumental noise. (b) A contour plot of the local conditional probability density function, \( \log[p(s_{o_i} | v_i)] \), as a function of \( v_i = (v_1, v_2) \). The NWP wind vector is marked \( \ast \) and the two humps correspond to the two closest points on the manifold in (a).

2.1. Forward models

In recent work (Bullen et al., 2003) we developed a forward model for a single scatterometer radar beam based on the combination of a radial basis function network and a truncated Fourier series:

\[
s_o = a_0 + a_1 \cos(\chi) + a_2 \cos(2\chi) + a_3 \cos(3\chi) + a_4 \cos(4\chi) + \epsilon ,
\]

where \( a_0, a_1 \ldots a_4 \) are the outputs from the radial basis function network, \( \chi \) is the wind direction relative to the satellite azimuth angle and \( \epsilon \) is the additive error. The inputs to the radial basis function network are wind speed and beam incidence angle, and wind speed and relative direction are derived from the wind vector, \( v \). The parameters of the network were estimated in a Bayesian framework which accounted for “input” noise on \( v \) and utilised a radial basis function network to approximate the noise variance, again fitted in a Bayesian framework (Bishop and Qazaz, 1997). The forward, or likelihood, model
provides an estimate of \( p(s^o \mid v) \), with the assumption that the errors on each beam are uncorrelated, and can be locally inverted (using non-linear optimisation) to retrieve \( v \), however care must be taken due to the presence of multiple local solutions, resulting from the Fourier form of the forward model and the observation noise, as illustrated in Fig. 2.

This ambiguity can be understood when imagining looking at waves side on (small gravity waves are almost totally symmetric). With a static image it is almost impossible to tell the direction of motion; this intuition carries over well into the microwave region of the spectrum. When observation noise is also taken into account there are typically between 2 and 4 local solutions, thus any inversion process must recognise this: a simple gradient based optimisation will only discover a single mode. In operational practice, multiple initialisations are used to determine all \( v \) which maximise \( p(s^o \mid v) \), and the solutions thus determined are used by meteorological agencies as inputs to their systems for estimating the current state of the atmosphere (Offiler, 1994).

For practical applications, where many thousands of scatterometer observations must be processed quickly, the non-linear optimisation from multiple starts is prohibitively expensive, and only provides the most probable \( v \), not \( p(s^o \mid v) \), which is desired. In some situations it is possible to design direct local inverse models (Ward and Redfern, 1999) but in this case the presence of multiple solutions for a given \( s^o \) observation precludes a simple approach.

### 2.2. Direct inverse models

Repeating the application of Bayes theorem to the marginal likelihood at each observation, it is possible to rewrite eq. (1) to use direct local inverse models, \( p(v \mid s^o) \), by noting that:

\[
p(s^o \mid v_i) \propto \frac{p(v_i \mid s^o_i) p(v_i)}{p(v_i)} 
\]

To model the multi-modal \( p(v_i \mid s^o_i) \) we have employed mixture density networks (Bishop, 1995), while the marginal, \( p(v_i) \), is the marginal from the prior model eq. (2).

Mixture density networks are conditional mixture models,

\[
p(v_i \mid s^o_i) = \sum_{k=1}^{K} \alpha_k(s^o_i) \phi_k(v_i \mid s^o_i),
\]

where in this work \( \phi \) is a Gaussian,

\[
\phi_k(v_i \mid s^o_i) = \frac{1}{2\pi \sigma_k^2(s^o_i)} \exp \left[ - \frac{||v_i - \mu_k(s^o_i)||^2}{2\sigma_k^2(s^o_i)} \right],
\]

and the parameters of the mixture model (mixing coefficients, \( \alpha_k \), means, \( \mu_k \) and variances, \( \sigma_k^2 \)) are functions of the outputs of a neural network, which in this case has inputs \( s^o_i \) (Evans et al., 2000). The mixing coefficients are constrained to sum to one to ensure the output is a valid probability distribution. In previous work (Evans, 2001) the mixture density networks were fitted directly using NWP data, however it is impossible to separate the contribution of the uncertainty deriving from the use of noisy NWP estimates and the uncertainty coming from the scatterometer observation noise. In this paper we use a mixture density network with parameters estimated using data generated from the forward model (Section 2.1) with simulated observational noise.
This direct inverse model efficiently provides an estimate of $p(v_i | s_{oi})$ using a single forward propagation, which is a significant speed improvement over the non-linear optimisation based methods. Look up table based methods, as used in many meteorological agencies, are of a similar speed but using mixture models will approximate the distribution of $v_i$.

### 3. Vector Gaussian process priors

A stationary vector Gaussian process model is used to represent wind fields, based on the decomposition of a vector field into purely divergent and purely rotational flow, known as Helmholtz’s theorem (Daley, 1991). We do not give details here, but the full development can be found in (Cornford, 1998). Essentially, a scalar modified Bessel (or Matérn) covariance function based Gaussian process (Cressie, 1993; Handcock and Wallis, 1994) was applied independently to both the stream function, $\Psi$, and velocity potential, $\Phi$, rather than directly to the wind vector components. The wind vector can be written in terms of derivatives of $\Psi$ and $\Phi$,

$$v = \nabla \times \Psi + \nabla \Phi,$$

and thus the covariance functions for the wind vector components are defined in terms of second order derivatives of the $\Psi$ and $\Phi$ covariances

$$C_{lt}(r) = -\frac{1}{r} \frac{d}{dr} C_{\Psi\Psi} - \frac{1}{r} \frac{d^2}{dr^2} C_{\Phi\Phi} - \frac{1}{r} \frac{d}{dr} C_{\Psi\Phi},$$

$$C_{tt}(r) = -\frac{1}{r} \frac{d^2}{dr^2} C_{\Psi\Psi} - \frac{1}{r} \frac{d}{dr} C_{\Phi\Phi} - \frac{1}{r} \frac{d}{dr} C_{\Psi\Phi},$$

$$C_{lt} = C_{tl} = 0,$$

where $r$ is the distance between two observations. Thus given $C_{\Psi\Psi}$ and $C_{\Phi\Phi}$ we can compute the wind covariances, which are not isotropic in general, based on simple scalar isotropic covariance models for $\Psi$ and $\Phi$. Computing the covariances in this way ensures that the joint covariance matrix, $K_v$, is positive definite:

$$K_v = \begin{bmatrix} C_{uu} & C_{uv} \\ C_{vu} & C_{vv} \end{bmatrix} = \begin{bmatrix} \cos(\phi) & -\sin(\phi) \\ \sin(\phi) & \cos(\phi) \end{bmatrix} \begin{bmatrix} C_{lt} & 0 \\ 0 & C_{tt} \end{bmatrix} \begin{bmatrix} \cos(\phi) & \sin(\phi) \\ -\sin(\phi) & \cos(\phi) \end{bmatrix}$$

where $\cos(\phi)$ is a square matrix of the cosine of the angle (relative to the x-axis) between each observation pair. This formulation allows control over the ratio of divergence (given by the variance of $\Psi$) to vorticity (given by the variance of $\Phi$) in the resulting wind field.

The parameters of the zero mean modified Bessel covariance function are estimated using a sector of the North Atlantic from $52.5^\circ W$, $40^\circ N$ to $10^\circ W$, $60^\circ N$, since this is the most reliably observed ocean region. A large amount of data from the European Centre for Medium range Weather Forecasting (ECMWF) is used; this consists of gridded analysis data on a regular 2.5 degree latitude-longitude grid. In order to obtain reliable climatological estimates of the parameters in the wind field model, three complete years of data (1995–97) were used. To account for seasonality, a separate prior wind field model is fitted for each month, using the three years data, thus the prior is strictly tuned to the North Atlantic region. The prior distribution of the covariance parameters are chosen by expert assessment. The maximum a posteriori probability values of the parameters are used in subsequent experiments. These values suggested that the wind fields are once differentiable. Thus
the modified Bessel covariance function simplifies to a polynomial-exponential covariance function which has the form:

\[ C_\cdot(r) = E^2 \left(1 + \frac{r}{L} + \frac{r^2}{3L^2}\right) \exp\left(-\frac{r}{L}\right) + \eta^2 \]  

(6)

where \(L\) is a characteristic length scale parameter, \(E^2\) is the process variance and \(\eta^2\) is the noise variance. This form is much quicker to compute and is used for this reason. In the text we treat the maximum \textit{a posteriori} probability parameters of the Gaussian process as known and fixed since the posteriors for all but the differentiability parameters were strongly peaked around the maximum \textit{a posteriori} probability values. While not in the fully Bayesian spirit this renders the problem more tractable in real time.

4. Bayesian approaches

In this section we introduce the practical methods which are applied to approximate the full posterior distribution from eq. (1), based on sampling and a sequential Bayes algorithm.

4.1. Sampling

We turn our attention to drawing samples from the posterior distribution but to improve speed we replace the product term in eq. (1) using eq. (4) to give:

\[ p(V_j | S_0) \propto \left( \prod_i p(v_i | s_{i0}) \right) p(V) \]  

(7)

The likelihood term involves a scaled mixture model, thus the posterior cannot be computed analytically due to the factorial growth in complexity. We apply Markov Chain Monte Carlo (MCMC) techniques in order to explore the posterior distribution. MCMC techniques are powerful when applied to such distributions as they make no assumptions about the form of the distribution.

Earlier work has shown that the posterior distribution is multi-modal, and typically it is dominated by two well separated modes (Evans, 2001). Sampling from the posterior distribution is thus implemented by an extension of the Metropolis-Hastings algorithm (Metropolis et al., 1953; Hastings, 1970) incorporating knowledge of the dominating modes located in a mode finding exercise.

Due to the number of local modes an exhaustive search is impractical. Instead, a mixture of probabilistic and deterministic methods are applied. Modes in the posterior distribution are identified by several hundred optimiser searches, where the optimiser is initialised to a random sample from the prior distribution, and run until it converges. This method identifies several modes. However, two of the modes dominate the posterior distribution, and are approximately symmetric. The remaining modes may be ignored as it has been shown that these are local maxima within the basin of attraction of one of the two dominating modes (Evans, 2001).

Assume two posterior modes have been identified, labelled \(V_1\) and \(V_2\). The vector \(J_{12} = V_2 - V_1\) defines a jump in state space from \(V_1\) to \(V_2\). To propose a jump from \(V_1\) to \(V_2\), we modify the proposal distribution to incorporate the jump: \(q(V^* | V_t + J_{12})\). To sample within the current mode we set the jump to be zero. Assume we are in mode 1, then a new sample is generated by the following algorithm:
(a) Propose a mode jump with probability 0.5; \( J = 0 \) or \( J = J_{12} \).

(b) Propose a new state, \( V^* \), by drawing a sample conditionally on the current state \( V_t \):

\[
V^* = V_t + J + N(0, \sigma^2 K_v).
\]

(c) Compute the acceptance probability:

\[
\alpha = \min \left\{ 1, \frac{p(V^*|S^o)q((V_t + J)|V^*)}{p(V_t|S^o)q(V^*|(V_t + J))} \right\}.
\]

(d) Draw sample \( \alpha_{test} \) from uniform distribution \([0, 1]\).

(e) If \( \alpha > \alpha_{test} \) then set \( V_{t+1} = V^* \); else \( V_{t+1} = V_t \).

To ensure that it is possible to jump back from \( V_2 \) we must also propose a jump \( V_1 = V_2 - J \) in our sampler scheme with equal probability to the jump from \( V_1 \) to \( V_2 \); this maintains detailed balance for the sampler. The above method is equivalent to combining the proposal distributions by the mixing method described in (Brooks, 1999).

The practical advantage of this method is that the mode finding is completed before the sampling commences. This means that specialist mode finding methods may be employed for each particular problem, and the computational effort required to find the modes is not repeated many times during a sampling run. The presented algorithm is similar to existing methods for sampling from multi-modal distributions, such as simulated tempering (Marinari and Parisi, 1992), tempered transitions (Neal, 1995) and mode jumping kernels for MCMC (Tjelmeland and Hegstad, 2001). However, these all rely on simultaneously identifying modes and sampling and we have found that in practice, with the well separated modes in this problem, the existing methods do not converge in any reasonable time, and initial mode finding is the only computationally feasible method.

If the aim is to efficiently explore each mode individually (or where the problem has been rendered uni-modal by the use of a non-zero mean prior) the hybrid Monte Carlo algorithm (Duane et al., 1987) is used to explore of the uncertainty within the modes.

### 4.2. A sparse, sequential Bayes algorithm

We next turn to our second approach for tackling the intractable computations with the posterior distribution,

\[
p(V|S^o) \propto \prod_{i=1}^{N} p(s_{oi}^o|v_i) p(V).
\]

Our method is based on an approximation to eq. (10) which allows us to reduce the problem of computing \( 2N \)-dimensional integrals over the wind fields \( V \) by a sequence of two dimensional integrations. We use the fact that we can build up the posterior eq. (10) sequentially, by processing the observations at different spatial locations one after the other in an arbitrary sequence. Our approximation will use the two following main ideas.

(a) As we advance through the sequence of observations, we will update a locally optimised Gaussian approximation to the posterior.

(b) Since this first approximation has the unpleasant feature of depending on the chosen data sequence, a further improvement, by subsequent repeated cycles through the data sequence is performed until sufficient convergence to a fixed point is achieved. The second step is nontrivial, because re-processing a data point for a second time requires an approximate deletion of its likelihood from the approximate posterior.
We will begin by explaining the first sequence of approximate posterior updates. Assuming that we have already processed \(t\) data points, the next observation with its likelihood, \(p(s_{t+1}^o|v_{t+1})\), leads to the exact update of the posterior distribution,

\[
p_{t+1}(V|S_{t+1}^o) = \frac{1}{Z_{t+1}} p(s_{t+1}^o|v_{t+1}) p_t(V|S_t^o),
\]

where \(S_t^o = \{s_{t}^o, s_{t-1}^o, \ldots, s_1^o\}\) denotes the set of observations which have been processed up to time \(t\) and \(Z_{t+1}\) is a normalisation constant. Having observed all \(N\) data points, we obtain \(p_{2N}(V|S^o)\) as the desired full posterior eq. (10).

Initialising the sequential update with the tractable Gaussian prior distribution \(p_0(V) = p(V)\) from eq. (2), the first observation leads to a non-Gaussian joint density \(p_1(V|S_1^o)\) for the set of wind vectors \(V\). The basic idea of our approximation is to project \(p_1\) to a sensibly defined “closest” Gaussian distribution \(q_1\). By using \(q_1\) in place of \(p_1\) on the right hand side of the update eq. (11) we get, again, a non-Gaussian approximation \(\tilde{p}_2(V|S_2^o)\) to the true posterior \(p_2(V|S_2^o)\). We can repeat the projection method to obtain a tractable Gaussian approximation \(q(V|S_2^o)\). Performing the update and projection steps for any observation in the sequence,

\[
\begin{align*}
\tilde{p}_{t+1}(V|S_{t+1}^o) & = \frac{1}{Z_{t+1}} p(s_{t+1}^o|v_{t+1}) q_t(V|S_t^o) \\
q_{t+1}(V|S_{t+1}^o) & = q_{t+1}(V|S_{t+1}^o)
\end{align*}
\]

for \(t = 0, \ldots, N-1\) leads to a sequence, \(q_t\), of Gaussian approximations to the true posteriors \(p_t\). To minimise the instantaneous loss of information in the projection step, we choose \(q_{t+1}\) to be the closest distribution to \(\tilde{p}_{t+1}\) in the family \(\mathcal{G}\) of Gaussian distributions where closeness is measured by the (non-symmetric) relative entropy or Kullback-Leibler (KL) divergence (Cover and Thomas, 1991). This means, we set

\[
q_{t+1}(V|S_{t+1}^o) = \arg\min_{q \in \mathcal{G}} \int dV \, \tilde{p}_{t+1}(V|S_{t+1}^o) \ln \frac{\tilde{p}_{t+1}(V|S_{t+1}^o)}{q(V)}.
\]

Note, that our choice of the order of original and approximating distributions is different from the one used in the variational approximation (see e.g. Ghahramani and Beal in (Opper and Saad, 2001)) frequently used in the machine learning community. We believe that our choice is more sensible because the log-ratio in eq. (13) is weighted by the less approximate Gaussian density \(q_{t+1}\) which has the same first and second moments as \(\tilde{p}_{t+1}\). The idea of approximating a posterior by sequentially propagating moments has been discovered independently in different scientific communities and applied to inference problems with Gaussian processes in Csató and Opper (2002).

The only remaining nontrivial task is the computation of the moments, i.e. the mean vector and of the covariance matrix for the non-Gaussian distribution \(\tilde{p}_{t+1}(V|S_{t+1}^o)\). However, since all variables \(v_t\), except for \(v_{t+1}\), appear only in the Gaussian density \(q_t\) in eq. (12), the corresponding multidimensional integrals are easily reduced to two-dimensional non-Gaussian ones over the vector \(v_{t+1}\). In many cases of interest, such as the Gaussian mixture model specified by the direct inverse model introduced in Section 2.2, these can be performed analytically (Csató et al., 2001). Obviously, our approximation becomes exact when the likelihood model is Gaussian, i.e. when \(\ln p(s_{t+1}^o|v_{t+1})\) is a quadratic form in \(v_{t+1}\).
A simple representer lemma, proved in (Csató and Opper, 2002), shows how the means and the covariances for the set of fields at all points can be represented using a 2t dimensional ‘mean ’ vector and a 2t \times 2t dimensional ‘kernel’ matrix, which both depend only on the t spatial locations which have been “visited” by the algorithm so far. This ‘mean ’ vector and the ‘kernel’ matrix are the parameters which are processed by the sequential algorithm. The algorithm recomputes their elements and enlarges their dimensionality when a new data point is added. It should be noted that although we process a single observation at a time, at each instant our method updates an approximate joint Gaussian density of the wind field at all observation points.

The propagation of a Gaussian approximation to the posterior distribution is in the spirit of the Kalman filter algorithm (Kalman and Bucy, 1961). There is however a crucial difference which comes from the fact that we do not introduce a local approximation to the original likelihood (e.g. linearisation). The integration in eq. (12) is a global smoothing operation which allows us to deal with likelihoods that are non-smooth or even non-continuous functions of the parameters.

Although we have dealt with the problem of high dimensional intractable integrals, the major problem for a practical application of this sequential Bayesian procedure to realistic problems is the increase in computational complexity with the number of observations, N. The increase in the size of the matrix used by the algorithm makes an application to large systems infeasible. Hence, a further sparse approximation is necessary. The simplest idea would be to discard a new data point if its influence on the entire posterior process is sufficiently small. This influence could be measured by the change of the approximated posterior density when the data point is being processed. To quantify the change, one could use an information theoretic measure like the relative entropy defined in eq. (13). Discarding a substantial fraction of observations would then lead to a slower increase of the ‘kernel’ matrix size but may on the other hand lead to a substantial loss of information.

An alternative representation, which gives insight into the sparsity heuristic we have developed is to consider an approximate likelihood, as opposed to projecting onto an approximate posterior. At step t the density has the form

$$q_t(V|S^o_t) \propto Q_t(S^o_t|V_t) p_0(V).$$

(14)

The effective “Gaussian” likelihood can again be written in a product form corresponding to individual sites

$$Q_t(S^o_t|V_t) = \prod_{i=1}^{t} L_i(S^o_t, v_i),$$

(15)

where the logarithm of $L_i$ is a quadratic form in the wind field $v_i$.

With slightly more effort, we can still keep some amount of the information contained in the discarded observations without increasing the size of the matrix. This is achieved by replacing the likelihood $Q_t$ in eq. (14) by a further optimised sparse approximation $Q^s_t$ which is still the exponential of a quadratic form but depends on a small set of observations only. The sparse approximation is performed sequentially, together with the projection and update steps of the Bayes sequential algorithm introduced above. Suppose that at step t, we have a sparse approximation to the likelihood $Q^s_t(S^o_t|V^s)$, which depends only on a subset, $V^s$, of wind vectors, which we call basis vectors. The update and projection step applied to a new data point $v_{t+1}$ leads to an updated Gaussian posterior $q_{t+1}$ and a corresponding effective Gaussian likelihood $Q_{t+1}(S^o_{t+1}|V^s, v_{t+1})$. 
We aim at constructing an approximation to the posterior which is of the form

\[ q_{t+1}^*(V|S_{t+1}^o) \propto Q_{t+1}^*(S_{t+1}^o|V^*) p_0(V), \]

for which the new likelihood \( Q_{t+1}^* \) is again sparse, that is a function of only the basis vectors, \( V^* \). To make \( q_{t+1}^* \) and \( q_{t+1}^o \) as close as possible the optimisation is most naturally performed when we measure the closeness of densities by the KL divergence

\[ \int dV \frac{q_{t+1}^o(V|S_{t+1}^o)}{q_{t+1}^*(V|S_{t+1}^o)} \ln \frac{q_{t+1}^o(V|S_{t+1}^o)}{q_{t+1}^*(V|S_{t+1}^o)}. \]  

(16)

Note, that compared to eq. (13) the order of original and approximated distributions has been interchanged. One can show that the optimal likelihood function is obtained by replacing \( v_{t+1} \) by its minimal mean square prediction \( \tilde{v}_{t+1}(V^*) \) given the basis vectors under the prior distribution. This means that we replace \( v_{t+1} \) by its projection on the space spanned by the basis vectors \( V^o \), where the metric is induced by the inner product defined by the covariance kernel, \( K_v \), from eq. (5). Hence, we get

\[ Q_{t+1}^*(S_{t+1}^o|V^*) = Q_{t+1}(S_{t+1}^o|V^*, \tilde{v}_{t+1}(V^*)). \]

(17)

When the minimal KL divergence is smaller than some given tolerance \( \epsilon \), we replace \( q_{t+1} \) by \( q_{t+1}^o \) and the size of the basis set remains the same. Otherwise, we augment the basis vector set by the vector \( v_{t+1} \). If the number of basis vectors increases beyond a maximally tolerable number, we can apply the sparsification method to discard old basis vectors which achieve a small information score using eq. (17).

In the second step of our approximation scheme, we deal with an inherent problem in the sequential method as defined thus far. This stems from the dependence of the final prediction on the order in which the observations are processed, which is arbitrary. It is thus highly desirable to be able to sweep through the data several times to reﬁne the approximation until convergence is achieved. A naive repetition of the projection and update steps eq. (12) in a second sweep through the data would be highly inconsistent. It would violate the assumption, which is inherent in the sequential construction of the posterior, that the local likelihood for the ﬁeld \( v_{t+1} \) has not yet been included in the posterior.

A method for solving this problem within the general framework of sequential Bayesian methods has been suggested by Minka (2000). In the expectation propagation approach, Minka shows how to approximately delete the information contained in an observation from the posterior before the update / projection step is performed again on this data point. If we want to re-process an observation, say \( s_{t+1}^o \), we replace the update step of the posterior in (12) by

\[ \tilde{q}_{t+1}(V|S^o) \propto p(s_{t+1}^o|v_i) \tilde{q}_{t+1}(V|S^o) \]

(18)

where \( \tilde{q}_{t+1} \) is a Gaussian approximation to the posterior from which the observation \( v_i \) was deleted. \( \tilde{q}_{t+1} \) is constructed from the current Gaussian approximation \( q(V|S^o) \) to the posterior by dividing it with an effective local Gaussian likelihood, \( L_i \), which is defined analogous to (15),

\[ \tilde{q}_{t+1} \propto q(V|S^o)/L_i(S^o, v_i). \]

(19)

After we get the new Gaussian posterior approximation from the projection step, we must also recompute the local likelihood \( L_i \).

It has been shown that when this procedure converges, the final approximation is equivalent to the so called TAP (named after Thouless, Anderson and Palmer) approximation. The
latter method was originally developed in the field of statistical physics (Mezard et al., 1987) and has recently been successfully applied to approximate inference (Opper and Winther, 2001) for a variety of probabilistic models (for a review see Opper and Saad (2001)). It should be noted that upon convergence it is possible to extract information from the approximate posterior which goes beyond simple Gaussian statistics. In fact, if we marginalise the updated approximation \( \tilde{p}_i(V|S^o) \) with respect to the wind vector \( v_i \), the inclusion of the exact likelihood \( p(s^o_i|v_i) \) in (18) leads to a non-Gaussian approximation to the marginal posterior \( p(v_i|S^o) \) which can show nontrivial behaviour like multi-modality.

A heuristic argument adopted from statistical physics may indicate when this approximation scheme might be expected to work well. If we restrict ourselves to the accuracy of the approximated marginal posterior, we can see that the crucial assumption is the Gaussianity of the marginal \( \tilde{q}_i(v_i|S^o) \) for observation \( s^o_i \) removed. If the Gaussian process covariance has a sufficiently large length scale with respect to the sampling design, \( v_i \) is influenced weakly by a large number of neighbouring fields \( v_j \) and a heuristic central limit argument suggests that a Gaussian approximation is appropriate.

![Fig. 3. Hybrid Monte Carlo samples using the inverse model from a North-West Atlantic wind field, on 10th June 1998, showing the two dominating modes. The black arrows define the mean wind vectors, while the grey ellipses around the tips of the arrows define the marginal covariance (plotted at 5 standard deviations to aid visualisation) at each observation.](image)
Bayesian approaches to scatterometer inverse problems

Table 1. A summary of the results comparing the forward and direct inverse approaches to local modelling, showing their broad equivalence in terms of errors on an independent test set. VRMSE: vector root mean squared retrieval error, Spd. std.: standard deviation of the retrieved wind speed error, Dirn. std.: standard deviation of the retrieved wind direction error, within 20°: the percentage of retrieved most probable wind vectors within 20° of the “true” wind, Time: the CPU time taken to retrieve each wind vector from the s“ observations.

<table>
<thead>
<tr>
<th></th>
<th>VRMSE (ms⁻¹)</th>
<th>Spd. std. (ms⁻¹)</th>
<th>Dirn. std. (°)</th>
<th>within 20°</th>
<th>Time (s pattern⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Forward</td>
<td>2.7</td>
<td>1.7</td>
<td>16</td>
<td>53</td>
<td>1.05</td>
</tr>
<tr>
<td>Inverse</td>
<td>2.6</td>
<td>1.8</td>
<td>17</td>
<td>44</td>
<td>0.03</td>
</tr>
</tbody>
</table>

can be chosen to be a small fraction of the overall number of data without losing too much predictive power.

For some models and likelihoods the integrals required by the algorithm can be computed analytically (e.g. the inverse model, where the likelihood corresponds to a Gaussian mixture model), however for the non-linear forward model this is not possible. In this work we linearise the forward model, eq. (3), about the marginal posterior mean at time t, which leads to a Gaussian posterior. The linearisation means that it is important that the predicted marginal posterior distribution, q_t(v_{t+1}) is reasonably close to a local maximum a posteriori probability solution, thus when using the forward (likelihood) model a non-zero mean prior is required.

5. Results and discussion

We illustrate the methods in Sections 4.1 and 4.2 on the retrieval of two typical wind fields. First, we evaluate the performance of the local forward and inverse models.

5.1. Forward models and direct inverse models

A test set of 60,000 independent scatterometer observations was processed using a local inversion of the forward model, by finding the most probable wind vector given the scatterometer observation with a uniform prior over v_i, and the direct inverse model. This allows us to assess the accuracy of the likelihood models with respect to the NWP observations, which is the closest thing we have to “ground truth”. The results in Table 1 show that both models are very comparable in performance, as might be expected. The forward model is considerably better at “guessing” the correct direction. Seeking the correct solution within 45°, that is in the correct quadrant, means the results rise to 63% and 54% respectively, showing the inverse model also has skill in determining the correct from the ambiguous solution. The penalty to pay for using the forward model comes in the time taken to process one pattern: it is nearly two orders of magnitude slower than the direct inverse model, due to the requirement for non-linear optimisation.

5.2. Sampling and the sequential Bayes algorithm

The results of the hybrid Monte Carlo sampling from eq. (7) using a zero mean prior from the two dominating modes is shown in Fig. 3; (a) shows the “true” mode, while (b) illustrates the ambiguous mode. There were 10,000 samples drawn from each mode, the results for the
figures were computed on the last 5,000 samples to account for burn-in and convergence was assessed using multiple chains. As expected, with wind fields that have moderately high wind speeds, there is low posterior variance in wind speed and direction. Uncertainty at the edges is greater than that in the middle of the wind field, and this is due to the reduced impact of neighbours at the edges, as well as lower wind speeds in the upper right hand corner, which result in a lower signal to noise ratio in scatterometers.

![Graphical representation of wind field](image)

**Fig. 4.** (a) Hybrid Monte Carlo samples using the inverse model from a South Atlantic wind field, on 25th January 1999. (b) the corresponding UK Met. Ofce wind vector estimates using the scatterometer data and NWP predicted winds.

Fig. 4 (a) shows the results of sampling from the most probable mode of the posterior distribution eq. (7). An interesting feature of this plot is the increased uncertainty in the local models where there is low wind speed. This correlates with the physical characteristics of the data collection system, where it is known that at low wind speeds, wind direction is especially difficult to estimate from $s^o$. This is evident by the larger ellipses in the upper right and lower left quadrant of the wind field. At low wind speeds, the two dominant solutions in local models are not completely separated. Thus depending on the certainty in the neighbourhood, it is possible for wind vectors to swap into their ambiguous state. Also, this plot highlights how spatial consistency in the measurements increases certainty in the posterior distribution.

In order to assess the probability mass under the two dominating modes in both June
and January wind fields 40,000 samples were sub-sampled from a sample chain of 2,000,000 points obtained using the mode jumping sampler described in Section 4.1. For each example the two maximum a posteriori probability modes were supplied to the sampler. After burn-in, each sampler remained in the dominant mode, suggesting it is very likely that this is the correct mode. This can be explained by the factorising likelihood in eq.(7). The likelihood is a product of the local model probability distributions. If the local mass associated with the local wind vectors is 0.51 for the first mode and 0.49 for the second, then when these are combined in the likelihood, the total likelihood of the first mode would be

$$P(m_1) = (0.51)^N/(0.51^N + 0.49^N)$$

where $N$ is the number of wind vectors. Hence, when $N = 200$, $P(m_1) = 0.9997$, and the majority of the mass of the posterior is in the first mode although it is almost evenly distributed between each mode in the local model.

Note the prior would tend to reduce the effective $N$ by introducing correlation. In other experiments (Evans, 2001) we have shown that for smaller wind fields with $N = 100$ using the probability mass from sampling allows us to choose the correct wind field 73% of the time, a small improvement over using maximum a posteriori probability.

Fig. 5. The results of sampling from the posterior distribution eq. (1) for the wind field in Fig. 4 using the forward model and hybrid Monte Carlo sampling (a), and the sparse, sequential Bayes algorithm (b). In contrast to previous figures these results use a non-zero mean Gaussian process prior, which represents the error statistics of the NWP forecast wind field, which forms the mean. In (b) only the (50) basis vectors are drawn in black, the remaining wind vectors are interpolated, shown in grey.

Fig. 5 shows the results of sampling using the forward model defined in Section 2.1
and using the sequential Bayes algorithm. In this case a non-zero mean prior is used, which renders the posterior distribution unimodal. This prior uses the forecast NWP wind vectors as the mean function, with the covariance, which has the same form as described in Section 3, describing the error (or increment) structure of this field.

It can be seen (Fig. 5) that the retrieved distributions are broadly similar, but that the sequential Bayes result (Fig. 5(b)) is smoother and has larger posterior uncertainty. This can be traced to the approximate nature of the sparse representation; we have run the experiment allowing more basis vectors to be used and the results tend to the sampling solution (Fig. 5(a)). In previous work (Csató et al., 2001) we have shown that sparse representations can well approximate the mean function, but that the covariance estimates degrade much more quickly as sparsity increases.

The key benefit of the sequential Bayes algorithm is speed. To obtain the samples from the posterior distributions using both inverse and forward models took many hours of CPU time, for just one wind field. This is impractical for operational use, where approximately 200 wind fields of the size shown are observed by the scatterometer each day. The sequential Bayes algorithm, on the other hand proceeds very quickly; 20 sweeps through the data using the expectation propagation method take less than two minutes of CPU time, on a 2 GHz Pentium machine. Although we do not show the results of applying sequential Bayes algorithm to the posterior distribution evaluated using the direct local inverse model, as the results are very similar to the forward model results, it is even faster since the integrals required for the update step in the algorithm can be computed analytically. When using the forward model it is necessary to linearise the model about the current state estimate to derive an expression for the integral, and this approximation means more sweeps through the data are required for the algorithm to converge, as well as each step being more computationally expensive.

6. Conclusions

In Section 2 we have shown local forward and direct inverse (likelihood) models for scatterometer wind retrieval. We have described a Bayesian solution to the retrieval of wind fields from scatterometer data and have applied novel sampling and sequential Bayes algorithms to this problem.

For the resulting wind fields to be used sensibly it is necessary to supply a mean prediction and the associated uncertainty. For the wind field retrieval problem this necessitates sampling or use of the sequential Bayes algorithm. The fact that the two methods give consistent results gives us confidence in the use of the sequential Bayes algorithm, which can use either direct inverse or forward likelihood models.

One drawback with using the forward models in wind field retrieval is that it is not possible to start with a zero mean prior: one either has to start with the NWP winds as the initial field, which results in a solution similar to Fig. 4(a) or use a non-zero mean prior as is done in Fig. 5. This is because the forward (likelihood) model is non-linear, thus both hybrid Monte Carlo sampling and the sequential Bayes algorithm use the Jacobian of the forward model, evaluated at the current estimate of the state. With a zero mean initial state this Jacobian is poorly defined and the implied linearisation is a very poor approximation to the true model. An alternative we are exploring is to use “local” importance sampling to evaluate the 2D integrals at each step in the sequential Bayes update using the forward model.
In practice the sequential Bayes algorithm allows us to treat large wind fields, much bigger than the ones we have dealt with in the examples shown, in a consistent framework, which minimises the loss of information caused by the approximation, while allowing a fully Bayesian representation of the posterior distribution. While sampling does allow us to account for the non-Gaussian nature of the posterior distribution, all practical uses of the wind fields would utilise only the first two moments in any case, so this restriction to a Gaussian posterior is not too restrictive.

In theory, the sequential Bayes algorithm allows us to compute a good approximation to the total probability of all observations, that is the Bayesian evidence. This would give us a yardstick for comparing models with different hyper-parameters, and allows estimation of hyper-parameters in the posterior covariance function. It would be interesting to explore the differences between the prior covariance function hyper-parameters as estimated from NWP models and those estimated directly from the scatterometer data, which contains more detail in the wind fields.

In future work we plan to use additional sources of information, such as cloud drift winds, to obtain completely independent (of NWP models) estimates of the near surface wind field, by combining the multiple sources of information in the Bayesian framework proposed. This will be particularly important in the retrieval of wind fields when NWP forecasts are not available or desirable, for instance if the retrieved winds are to be used subsequently for estimating the state of the atmosphere in NWP data assimilation (where the impact of the NWP forecast in the prior could be very detrimental). An alternative to using more information to reduce the effects of ambiguity in the observations would be to modify the sequential Bayes algorithm to allow estimation in Gaussian process mixture models, which would enable us to cope with multi-modal or non-Gaussian posteriors in an efficient manner.

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References


