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Comparison of the Ensemble Transform Kalman Filter with the Ensemble Transform Kalman Smoother

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Abstract

Data assimilation aims to produce initial conditions for a weather forecast that are as close as possible to reality. The Ensemble Kalman Filter (EnKF) is a data assimilation method that forecasts a statistical sample of state estimates using a linear or nonlinear model. This forecast state is then updated with observations according to given errors in the observations and the forecast state. This update is the best estimate of the state of the system, called the analysis. Various formulations of the EnKF are discussed, which di er in the analysis step. The ETKF has been chosen for implementation.

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Contents

1	Intr	oductio	on 1	I			
	1.1	What i	s data assimilation?	I			
	1.2	Backgr	ound	2			
	1.3	Goals f	For the project	3			
	1.4	Outline	e of the report	1			
2	For	Formulations of the Kalman Filter 7					
	2.1	Backgr	ound information	7			
		2.1.1	The forecast model	7			
		2.1.2	Statistics	7			
	2.2	The Ka	alman Filter	3			
		2.2.1	The observation operator	3			
		2.2.2	Forecast equations)			
		2.2.3	Kalman gain matrix)			
		2.2.4	Analysis equations)			
		2.2.5	Advantages and Disadvantages of the KF	I			
		2.2.6	Summary of the KF	I			
	2.3	The Er	nsemble Kalman Iter (EnKF)	I			
		2.3.1	EnKF terms and notation)			
		2.3.2	Forecast equations)			
		2.3.3	Analysis equations	3			
		2.3.4	Advantages and Disadvantages of the EnKF	5			
		2.3.5	Summary of the EnKF	5			
	2.4	The Er	nsemble Square Root Filter	ό			
		2.4.1	The observational ensemble perturbation matrix	Ś			
		2.4.2	The Kalman gain matrix 16	ć			
		2.4.3	Analysis step	7			

		2.4.4	Advantages and Disadvantages of the EnSRF	17	
		2.4.5	Summary of the EnSRF	18	
	2.5	The E	nsemble Transform Kalman Filter (ETKF)	18	
		2.5.1	The ETKF equations	18	
		2.5.2	Summary of the ETKF	19	
	2.6	Smoot	hing - What is it and why is it important in data assimilation?	19	
	2.7	The K	alman Smoother (KS)	21	
		2.7.1	Summary of the KS	23	
	2.8	The E	nsemble Square Root Smoother (EnSRS)	23	
		2.8.1	Experimental results with the EnSRS	24	
		2.8.2	Summary of the EnSRS	25	
	2.9	The E	nsemble Transform Kalman Smoother and an approximation	25	
		2.9.1	ETKS using a smoothed forecast state (ETKS)	25	
		2.9.2	ETKS using a xed square root matrix (ETKS2)	27	
		2.9.3	Expected di erences in ETKS and ETKS2 results	27	
		2.9.4	Summary of the ETKS and ETKS2	27	
3	The Swinging Spring model 29				
	3.1	The S	winging Spring	29	
		3.1.1	Summary of the swinging spring system	33	
	3.2	Nume	rical method of integration for the swinging spring	33	
	3.3	Norma	al mode oscillations	34	
		3.3.1	Initialization	34	
		3.3.2	Normal mode initialization (NMI) of the swinging spring system	34	
		3.3.3	Summary of the numerical integration of the swinging spring and NMI	35	
4	Imp	lemen	ting the ETKF, ETKS and ETKS2	38	
	4.1	Impler	mentation of the ETKF	38	
4.2 Implementation of the ETKS and ETKS2 methods		Impler	mentation of the ETKS and ETKS2 methods	39	
		4.2.1	Description of the ETKS and ETKS2 algorithms	40	
		4.2.2	ETKS algorithm	41	
		4.2.3	ETKS2 algorithm	42	
		4.2.4	Statistical signi cance method and the ensemble mean error	43	
		4.2.5	Summary of ETKF and ETKS/ETKS2 implementations and statis-		
			tical signi cance	44	

5	Res	ults	46			
	5.1	Comparison of the ETKF with xed-interval ETKS2 using perfect observations		46		
		5.1.1	ETKF	46		
		5.1.2	Fixed-interval ETKS2	49		
	5.2	Comparison of the ETKF with the ETKS2 using imperfect observations		50		
		5.2.1	ETKF	50		
		5.2.2	Fixed-interval ETKS2	52		
		5.2.3	Ensemble error of the ETKF and ETKS2 using di erent ensemble			
			sizes with imperfect observations	52		
		5.2.4	ETKS2 with varying lags using imperfect observations	54		
	5.3	3 Summary of results		55		
6	Con	Conclusions and Future work6.1Summary and discussion6.2Limitations of the experiments		66		
	6.1			66		
	6.2			69		
		6.2.1	The Swinging Spring system	69		
		5	50			
		5.2.1 ETKF9150F50500(.)-50000(.)-500(.)-500(.)-5 en7J50t0 5Li-1144(E9-500(

List of Symbols

Matrices and vectors

- *h* Observation operator
- H Linear Observation operator
- I Identity matrix
- K Kalman gain matrix
- *M* Linear dynamical model
- *m* Non-linear dynamical model
- *P* State error covariance matrix
- *Q* Model error covariance matrix
- *R* Observation error covariance matrix
- R_e Observation error covariance matrix for perturbed observations
- *S* Ensemble version of innovations covariance matrix
- 7 Post-multiplier in deterministic formulations of the EnKF
- *U; V; W* Orthogonal matrices
- <u>x</u> State vector
- X Ensemble perturbation matrix
- <u>y</u> Observation vector
- Y Observation ensemble perturbation matrix
- <u>d</u> Observation vector for perturbed observations (ETKF)
- *D* Observation ensemble perturbation matrix for the ETKF
- <u>z</u> Arbitrary vector
- *Z* Arbitrary matrix
 - Random observation error vector
 - Random model error vector
 - Diagonal matrix of eigenvalues
 - Matrix of singular values

Sub and superscripts

- f Forecast value
- a Analysis value
- *e* Ensemble value (unless otherwise stated)
- *i* Ensemble Member
- j Time index
- t True value
- 100 Average over 100 runs

Numbers of components

- *n* Number of state components
- *p* Number of observation components
- N Number of ensemble members
- *I* Number of time-steps in time-interval

Operators

Standard deviation

- < z > Expectation of value
- z Mean value
- z^{ℓ} Deviation from the mean
- *ż* Scaled value

Abbreviations

KF	Kalman Filter
EnKF	Ensemble Kalman Filter
SRF	Square Root Filter
EnSRF	Ensemble Square Root Filter
ETKF	Ensemble Transform Kalman Filter
KS	Kalman Smoother
SRS	Square Root Smoother
EnSRS	Ensemble Square Root Smoother
etks	Ensemble Transform Kalman Smoother
etks2	Approximation of Ensemble Transform Kalman Smoother
NWP	Numerical Weather Prediction
NMI	Normal Mode Initialization
LNMI	Linear Normal Mode Initialization
NNMI	Non-linear Normal Mode Initialization
SVD	Singular Value Decomposition

List of Figures

1.1	Diagram of an analysis curve of a lter and a xed-interval smoother for the variable to be measured V in a time window. Analysis at time t_k , shown on the red line shows use of just past observations for the lter but all available observations for the xed-interval smoother	5
1.2	Diagram of an analysis curve of a xed-lag smoother for the variable to be measured V in a time window. Analysis at time t_k , shown on the red line	0
	shows use of the past and one future observation	6
3.1	Coordinates and forces for the swinging spring. Coordinates are given by angle , radius r and bob mass m . The gravitational force is given by mg and the elastic force is given by $k(r = l_0)$ where k is elasticity and r is unstretched	
2.2	length.	30
3.2	Coordinates and their fourier transforms of slow and fast variables for an uninitialized swinging spring. Parameter values give rotational and elastic frequencies of $f = 0.5$ and $f_r = 5$ respectively. Initial conditions are	
	$(; p; r; p_r) = (1; 0; 1:05; 0).$	32
3.3	Coordinates and their fourier transforms of slow and fast variables for a swinging spring with linear normal mode initialization. Parameter values give rotational and elastic frequencies of $f = 0.5$ and $f_r = 5$ respectively.	
	Initial conditions are $(; p; r; p_r) = (1; 0; 1; 0)$.	36
3.4	Coordinates and their fourier transforms of slow and fast variables for a swinging spring with nonlinear normal mode initialization. Parameter values	
	give rotational and elastic frequencies of $r = 0.5$ and $r_r = 5$ respectively. Initial conditions are $(\cdot, p, \cdot, r, p_r) = (1.0.0.99540.0)$	37
	(1, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0, 0,	57

- 5.2 ETKF, perfect observations, absolute error trajectory of the ensemble mean (*N*

5.10	Fraction of analyses with ensemble mean within one ensemble standard de-	
	viation of the truth. Computed from 100 runs of the ETKF with di erent	
	random initial conditions. The observations are imperfect and include ran-	
	dom observation errors.	55
5.11	Time averaged errors of the ensemble mean for di erent lags. Ensemble	
	mean averaged from 100 runs of the ETKS2 with di erent random initial	
	conditions	55

Chapter 1

Introduction

1.1 What is data assimilation?

Data assimilation incorporates observational data into a numerical model to produce a model state that aims to be as close as possible to reality. In order to produce a weather forecast initial conditions are required in the numerical model. Numerical weather prediction (NWP) would not function properly by just inserting measured observations into a model. One main reason is that there are too few observations to determine the state of the system everywhere. Some regions are data rich such as Eurasia and North America, but others are poorly observed (Kalnay, 2003). In modern NWP the number of degrees of freedom of a model is of $O(10^7)$. But the number of conventional observations that can be inserted directly into a model is $O(10^4)$ (Kalnay, 2003). Many types of data such as satellite and radar observations are indirect, meaning they cannot be simply inserted into a model. In NWP it is necessary to have a complete rst guess estimate of the state of the atmosphere at all the grid points in order to generate the initial conditions for the forecasts (Bergthorsson and Doos, 1955). This is done by updating a prior estimate of the state of the s

Data assimilation methods are either sequential, variational or both. Sequential refers to the time behaviour of the scheme. This means that the analysis is solved by updating a forecast by assimilating available observations at each time-step in a time-window (Bannister, 2007). An example is the Kalman Iter (Kalman, 1960). Variational methods mean that iterations are used to minimize a cost function. The cost function is a function of the model state and is a measure of the di erence between the background state and the observations. The model state that minimizes the cost function is used as the analysis state

(Bannister, 2007). An example of a variational method is 4D-Var (Dimet and Talagrand, 1986). Variational methods are most commonly used in operational NWP but are not the subject of this report.

1.2 Background

This report focuses on a sequential form of data assimilation called the Ensemble Transform Kalman smoother (Evensen and van Leeuwen, 2000). This is one of the more recent of a series of modi cations of the Kalman Iter (KF), introduced by Kalman [1960]. The KF uses an initial single analysis state estimate. This is forecasted to the next time-step using a linear model. Then the forecast is updated by assimilating the observations at that time, to produce the analysis state. A weighting is given to the observations by a Kalman gain matrix, according to a ratio between estimated errors in the forecast and estimated errors in the observations. The estimate of the errors in the forecast and analysis states are also updated at each time-step using the forecast model and the Kalman gain matrix. This step by step process is repeated until the latest analysis step reaches the end of the time window. The KF is not used in operational NWP since it cannot process non-linear models and is computationally very expensive (Burgers et al., 1997). More detail on the KF and the KF equations are given in section 2.2. The Ensemble Kalman Filter (EnKF, section 2.3), Ensemble Square Root Iter (EnSRF, section 2.4) and the Ensemble Transform Kalman Filter (ETKF, section 2.5) all use an ensemble of state estimates. Each ensemble member is forecast individually. A ratio between the ensemble forecast covariances and the observation covariances are used in the Kalman gain to assimilate the observations in the analysis stage. The ensemble mean analysis state and the analysis ensemble perturbation matrix are calculated from this single gain matrix. The ensemble analysis can then be calculated from the mean and the perturbations. Although there is added expense in updating an ensemble of state estimates rather than just one, overall they are more computationally e cient than the KF since you only have to store each ensemble member, not the full error covariance matrix. They can also use non-linear forecast models and are more suited to parallel processing. These features make them more desirable for operational NWP.

Smoothing di ers from Itering in that future observations are assimilated in the analysis step (Ravela and McLaughlin, 2007). This would imply that the smoothed analysis would be more accurate than the Itered analysis since the smoothed analysis is assimilating extra observations (Ravela and McLaughlin, 2007). Smoothing can be divided into two types, xed-interval and xed-lag. Fixed-interval smoothing uses all the observations in a time interval to update all the analysis states. Fixed-lag smoothing only updates a xed number of states prior to the current observation time, which saves computational cost (Cohn et al., 1994). The xed-lag Ensemble Transform Kalman Smoother 2 (ETKS2, see section 2.9.2) introduced in this report is based on the linear smoothing method of Jazwinski [1970] and the xed-lag method of Cohn et al. [1994]. The implementation is similar to the implementation of the ETKF by Livings [2005].

A schematic showing the comparison between a lter and a xed-interval smoother is shown in Figure 1.1. In the schematic the smoothed solution (black curve on smoother diagram) is closer to the truth than for the Iter (black curve on Iter graph). A speci c analysis time t_k is highlighted to show that the lter analysis uses only past and present observations while the xed-interval smoother analysis uses all the observations in the time window at each step. Also the Iter analysis curve tends towards the true solution as more observations are assimilated. The xed-interval smoother analysis curve oscillates at approximately equal distance from the true solution over time, since all the observations are used in the time window at each step. The xed-lag smoother diagram (Figure 1.2) shows that some of the future observations are used at analysis time t_k . In this case the lag = 1 meaning the rst closest future observation is assimilated. Thus the solution is less accurate than the xed-interval smoother but more accurate than the Iter. Experiments using smoothing methods such as the xed-lag Kalman Smoother of Cohn et al. [1994] and the xed-lag Ensemble Square Root Smoother (see section 2.8) of Whitaker and Compo [2002] have been compared with the KF and EnSRF respectively and show an improvement in accuracy.

The ETKF forecast model implemented by Livings [2005] uses the swinging spring equations with a pseudo-random initial ensemble (see the implementation Chapter 4 for more details). The NNMI solution is used as the true solution. Livings [2005] compared the results for perfect and imperfect observations as well as for di erent ensemble sizes. The experimental part of this project aims to implement the ETKS2 and experiment with perfect and imperfect observations, as well as di erent ensemble sizes. The results can then also be compared with the ETKF implementation of Livings [2005].

1.3 Goals for the project

Implement the xed-lag ETKS2

Compare results of the xed-lag ETKS2 with the ETKF for

- { Perfect/Imperfect observations
- { Di erent lag lengths

{ Di erent ensemble sizes

1.4 Outline of the report

Chapter 2 describes the Kalman Filter (KF) and di erent stochastic and deterministic formulations of the KF. The Ensemble Square Root Filter is discussed and the Ensemble Transform Kalman Filter (ETKF) is given as an alternative Square Root Filter. The ETKF is the method used in the experiments. An introduction to smoothing and di erent smoothing types is given. The Kalman Smoother (KS) is derived, using the linear smoothing method of Jazwinski [1970] and a Bayesian Maximum likelihood approach. The xed-lag Ensemble Square Root Smoother is given and the ETKS follows on from the KS equations. An approximation to the ETKS called the ETKS2 is introduced as the method to be used in the experiments.

Chapter 3 describes the swinging spring model to be used in the experiments, as well as introducing the concept of initialization.

Chapter 4 gives the implementation of the ETKF and ETKS2 to be used in the experiments, including algorithms. An ETKS implementation is suggested although not used. Results from ETKF and ETKS2 experiments are discussed in chapter 5 and comparisons made. The conclusion in Chapter 6 summarises the report, as well as suggesting some limitations in the experiments and possible future work.



Figure 1.1: Diagram of an analysis curve of a lter and a xed-interval smoother for the variable to be measured V in a time window. Analysis at time t_k , shown on the red line shows use of just past observations for the lter but all available observations for the xed-interval smoother.



Figure 1.2: Diagram of an analysis curve of a xed-lag smoother for the variable to be measured V in a time window. Analysis at time t_k , shown on the red line shows use of the past and one future observation

Chapter 2

Formulations of the Kalman Filter

This chapter describes the necessary background on Itering and smoothing methods that is important in order to understand the Ensemble Transform Kalman Filter (ETKF) and Ensemble Transform Kalman Smoother 2 (ETKS2) that are used in the experimental work for this report. The meaning of the notation introduced in each section will remain in e ect unless specified otherwise.

2.1 Background information

2.1.1 The forecast model

The forecast model of the system that is being estimated is described here and applies to all formulations of the Kalman Filter (KF). Let *n* be equal to the state space dimension and *p* be equal to the observation space dimension. The state vector at time t_k is represented by \underline{x}_k . The observation vector at the same time is represented by \underline{y}_k . Let the true state of the system \underline{x}_k^t be related to the dynamical model operator *m* (such that $m : <^n ! <^n$) by

$$\underline{x}_{k}^{t} = m_{k}(\underline{x}_{k-1}^{t}) + k = 1$$

Here m_k can be linear or non-linear and $_{k-1}$ (a vector of dimension *n*) is the random model error, which is unbiased.

2.1.2 Statistics

Estimating the model errors and the forecast, analysis and observational error covariance matrices are essential to all the formulations of the KF. The random model error $_{k-1}$ is normally distributed as $N(0; Q_{k-1})$, where Q_{k-1} is the covariance matrix. Q_{k-1} is a measure

of the error correlation between di erent variables in the modelled state $(m_k(\underline{x}_{k-1}^t))$ and can be written in the form

$$Q_{k-1} = \left\langle \begin{array}{cc} k & 1 & T \\ k & 1 & k \end{array} \right\rangle; \tag{2.1}$$

where the angle brackets represent the expected value. The normal distibution (or Gaussian distribution) of the error () should be symmetric about zero. This means that the error is expected to average to zero and h i = 0, which means that is unbiased. The variance is the measure of the spread around the expected value. A narrower distribution of errors around the expected value means a smaller variance. The covariance measures the error correlation between two variables:

$$COV(1; 2) = h(1 h_1 i)(2 h_2 i)i:$$
 (2.2)

An error covariance matrix relates errors in the state vector components. The diagonal represents the variances and the covariances are the o -diagonal elements of the matrix.

In deriving the Kalman Smoother in section 2.7 it is important to know the following

$$y_k = H\underline{x}_k + k$$

where $_k$ (of length p) is the random observational error and is assumed to be unbiased. The observations have an error covariance matrix R_k (of dimension (p - p)). Assuming the observation error does not vary with time then

$$\left\langle \begin{array}{c} {}_{k} {}_{k}^{T} \right\rangle = R:$$
 (2.4)

in the KF.

2.2.2 Forecast equations

In the Kalman Iter the current forecast state \underline{x}_{k}^{f} (of length *n*), is a forecast of the previous analysis \underline{x}_{k}^{a} (of *n*) using the linear model *M*. For simplicity, it will be assumed that the model is perfect. Firstly an initial analysis state \underline{x}_{0}^{a} and an initial analysis error covariance matrix P_{0}^{a} (of dimension *n n*) are estimated. Then the linear forecast model *M* is used to produce the forecast at the next time step. The forecast state \underline{x}_{k}^{f} is defined as

 \underline{x}_{k}^{f}

where R is the observational error covariance matrix. This equation for K comes from minimising the following cost function to nd the analysis:

$$J(\underline{x}) = (\underline{x} \quad \underline{x}^{f})^{T} (P^{f})^{-1} (\underline{x} \quad \underline{x}^{f}) + (\underline{y} \quad H\underline{x})^{T} (R)^{-1} (\underline{y} \quad H\underline{x}):$$
(2.9)

The rst term gives the distance between the model state \underline{x} and the forecast state (\underline{x}^{f}) weighted according to the inverse of the forecast error covariance (P^{f}) . The second term gives the distance between the model state and the observations \underline{y} weighted according to the inverse of the observational covariance matrix R. The minimiser of J is the analysis state and satis es the analysis equation 2.10 (Kalnay, 2003).

2.2.4 Analysis equations

In the analysis step, the Kalman gain is used to assimilate the observations to update the forecast. The analysis \underline{x}_{k}^{a} (of length *n*) is defined as

$$\underline{x}_{k}^{a} = \underline{x}_{k}^{f} + K_{k}(\underline{y}_{k} \quad H\underline{x}_{k}^{f}); \qquad (2.10)$$

where y_k of dimension p is the vector of observations. The Kalman gain is also used to update the forecast error covariance to produce the analysis error covariance P_k^a . This is discussed in Burgers et al. [1997].

 P_k^a is de ned as

$$P_k^a = \left\langle (\underline{x}_k^a \quad \underline{x}_k^t) (\underline{x}_k^a \quad \underline{x}_k^t)^T \right\rangle$$
(2.11)

Firstly, the analysis (equation 2.10) is substituted into equation 2.11 and $K(H\underline{x}_k^t \quad \underline{y}_k^t) = 0$ is added to give

$$P_{k}^{a} = \left\langle (\underline{x}_{k}^{f} \quad \underline{x}_{k}^{t} + K_{k}(\underline{y}_{k} \quad \underline{y}_{k}^{t} \quad H\underline{x}_{k}^{f} + H\underline{x}_{k}^{t}))(\underline{x}_{k}^{f} \quad \underline{x}_{k}^{t} + K_{k}(\underline{y}_{k} \quad \underline{y}_{k}^{t} \quad H\underline{x}_{k}^{f} + H\underline{x}_{k}^{t}))^{T} \right\rangle :$$

$$(2.12)$$
Factorising out $(\underline{x}_{k}^{f} \quad \underline{x}_{k}^{t})(\underline{x}_{k}^{f} \quad \underline{x}_{k}^{t})^{T}$ and $(\underline{y}_{k} \quad \underline{y}_{k}^{t})(\underline{y}_{k} \quad \underline{y}_{k}^{t})^{T}$ gives

$$P_{k}^{a} = (I \quad K_{k}H) \left\langle (\underline{x}_{k}^{f} \quad \underline{x}_{k}^{t}) (\underline{x}_{k}^{f} \quad \underline{x}_{k}^{t})^{T} \right\rangle (I \quad K_{k}H)^{T} + K_{k} \left\langle (\underline{y}_{k} \quad \underline{y}_{k}^{t}) (\underline{y}_{k} \quad \underline{y}_{k}^{t})^{T} \right\rangle K_{k}^{T}$$

Note that there are no cross-product terms since it is assumed that $(\underline{y}_k \quad \underline{y}_k^t)(\underline{x}_k^f \quad \underline{x}_k^t) = 0$. Substituting in P_k^f and R respectively gives

$$P_k^a = (I \quad K_k H) P^f (I \quad H^T K_k^T) + K_k R K_k^T$$
(2.13)

Expanding out the brackets and re-arranging then gives

$$P_k^a = P^f \quad K_k H P^f \quad P^f H^T K_k^T + K_k (H P^f H^T + R) K_k^T$$
$$= (I \quad K_k H) P_k^f: \qquad (2.14)$$

2.2.5 Advantages and Disadvantages of the KF

The Kalman Iter is a good method in that it produces unbiased analysis and forecast state updates and hence their covariances are exact. Also the Iter is optimal since the Kalman gain is produced from minimizing a cost function to nd the optimal solution. However, the Kalman Iter is not currently used in operational weather forecasting. One reason is that the Kalman Iter can only be used for linear systems but NWP models are non-linear. Also, it is very computationally expensive to implement (Ehrendorfer, 1992), mostly due to its propagation of large forecast error covariance matrices using the model. In NWP models, the number of state variables is $O(10^7)$ (UKMO, 2009). Hence the corresponding error covariances will have $O(10^7 \ 10^7)$. Observations of $O(10^6)$ (UKMO, 2009) must then be assimilated at each analysis step. This requires too much storage to be practical. On top of this the Kalman gain requires the inversion of a large matrix, which is expensive and ine cient.

2.2.6 Summary of the KF

To summarise, the Kalman Iter uses a linear forecast model to produce a forecast (equation 2.5), which is updated by giving a weighting to the observations according to a ratio between forecast and observational error covariances. This is done using the Kalman gain (equation 2.8). This updated forecast is the analysis (equation 2.10). The forecast covariance matrix is also updated using the linear forecast model (see equation 2.7). The analysis covariance matrix (equation 2.14) comes from updating the forecast covariance matrix using the Kalman gain.

The KF is not currently used in NWP as it is too expensive to implement and can only be used for linear systems.

2.3 The Ensemble Kalman Iter (EnKF)

The Ensemble Kalman Filter was introduced by Evensen [1994]. Much of the following developments of the EnKF are discussed in Evensen [2003]. The EnKF uses many of the ideas of the KF but can can be applied to non-linear systems. While the KF uses a single

state estimate, the EnKF uses a statistical sample of state estimates, called an ensemble. Like with the KF there is a forecast step and an analysis step. In the forecast step the sample mean and error covariances are calculated from an ensemble of forecast states. These are used to calculate a single Kalman gain matrix. This Kalman gain matrix is used to assimilate the observations in to the forecast ensemble state and produce the analysis ensemble state.

2.3.1 EnKF terms and notation

The following equations 2.15 to 2.19 come from Evensen [2003], but using a new notation. Firstly, assume that an ensemble of state estimates of size N will be used. Therefore \underline{x}_i (of length n) represents the state vector for each ensemble member (i = 1; ...; N). The ensemble mean (of length n) is:

$$\underline{X} = \frac{1}{N} \sum_{i=1}^{N} \underline{X}_{i}:$$
(2.15)

Define the ensemble error covariance P_e (of dimension n n) as

$$P_e = \frac{1}{N-1} \sum_{i=1}^{N} (\underline{x}_i)$$
)e the iobservations in two



The forecast state \underline{x}_{i}^{f} at time t_{k} is dened as

$$\underline{x}_{i;k}^{f} = m(\underline{x}_{i;k}^{a})$$

The observation ensemble \underline{d}_i (of length p) has ensemble members i = 1; ...; N. Here i is pseudo-random observation errors normally distributed as N(0; R) (with mean zero and covariance R). An observation state matrix D_e can be defined with columns \underline{d}_i , i = 1; ...; N. An observation ensemble perturbation matrix D analogous to equation 2.18 (of dimension p = N) has columns \underline{d}_i^{l} , i = 1; ...; N. Also the observation ensemble covariance matrix is now defined as

$$R_e = DD^T$$
 (2.26)

The new Kalman gain has R_e in place of R:

$$K_{e} = P_{e}^{f} H^{T} (H P_{e}^{f} H^{T} + R_{e})^{-1}$$
(2.27)

The EnKF analysis step is similar to the KF, but uses the ensemble Kalman gain and the observation ensemble matrix as follows

$$\underline{x}_{i}^{a} = \underline{x}_{i}^{f} + K_{e}(\underline{d}_{i} \quad H\underline{x}_{i}^{f}):$$

$$(2.28)$$

for i = 1; ...; N. The ensemble mean can then be calculated as

$$\overline{\underline{X}^{\overline{d}}} = \overline{\underline{X}^{\overline{f}}} + K_{e}(\overline{\underline{d}} + H\overline{\underline{X}^{\overline{f}}_{e}}):$$
(2.29)

Here \underline{d} is replacing the actual observation \underline{d} . This can be accepted as it is, considering that \underline{d} tends to \underline{d} as the ensemble size increases. However, imposing the constraint $\overline{d} = 0$ on the vectors will ensure $\underline{d} = \underline{d}$ (Evensen, 2003). The ensemble perturbation matrix is

$$X^{a} = (I \quad K_{e}H)X^{f} + K_{e}D:$$
(2.30)

Thus

$$P_{e}^{a} = (I \quad K_{e}H)P_{e}^{f} + (I \quad K_{e}H)X^{f}D^{T}K_{e}^{T} + K_{e}D(X^{f})^{T}(I \quad K_{e}H)^{T}$$
(2.31)

The rst term in this expression is the desired result. Evensen [2003] discusses how this desired result can be achieved, but uses a di erent notation. As long as the distributions used to generate the model state ensemble and observation state ensemble are independent then the random vectors $_i$ are independent of $\underline{x}_i^f \quad \overline{\underline{x}}^f$. This implies that $X^f D^{fT}$ and hence the second and third terms tend to zero as the ensemble size increases. Equation 2.31 can be left as it is, but in order to make the 2nd and 3rd terms vanish altogether then the constraint $(\underline{x}^f \quad \overline{\underline{x}}^f)^T = 0$ can be imposed. Information on how to minimize the

calculation of large covariance matrices in the analysis step when implementing the scheme is also discussed in Evensen [2003].

2.3.4 Advantages and Disadvantages of the EnKF

The EnKF is able to use non-linear models, which is necessary in NWP. Using an ensemble of state estimates should improve the quality of the forecast provided that the members have been sampled properly. For example, care must be taken to ensure that the size of the sample is statistically representative of the model (Kalnay, 2003). Many problems can develop from undersampling, some of which have been investigated by Petrie [2008]. Inbreeding is one such problem. This means that the analysis error covariances are systematically underestimated after each of the observation assimilations. The obvious expense from the EnKF comes from maintaining the ensemble of state estimates. However, the covariance matrices are no longer evolved using the forecast model like they are in the KF, which is cheaper to implement. Also, a single Kalman gain is applied to each state estimate, which is less expensive than if separate Kalman gains were used for each member.

2.3.5 Summary of the EnKF

To summarise, The EnKF uses a statistical sample of state estimates that are forecasted forewards in time using a non-linear forecast model (equation 2.20). Unlike the KF, the forecast ensemble covariance matrix is not updated using the forecast model, but is updated using the matrix square of the forecast ensemble perturbation matrices (equation 2.21). In order that the analysis error covariance matrix can be calculated correctly observation perturbations are introduced (equation 2.25). Filters such as the EnKF that use perturbed observations are called stochastic liters. The forecast ensemble can then be updated using a single Kalman gain matrix (equation 2.27) to produce the analysis (equation 2.28). The desired result for the analysis error covariance matrix can be achieved by imposing the constraint $(\underline{x}^f \ \underline{x}^f)^T = 0$. This means that the last term on the right hand side of equation 2.31 vanishes.

Unlike the KF the EnKF can use non-linear models, necessary for NWP. Also an ensemble of state estimates should improve the quality of the forecast over one state estimate provided that the members have been sampled properly. The EnKF has added expense in maintaining an ensemble of state estimates instead of one for the KF. However, calculating the Kalman gain matrix and error covariance matrices are less expensive in the EnKF.

2.4 The Ensemble Square Root Filter

The Ensemble Kalman Square Root Filter (EnSRF) presented here uses the formulation in Tippett et al. [1999]. The EnKF is an example of a liter that uses perturbed observations. This implies that the observation ensemble is created by adding random vectors to the actual observations for each member. Therefore the EnKF is called a stochastic liter. Filters that do not use perturbed observations are called deterministic liters (Hamill, 2006). The EnSRF is a deterministic liter and di ers in the analysis step to the EnKF (section 2.3).

2.4.1 The observational ensemble perturbation matrix

Since the EnSRF is deterministic, the observational ensemble perturbation matrix Y is di erent than for the EnKF (section 2.3). The state members (\underline{x}_i) can be mapped to observation space by

у

these problems are eliminated when the square root analysis algorithm is used as it does not use perturbed observations (Evensen, 2004). Square root Iters (SRFs) have also been shown to be computationally more e cient and less expensive (Hamill, 2006). This is partly because they do not calculate the forecast error covariance matrix, only the ensemble state perturbations (Hamill, 2006).

2.4.5 Summary of the EnSRF

To summarise, the EnSRF is a deterministic lter, which means it does not use perturbed observations. This implies the observational ensemble perturbation matrix Y can be written in terms of the ensemble state perturbations in equation 2.33. Like for the EnKF, a statistical sample of state estimates are forecasted forwards in time using a non-linear forecast model. The forecast error covariance matrix is updated using the forecast ensemble perturbation matrices (equation 2.21). The Kalman gain used in the analysis stage now contains the EnSRF version of Y and the observational error covariance R is the same R used for the KF (equation 2.4). The analysis error covariance (see equation 2.39) can now be updated using the analysis perturbation matrices. The analysis perturbation matrix (equation 2.40) is updated using the square root matrix T (equation 2.41).

Unlike the EnKF, the EnSRF does not use pertubed observations, which can introduce sampling errors. SRFs such as the EnSRF have also been shown to be more e cient and less expensive than the EnKF (Hamill, 2006).

2.5 The Ensemble Transform Kalman Filter (ETKF)

2.5.1 The ETKF equations

The Ensemble Transform Kalman Filter was introduced by Bishop et al. [2001]. It is a form of square root lter. It is the chosen ltering method to be used in experimental work in this report (see the implementation section 4). The ETKF uses the identity

$$I \quad (Y^{f})^{T} S^{-1} Y^{f} = (I + (Y^{f})^{T} R^{-1} Y^{f})^{-1}$$
(2.43)

This can be veri ed as follows. Firstly multiply $I = (Y^f)^T S^{-1} Y^f$ by $(I + (Y^f)^T R^{-1} Y^f)$

$$I \quad (Y^{f})^{T}S^{-1}Y^{f} \quad (I + (Y^{f})^{T}R^{-1}Y^{f})$$

=
$$I \quad (Y^{f})^{T}S^{-1}Y^{f} + (Y^{f})^{T}R^{-1}Y^{f} \quad ((Y^{f})^{T}S^{-1}Y^{f})((Y^{f})^{T}R^{-1}Y^{f})$$

and then replacing S by equation 2.36 to give

$$I + (Y^{f})^{T} (Y^{f} (Y^{f})^{T} + R)^{-1} \begin{bmatrix} Y^{f} + (Y^{f} (Y^{f})^{T} + R)R^{-1}Y^{f} & Y^{f} (Y^{f})^{T}R^{-1}Y^{f} \end{bmatrix}$$

= $I + (Y^{f})^{T} (Y^{f} (Y^{f})^{T} + R)^{-1} \begin{bmatrix} Y^{f} + Y^{f} (Y^{fT})R^{-1}Y^{f} + Y^{f} & Y^{f} (Y^{f})^{T}R^{-1}Y^{f} \end{bmatrix}$
= $I + (Y^{f})^{T} (Y^{f} (Y^{f})^{T} + R)^{-1} = 0$
= I :

This is important since calculating R^{-1} is usually simpler than calculating S^{-1} , since R is usually diagonal while S has a more complicated structure. An eigenvalue decomposition can then be calculated by

$$Y^{fT}R^{-1}Y^{f} = U \quad U^{T}; (2.44)$$

where U is orthogonal and is diagonal. The identity in equation 2.43 can now be written as

$$I \quad (Y^{f})^{T} S^{-1} Y^{f} = U(I +)^{-1} U^{T}$$
(2.45)

The nal term U^T is required here to make the Iter unbiased (Livings et al., 2008). Hence the square root is given by

$$T = U(I +)^{\frac{1}{2}}$$
 (2.46)

This *T* is the square root that gives the ETKF. This is more e cient to compute than the square root in the EnSRF (matrix square root of 2.41) since (I +) is diagonal, making the calculation of (I +)¹ simple. According to Hamill [2006] the ETKF was veri ed to be faster to compute than the EnSRF of Whitaker and Hamill [2002].

2.5.2 Summary of the ETKF

To summarise, the ETKF is a form of square root Iter that uses the identity $I (Y^{f})^{T}S^{-1}Y^{f} = (I + YYot)$

observations in use, then theoretically the smoothed analysis at the end of the window is the same as the Itered analysis at the end of the window since in both cases all the observations have been assimilated. However, the analysis in the middle of the window will not be the same since the Iter will only use the observations in the rst half of the window while the smoother will use observations in both the rst and second half of the window. Thus at all points inside the window you would expect to get more accurate analyses for the smoother than for the Iter. However, a great deal more work will be required to get this since past, present and future observations in the window will have to be assimilated at each step for the smoother, as opposed to just past and present observations for the Iter (Ravela and McLaughlin, 2007).

The following descriptions of di erent types of smoothing methods come from Ravela and McLaughlin [2007]. There are two types of smoothing, xed interval and xed lag smoothing. Fixed interval smoothing updates all desired states within a time interval [0,T] using all the available observations. Fixed lag smoothing only updates a xed number of states prior to the current observation time. Therefore it only requires updates in a lag window W (where W < T) before the most recent measurement. Fixed lag smoothing and their applications have generally proven to be computationally faster, since fewer prior model states are updated using the current observations. For example, xed lag smoothing methods have been tested for the Goddard Earth Observing System (GEOS). It has been tested both for retrospective analysis of climate and for short term forecasting (Zhu et al., 1999).

A schematic of a lter analysis (black curve in lter diagram) and a smoother analysis (black curve on smoother diagram) is shown in Chapter 1 in Figure 1.1. The analysis time t_k is highlighted to show that the lter uses just past and present observations while the xed-interval smoother uses all available observations. The xed-lag smoother analysis is shown in Figure 1.2 and only assimilates one (*Wabj*ile thelagal sm267(time)]T--4oscil(one)-3267t-3267mation fester, 6

The observation distribution is

$$p(\underline{y}_{k+1:k+1}|\underline{x}_k) = \prod_{i=1}^{l} N(HM_{k! \ k+i}\underline{x}_k; R):$$
(2.51)

Substituting equations 2.50 and 2.51 into 2.49 and taking logs gives the cost function

$$\frac{1}{2}(\underline{x}_{k} \quad \underline{x}_{k}^{a})^{T}P_{k}^{a-1}(\underline{x}_{k} \quad \underline{x}_{k}^{a}) + \sum_{i=1}^{I}(\underline{y}_{k+i} \quad HM_{k! \ k+i\underline{x}_{k}})^{T}R^{-1}(\underline{y}_{k+i} \quad HM_{k! \ k+i\underline{x}_{k}}): (2.52)$$

Using one future observation time I = 1, the minimizer of equation 2.52 satis es

$$\underline{x}_{k}^{s} = \underline{x}_{k}^{a} + K_{kjk+1}^{s} (\underline{y}_{k+1} \quad HM_{k! \ k+1} \underline{x}_{k}^{a}); \qquad (2.53)$$

where \underline{x}_{k}^{s} is the smoothed analysis. Note that the ltered analysis must have already been calculated and stored before calculating 2.53. Here \mathcal{K}_{kjk+1}^{s} is the Kalman gain at time t_{k} using observations up to t_{k+1} .

$$K_{kjk+1}^{s} = P_{k}^{a} M_{k! \ k+1}^{T} H^{T} (HM_{k! \ k+1} P_{k}^{a} M_{k! \ k+1}^{T} H^{T} + R)^{-1}$$
(2.54)

and

$$P_k^s = (I \quad K_{kjk+1}^s H M_{k! \ k+1}) P_k^a$$
(2.55)

Now consider the smoothed solution using two future observations, I = 1/2. The minimizer of equation 2.52 now satis es

$$\underline{x}_{k}^{s} = \underline{x}_{k}^{a} + K_{kjk+1}^{s}(\underline{y}_{k+1} \quad HM_{k! \ k+1}\underline{x}_{k}^{a}) + K_{kjk+2}^{s}(\underline{y}_{k+2} \quad HM_{k! \ k+2}(\underline{x}_{k}^{a} + K_{kjk+1}^{s}(\underline{y}_{k+1} \quad HM_{k! \ k+1}\underline{x}_{k}^{a}))) \quad (2.56)$$

Now consider the solution using *L* future observations, I = 1;2;...L for the general xed-lag smoother. The same notation for the xed-lag smoother of Cohn et al. [1994] is introduced. The smoothed analysis will now be represented by $\underline{x}_{kjk+l'}^a$ where the subscript kjk + l notation represents the value at time t_k using observations up to t_{k+l} . Thus \underline{x}_{kjk+l}^a is calculated from

$$\underline{X}^{a}_{kjk+l} = \underline{X}^{a}_{kjk+l-1} + K_{kjk+l}(\underline{y}_{k+l} \quad HM_{kl-k+l}\underline{X}^{a}_{kjk+l-1}):$$
(2.57)
Here K_{kjk+l} is the Kalman gain at time t_k using observations up to t_{k+l}

covariance matrix can be computed from

$$(HP_{k+1;kjk+1-1}^{fa})^{T} = X_{kjk}^{a}$$

that sampling error in the estimate of the forecast-analysis error cross covariance matrix $(P_{k+l;kjk+l-1}^{fa})$ increases with *l*. This means a larger ensemble is required to take advantage of observations further removed from the analysis time. This is consistent with the fact that the quality of the analysis starts to degrade as the lag is increased beyond a certain point. This point is higher for more ensemble members (Whitaker and Compo, 2002).

2.8.2 Summary of the EnSRS

The EnSRS of Whitaker and Compo [2002] uses the xed-lag method of Cohn et al. [1994]. The xed-lag ensemble smoothed solution for I = 1 is the ensemble smoothed analysis state at time t_k using observations up to t_{k+I} . The Kalman gain (see equation 2.60) uses a cross covariance matrix between the forecast for the Kalman Iter update equation for time k + I and the lag I = 1 Kalman smoother analysis for time k. This is derived from a least squares sense (see Cohn et al. [1994]) and is used to produce the smoothed analysis ensemble. The cross-covariance and Kalman gain are also used to update the analysis covariance matrix P_{kjk+I}^a (see equation 2.65). Experimental results by Whitaker and Compo [2002] from the 40-dimensional model of Lorenz and Emanuel [1998] have shown that as expected ensemble members and higher lags. However, results suggested that sampling error in the estimate of the forecast-analysis error cross covariance matrix ($P_{k+I;kjk+I}^{fa}$) increases with I. Thus larger ensemble sizes are required to o set this increase in sampling error for higher lags.

2.9 The Ensemble Transform Kalman Smoother and an approximation

2.9.1 ETKS using a smoothed forecast state (ETKS)

The Kalman Smoother equations in section 2.7 can be modiled to produce the Ensemble Transform Kalman Smoother. The ETKS method described here follows on from the ETKF of Bishop et al. [2001] using the xed-lag smoother method of Cohn et al. [1994]. It is very similar to the smoothing method used in the experimental work for this report (section 2.9.2) and could be a useful method to investigate in the future.

Consider time t_k to be the current smoothing time and *I* to be the lag time. The solution for the smoother ensemble mean is equivalent to equation 2.62 in section 2.8. However, the Kalman gain is now written in a similar form to the KS. The only di erence being that the forecast model is evolved in the same way as for the xed-lag smoother of Cohn et al. [1994], from t_{k+l-1} to t_{k+l} instead of from t_k to t_{k+l} for the KS. *M* is now short for M_{k+}

Thus equations 2.68, 2.73 and 2.74 give the equations for the I-step xed lag ETKS.

2.9.2 ETKS using a xed square root matrix (ETKS2)

The ETKS2 described here is an approximation of the ETKS method described in section 2.9.1. The ETKS2 is the method used in the experimental work for this report (see implementation in section 4.2.1). The innovation vector used in the smoothing of the ETKS2 uses the original forecast states from the ETKF. In other words, the innovation vector in the ETKS2 is written as

$$(\underline{y}_{k+l} \quad H\underline{x}_{i;k+l}^{f}) \tag{2.75}$$

Note that the T matrix is calculated from the original forecast perturbations of the ETKF. T is the matrix square root of

$$T_{k+I}T_{k+I}^{T} = I \quad (Y^{f})_{k+I}^{T}S_{k+I}^{-1}Y_{k+I}^{f}.$$
(2.76)

$$S = Y_{k+l}^{f} (Y^{f})_{k+l}^{T} + R:$$
(2.77)

Here

$$Y_{k+l}^{f} = H X_{k+l}^{f}$$
(2.78)

where X_{k+l}^{f} is the original forecast perturbation matrix calculated in the ETKF. In other words, it is not the updated forecast perturbation matrix calculated by forecasting the smoothed analysis from the previous lag $(X_{k+ljk+l-1}^{f})$.

2.9.3 Expected di erences in ETKS and ETKS2 results

The innovation vector of the ETKS2 (equation 2.75) and the T matrix do not use the

terms of the square root matrix $T_{k+ljk+l-1}^{T}$ de ned by equation 2.41 in the EnSRF section 2.4. The ensemble analysis perturbation matrix can then be updated using the *T* matrix (equation 2.73).

The ETKS2 method di ers from the ETKS in that it does not use the forecast from the smoothed analysis in the innovation vector and the square root T matrix. The ETKS2 would therefore be expected to be less expensive to implement but less accurate for the lag l > 1.

Chapter 3

The Swinging Spring model

The swinging spring model is the forecast model that will be used for the ETKF and ETKS2 experiments in this report. This chapter begins by describing the swinging spring model and equations from Lynch [2002]. After a brief de nition of initialization and its rlevance in data assimilation, the application of normal mode initialization to the swinging spring model will be demonstrated using the implementation and results of Livings [2005].

3.1 The Swinging Spring

The Swinging Spring model of Lynch [2002] is illustrated by the following Figure (3.1) showing coordinates and forces for the swinging spring. The gure is based on the gure from Livings [2005].

A mass *m* is suspended from a xed point in a uniform gravitational eld, *g* by a light unstretched spring with length l_0 and elasticity *k*. The bob is constrained to move in the vertical plane and the string may stretch along its length but cannot bend. The bob is located using polar coordinates (*r*,). The spring length *r* is measured from the suspension point and the angle is measured from the downward path. The generalised momenta are radial momentum $p_r = mr$ and angular momentum, $p = mr^2$. The motion is assumed to be simple harmonic (SHM). The following Hamiltonian of the system gives the sum of potential and kinetic energies:

$$H = \frac{1}{2m}(p_r^2 + \frac{p^2}{r^2}) + \frac{1}{2}k(r - l_0)^2 \quad mgrcos \tag{3.1}$$



Figure 3.1: Coordinates and forces for the swinging spring. Coordinates are given by angle , radius r and bob mass m. The gravitational force is given by mg and the elastic force is given by $k(r - l_0)$ where k is elasticity and r is unstretched length.

From 3.1 one can derive the following equations of motion:

- = *p*

and $(r; p_r)$. The angular motion satis es

• +
$$\frac{g}{I} = 0$$
: (3.10)

The elastic motion satis es

$$f^{\theta} + \frac{k}{m}r^{\theta} = 0 \tag{3.11}$$

where $r^{d} = r$ *I*. These are SHM equations and have angular frequencies

$$! = \sqrt{\frac{g}{l}} \tag{3.12}$$

and

$$!_{r} = \sqrt{\frac{k}{m}}$$
(3.13)

The ratio of frequencies is:

$$= \frac{l}{l_{r}} = \sqrt{\frac{mg}{kl}} = \sqrt{\frac{l_{0}}{l}}:$$
 (3.14)

For the purposes of this work, parameters were used such that <<1. The rotational mode represents low frequency Rossby waves and contains the slow variables (;p). The elastic motion represents high frequency gravity waves and contains the fast variables ($r;p_r$). The slow and fast variables will interact for nite oscillations. Following the parameters used by Lynch [2002]:

$$m = 1$$

 $g = {}^{2}$
 $k = 100 {}^{2}$
 $l = 1$

This gives motions with cyclic frequency $f = \frac{w}{2} = 0.5$ and $f_r = \frac{l_r}{2} = 5$. Therefore = 0.1. The initial conditions are $(; p; r; p_r) = (1/0; 1.05; 0)$. Using these initial conditions and the equations of motion, the Matlab programme by Livings [2005] was implemented using the numerical method described in section 3.2. The graphs in Figure 3.2 show the motion of the fast and slow variables. It is clear that most of the energy is concentrated around f = f for the slow variables and most of the energy for the fast variables is concentrated at $f = f_r$. But there is a clear sign that the slow and fast variables have interacted in the peak in the fourier transforms of r and p non-linear nature of the system in that energy is being transferred from the fast variables to the slow ones.



Figure 3.2: Coordinates and their fourier transforms of slow and fast variables for an uninitialized swinging spring. Parameter values give rotational and elastic frequencies of f = 0.5 and $f_r = 5$ respectively. Initial conditions are $(;p;r;p_r) = (1;0;1.05;0)$.

3.1.1 Summary of the swinging spring system

The swinging spring models the motion of a mass m suspended from a swinging spring using the method of Lynch [2002]. The equations of motion are given by equations 3.2 to 3.5. Linearising about a stable equilibrium point when the bob is at rest gives equations which split the motions of the spring into high frequency (equation 3.11) and low frequency (equation 3.10). Using the method described in the following section (3.2) the motions of the swinging spring were plotted for both high and low frequency variables. The plots revealed that there was some interaction between the high and low frequency motions, indicating the non-linearity of the system.

3.2 Numerical method of integration for the swinging spring

The following information about the implementation of the swinging spring in Matlab comes from the description given in Livings [2005] and the implementation was validated by Livings [2005]. The swinging spring equations could be integrated using the standard Matlab ode45 function. This uses an explicit Runge-Kutta (4,5) pair with a changeable step size. This means that a 5th order Runge-Kutta scheme is used to integrate the equations and the di erence between this and a 4th order scheme is an estimate of the truncation error.

The error tolerance can be used to determine a limit on the step size. Let **Reltol** and **Abstol** represent the relative tolerance and absolute tolerance for each state space coordinate. If y(j) is the *j*th component of the solution vector and e(j) is the corresponding error component from the Runge-Kutta pair, then the step size must be minimized such that:

je(*j*)*j* max(**Reltol** *jy*(*j*)*j*; **Abstol**):

A primary aim was to keep the truncation error small compared with the observation error. This was done by using 10⁻³ for **Reltol** and 10⁻⁶ for every component of **Abstol**. The parameter **MaxStep** acts as an upper bound on the step size. This can be used to ensure stability of the numerical method. For information on stability of Runge-Kutta methods with relation to linear systems see Lambert [1991], section 5.12. Using the Runge-Kutta(4,5) scheme the maximum step size *h* for stability is *h* = 0.3. This is explained in Livings [2005], section 4.3. Since the period of a general harmonic oscillator is T = 2, taking the upper bound **MaxStep** of the step size to be $\frac{T}{20}$ for an arbitrary harmonic oscillator ensures stability for the system (see Livings [2005]). For the swinging spring system, in the case of no initialization the low frequecy motion has period T = 2 and the

high frequency motion has period $T_r = 0.2$. Therefore taking the maximum step-size to be **MaxStep** = $\frac{T_r}{20} = 0.01$ ensures stability. Using this step-size the solution (Figure 3.2) is evidently stable.

3.3 Normal mode oscillations

3.3.1 Initialization

A form of initialization called normal mode initialization (NMI) will be applied to the swinging spring system in section 3.3.2. Firstly, it is important to understand the concept of initialization. The atmosphere consists of a range between very fast changing systems with cycles lasting less than a minute, such as localized pressure variations and very slow changing systems lasting millenia such as climate change (Kasahara, 1976). Weather fore-casters are often not interested in the very fast systems. For example, pressure variations with cycles less than a minute are not going to help predict the arrival of a low pressure system at least 24 hours away. The useless high-frequency changes are called noise. If noisy observations are incorporated into the basic equations in NWP the forecast may contain spurious large amplitude high frequency oscillations (Lynch, 2002). The elimination of this noise can be achieved by adjusting the initial elds in a weather forecast, a process called initialization (Lynch, 2002).

In data assimilation it is important to ensure that the forecast of the previous analysis does not allow large gravity waves to develop. Constraints in the analysis step can be made to ensure that this does not happen.

3.3.2 Normal mode initialization (NMI) of the swinging spring system

The following implementation of NMI to the swinging spring system comes from Lynch [2002]. Both linear and non-linear normal mode initialization are applied. Suppose it is known that high frequency oscillations are not present in the motion of the swinging spring, yet they are still present in the numerical solution due to observational errors. How can we adjust the initial conditions to get rid of the high frequency oscillations by setting their initial amplitude to zero. Nonlinear Normal Mode Initialization (NNMI) instead sets the initial rates of change of the fast variables to zero. Using the same parameters as for the uninitialized swinging spring (section 3.1), in this experiment LNMI is imposed by setting r(0) = I and p

LNMI the initial conditions are $(;p;r;p_r) = (1;0;1;0)$. For NNMI, to calculate $\underline{p}_r(0) = 0$ set $p_r(0) = 0$ in equation 3.4. For $\underline{p}_r(0) = 0$, we rst calculate -(0) from equation 3.2. This can be substituted in to equation 3.5 and re-arranged to give:

$$r(0) = \frac{I(1 \quad 2[1 \quad \cos(0)])}{1 \quad (-(0)=I_r)^2}$$
(3.15)

Also, to ensure that -(0) used in 3.15 is consistent with 3.2 and the new value of r(0) we must set:

$$p(0) = mr(0)^2 - (0) \tag{3.16}$$

Figure 3.3 shows the coordinates and fourier transforms of the swinging spring variables for LNMI. Compared with the uninitialized case (Figure 3.2) the high frequency oscillations have been largely supressed (r and p_r scales are one-tenth those in Figure 3.2). However, it is clear that the slow variables are exciting the fast variables as there is a large peak at f = 5, although the amplitudes are much smaller than in Figure 3.2).

The gure showing the NNMI case (Figure 3.4) is even better (p_r is $\frac{1}{4}$ the size than in Figure 3.3) since high frequency oscillations have been massively damped. There is a peak in the fourier transforms in both the LNMI and NNMI graphs at f = 1 = 2f. This is because the spring is stretched twice per angular cycle at the bottom of the swing. This is called 'balanced fast motion'.

3.3.3 Summary of the numerical integration of the swinging spring and NMI

The swinging spring equations were integrated using the standard Matlab ode45 function, as done by Livings [2005]. This uses an explicit Runge-Kutta (4,5) pair with a changeable step-size. It was important to choose a small enough step-size to ensure stability. As discussed by Livings [2005] this had to be smaller than $\frac{T}{20}$, where T is the period of the harmonic oscillator.

Initialization aims to prevent unwanted high-frequency noise from developing in a solution. Linear Normal Mode Initialization (LNMI) aims to get rid of the high frequency oscillations by setting their initial amplitude to zero. Nonlinear Normal Mode Initialization (NNMI) instead sets the initial rates of change of the fast variables to zero (Lynch, 2002). Both were imposed for the swinging spring equations using the method and initial conditions described by Lynch [2002]. It was discovered that NNMI was better at supressing the high-frequency oscillations due to the non-linear nature of the system.



Figure 3.3: Coordinates and their fourier transforms of slow and fast variables for a swinging spring with linear normal mode initialization. Parameter values give rotational and elastic frequencies of f = 0.5 and $f_r = 5$ respectively. Initial conditions are $(;p;r;p_r) = (1;0;1;0)$.



Figure 3.4: Coordinates and their fourier transforms of slow and fast variables for a swinging spring with nonlinear normal mode initialization. Parameter values give rotational and elastic frequencies of f = 0.5 and $f_r = 5$ respectively. Initial conditions are $(; p; r; p_r) = (1; 0; 0.99540; 0)$.

Chapter 4

Implementing the ETKF, ETKS and ETKS2

then the eigenvalue decomposition is

$$(\hat{Y}^f)^T \hat{Y}^f = U \quad U^T \tag{4.3}$$

where *U* is an *N N* orthogonal matrix. It is possible to avoid performing $(Y^f)^T Y^f$ and use a simpler matrix in the analysis equation, which is more e cient to compute. It also reduces the errors in the smallest eigenvalues (Golub and Van-Loan, 1996). This is done by using the Singular Value Decomposition (SVD) of Golub and Van-Loan [1996]:

$$(\hat{Y}^f)^T = U \quad V^T \tag{4.4}$$

Here, is the *N* p diagonal matrix satisfying = ^{*T*} and *V* is an p p orthogonal matrix. The ensemble perturbation matrix is updated by

$$X^{a} = X^{f}T = X^{f}U(I +)^{\frac{1}{2}}U^{T}$$
(4.5)

The matrix U^T has been added to the version in (Livings, 2005) to make it unbiased (Livings et al., 2008). The ensemble Kalman gain can now be written as

$$\begin{aligned} \mathcal{K}_{e} &= X^{f}(Y^{f})^{T}(Y^{f}(Y^{f})^{T} + R)^{-1} \\ &= X^{f}\hat{Y^{f}}^{T}(\hat{Y}^{f}(\hat{Y}^{f})^{T} + I)^{-1}R^{-\frac{1}{2}} \\ &= X^{f}U^{-(T-F)}(I^{T} + I)^{-1}V^{T}R^{-\frac{1}{2}}. \end{aligned}$$
(4.6)

The product vector

$$\underline{Z} = \begin{pmatrix} T \\ + I \end{pmatrix}^{-1} V^{T} R^{-\frac{1}{2}} (\underline{y} \quad \underline{y}^{f})$$
(4.7)

is built from right to left. This avoids storing the large K_e matrix. Also, it involves the

error over the time window should be less for the ETKS2. Both the ETKS and ETKS2 implementations have loops which perform a re-analysis process. Firstly it is important to have calculated the analyses and forecasts for the ETKF in the time interval, as this is used as initial data. The implementation for the ETKF is given in the previous section 4.1.

4.2.1 Description of the ETKS and ETKS2 algorithms

The time interval is [0; T] with *I* steps. Assume *H* does not vary with time. The subscript *i* represents the ensemble members i = 1/2/3

For the ETKS algorithm the new forecast ensemble members are then updated using the smoothed analysis as

Calculate the singular value decomposition (SVD) of $Y_{k+ljk+l-1}^{f}$:

$$(U; ; V) =$$

over 100 runs. The ensemble mean averaged over 100 runs is given as $\overline{\chi^a_{k;100}}$ (the *i* subscript is no longer there). The analysis ensemble mean averaged over 100 runs and averaged over the time-interval is given by $\overline{\chi^a_{k;100}}$, where \overline{k} is the average over the time-interval. $\overline{\chi^a_{k;100}}$ gives a single value in each coordinate.

The analysis ensemble mean absolute error is de ned as

$$\underline{\underline{e}}_{k}^{a} = (j\underline{\underline{x}}_{k}^{a} \quad \underline{x}_{k}^{t}j):$$
(4.15)

Using the same notation as $\underline{x}^{\underline{a}}_{\overline{k},100}$, the ensemble mean absolute error averaged over 100 runs and averaged over the time interval is $\underline{\underline{e}}^{\underline{a}}_{\overline{k},100}$ and gives a single value in each coordinate.

Statistical signi cance

The statistical signi cance of the analysis ensemble mean error averaged over 100 runs was

to calculate the expensive Kalman gain matrix. The ETKS and ETKS2 algorithms use the ETKF Iter run as initial conditions. Then the analyses are smoothed using future observations. The implementation for the smoothing process uses the features of the ETKF implementation to make it more e cient, such as he tproduct vector. The key di erence between the ETKS and ETKS2 algorithms is that the ETKS algorithm updates the forecast states from the smoothed analyses. This is important in calculating the smoothed innovation vector and square root matrix. The ETKS 2 uses the original Itered forecast state in the innovation vector and square root matrix.

Chapter 5

Results

The methodology described in chapter 4 is now applied to the Ensemble Transform Kalman Filter (ETKF - see section 4.1) and the Ensemble Transform Kalman Smoother 2 (ETKS2 - see section 4.2.3). The ETKS2 di ers from the ETKS (section 4.2.2) in that it does not smooth the forecast state in the innovation vector and square root matrix *T* but instead uses the Itered version. The NNMI trajectory from section 3.3 is used as the true trajectory. The model used in the forecast step is the same as the model used to generate the true trajectory. The ETKS2 is validated by checking the nal ensemble values for a seeded initial distribution are equivalent to the nal ensemble values of the ETKF. Also the ensemble mean error averaged over the time interval should be smaller for the ETKS2. The experiments di er in the analysis step, the ensemble size and whether the observations are perfect (noise free) or imperfect. The xed-lag smoother ETKS2 is also tested with di erent lags.

5.1 Comparison of the ETKF with xed-interval ETKS2 using perfect observations

5.1.1 ETKF

The following application of the ETKF uses the programme created by Livings [2005]. Firstly, the results using the ETKF will be observed for frequent, perfect observations on all four coordinates ($;p;r;p_r$). The rst observation is at time 0.1 and all subsequent observations have intervals of 0.1. Since the observations are perfect, the actual observation

true solution. The covariance matrix R is also used in generating the initial ensemble. An ensemble of pseudo random vectors is drawn from a normal distribution with the given covariance matrix as its covariance matrix and the true initial state as its mean. This ensemble is then translated slightly so that the ensemble mean coincides exactly with the initial state. This translated random ensemble is used as an initial ensemble (Livings, 2005).

For a fair comparison between the ETKF and ETKS2 it is necessary to use an identical (seeded) initial ensemble distribution for both runs. The result of an experiment with ensemble size N = 10 is shown in Figure 5.1. The graph shows the di erence between the ETKF analysis and the truth for all four coordinates. Error bars show the observations (centred on the true solution) with the radius representing the standard deviation passed to the Iter. The observations follow the truth since they are perfect. There is a decrease in the amplitudes of the Iter error over time for most of the ensemble members, most signi cantly over the rst few seconds. This is expected since the analysis becomes more accurate (close to the true solution) as more observations are assimilated. The values of the

E8s2=1t4sue3337C(liss)=Iy62gra:the11(O7(are)-268(assarly742(er)-4ly62tion)-358(py74ed)]TJ 1(issimipy745 Td [i

Ensemble member		р	r	p _r
1	-0.0170	0.0413	0.0004	-0.0001
2	0.0029	-0.0181	0.0000	-0.0004
3	-0.0083	0.0051	0.0001	0.0022
4	-0.0045	0.0083	0.0001	-0.0004
5	0.0038	0.0049	0.0001	0.0005
6	0.0055	-0.0080	0.000	-0.0001
7	0.0076	-0.0299	-0.0001	-0.0012
8	0.0050	-0.0001	0.0000	-0.0012
9	-0.0021	0.0191	0.0001	0.0017
10	0.0001	0.0116	0.0001	-0.0013

Experiment (perfect obs)			р			r	р	r
ETKF $\underline{e}_{\overline{k};\overline{100}}^{a}$	9 <i>:</i> 40	10 ³	2 <i>:</i> 53	10 ²	2 <i>:</i> 01	10 ⁴	1 <i>:</i> 43	10 ³
95% con dence	1 <i>:</i> 11	10 ⁴	2 <i>:</i> 81	10 ⁴	4 <i>:</i> 30	10 ⁷	3 <i>:</i> 44	10 ⁵
ETKS2 $\overline{\underline{e}_{k}^{a}}$	4:77	10 ³	1 <i>:</i> 34	10 ²	7 <i>:</i> 71	10 ⁵	7 <i>:</i> 48	10 4
95% con dence	7 <i>:</i> 31	10 ⁵	2 <i>:</i> 09	10 ⁴	1 <i>:</i> 43	10 ⁶	1 <i>:</i> 70	10 ⁶

Table 5.4: Time averaged errors of the ensemble mean the 95% con dence range. Computed from 100 runs of the ETKF and ETKS2 with di erent random initial conditions and perfect observations.

5.1.2 Fixed-interval ETKS2

The following application of the ETKS2 uses the implementation in section 2.9.2 and smoothes using all the available obervations, hence it is xed-interval. The ETKS2 uses the ETKF Iter runs calculated in the programme created by Livings [2005]. A new section of the programme calculates the smoothed analyses. Using the same seeded distribution as used for the ETKF in 5.1, the results for the ETKS2 run are shown in Figure 5.3. Clearly the initial errors for the ETKS2 are much less. Also there is no general trend in the magnitude of the errors in the time window. The magnitudes of the errors for one ensemble member (blue line) appear to increase over the time interval but this is an isolated case. At the end of the time window the ETKS2 errors for each ensemble member are the same as the ETKF errors. This is shown by comparing table 5.2 with table 5.5. This is the expected result since both the nal analysis of the ETKF and the nal analysis of the ETKS2 are calculated from the same forecast state and observations.

The standard deviation of the analysis ensemble for the ETKS2 averaged over 100 runs was averaged over the time interval for all four coordinates. The values are shown in table 5.3. Compared with the ETKF standard deviations in table 5.3 these are at least an order

averaged errors of the ensemble mean for the ETKS2 are smaller than for the ETKF in all four coordinates. The ETKS2 errors are all approximately half the size of the ETKF errors. Thus the ETKS2 solution is smoother and more accurate on average than the ETKF one.

Ensemble member		р	r	p _r
1	-0.0170	0.0413	0.0004	-0.0001
2	0.0029	-0.0181	0.0000	-0.0004
3	-0.0083	0.0051	0.0001	0.0022
4	-0.0045	0.0083	0.0001	-0.0004
5	0.0038	0.0049	0.0001	0.0005
6	0.0055	-0.0080	0.000	-0.0001
7	0.0076	-0.0299	-0.0001	-0.0012
8	0.0050	-0.0001	0.0000	-0.0012
9	-0.0021	0.0191	0.0001	0.0017
10	0.0001	0.0116	0.0001	-0.0013

Table 5.5: Errors of the ETKS2 10 ensemble members for the seeded distribution at t=6 using perfect observations.

5.2 Comparison of the ETKF with the ETKS2 using imperfect observations

5.2.1 ETKF

The following imperfect method for the system is equivalent to the method implemented by Livings [2005]. So far in the experiments, the observations have been noise-free, frequent and made on all four coordinates of the system. For the next experiment, these assumptions are relaxed so as to see the e ect on the ensemble statistics. For the next experiment the time interval between observations has been increased to 0.37. Not only is this larger than the previous interval of 0.1 but it has been chosen as it is not a sub-multiple of the oscillation periods T = 2 and $T_r = 0.2$. Instead of observing all four coordinates only is observed. The observation error standard deviation added to the Iter for is 0.1 as before, but now errors of this magnitude are really added to the observations. The initial ensemble is generated using a diagonal covariance matrix corresponding to the standard deviations. The standard deviations for the other coordinates are equivalent to the amplitudes of the uninitialised oscillations in Figure 3.2. The idea is that the initial ensemble is not in uenced by these coordinates. The initial ensemble is generated using pseudo-random vectors as

with the perfect observations. However, there is no nal translation to make the ensemble mean coincide exactly with the true initial state.

Firstly an ETKF run with a seeded initial ensemble distribution will be looked at for 10 ensemble members. This seeded distribution is the same used for the perfect case (Figure 5.1). It is shown in Figure 5.5. It is obvious when comparing gure 5.5 with 5.1 that the amplitudes of the errors are much larger for the imperfect case. Also, the errors do not decrease over time in the fast coordinates as they do for the perfect case, instead they continue to oscillate at a similar magnitude. In the slow coordinates there is a slight decrease in the errors over time. The timescale for the fast coordinates is also much slower in the imperfect case than the perfect case.

The time averaged analysis ensemble standard deviation over 100 runs for the ETKF using imperfect observations is shown in table 5.3. Compared with the ETKF using perfect observations the standard deviations of all four coordinates are higher. The higher standard deviations in the imperfect casvations

Coordinate	Standard deviation
	0.1
p	3
r	0:06
pr	1 <i>:</i> 5

Table 5.7: Observation error standard deviations passed to the lter in experiments with imperfect observations. These standard deviations are also used to generate the initial ensemble.

5.2.2 Fixed-interval ETKS2

Using the seeded initial distribution, the ETKS2 was run using imperfect observations. The ETKS2 run in Figure 5.7 shows no signi cant change in the error magnitudes over the time window. The errors are also smaller across all 4 coordinates than for the ETKF run in Figure 5.5.

Figure 5.8 shows the absolute value of the error over time using the ensemble mean smoothed analysis averaged over 100 Iter runs for the ETKS2 for the imperfect observations. Comparing Figures 5.8 and 5.6 it is noticeable that the ETKS2 errors are only signi cantly better than the ETKF errors for the slow coordinates. This would make sense sinse the slow coordinate is the only variable being observed. Hence the smoothing process in ETKS2 is going to be more bene cial in the slow coordinates since actual future observations are being assimilated. However, the acccuracy of the slow variables will have some in uence on the accuracy of the fast variables in the equations of motion (section 3.1). Comparing Figure 5.8 with Figure 5.4 shows that the ETKS2 has also lost a lot of accuracy when using imperfect observations instead of perfect ones. Table 5.6 shows that the time averaged errors of the ensemble mean for the ETKS2 are smaller than for the ETKF in all four coordinates when using imperfect observations. However, as discussed this is only signi cant for the slow coordinates.

5.2.3 Ensemble error of the ETKF and ETKS2 using di erent ensemble sizes with imperfect observations

Table 5.8 shows the time averaged errors of the ensemble mean (averaged over 100 runs) in all four coordinates of the ETKF for 2-10 ensemble members. The ETKS2 version is shown in table 5.9. As expected the ETKS average errors are smaller than the ETKF ones.

Both the ETKF and ETKS2 errors in all four variables decrease signi cantly when increasing the ensemble size from two to four. When the ensemble size is too small in

relation to the model problems can develop in the Iter. The ensemble must adequately span the model subspace. The smaller the ensemble is, the greater the chance of sampling errors and underestimated forecast error covariances or imbreeding (Ehrendorfer, 2007). Thus increasing the number of ensemble members from two to four improves the accuracy in the four-dimensional swinging spring model. However, the errors decrease less when increasing the size from four to six and when increasing the ensemble size further than 6 the average errors in all four coordinates are generally getting worse (see tables 5.8 and 5.9). Thus increasing the ensemble size above the number of model variables eventually increases the error. The following reasons given come from (Nichols, 2009). The state of the system lies in an *n*-dimensional space. The estimate of the state obtained by the ensemble method lies in the *p*-dimensional space spanned by the ensemble members. When p > n, then you have spanned the full space in which the states lie. As you increase the number of ensembles, you are adding dependent vectors to the basis for the space in which the state estimate lies. It is possible therefore that you end up using a very poor selection of vectors from the basis (nearly dependent) to form your new state estimate. The more vectors added to the basis (the larger the number of ensembles) the more likely this is to happen. This e ect is exacerbated by the round-o errors introduced in the calculations. These mean that your computation of the state estimate can be very poor, even if the basis is not a linearly dependent set, but is close to being dependent. Round-o error can also cause the basis derived from the ensembles to become dependent before you have reached the point where p > n, even though in exact arithmetic this should be independent.

As well as measuring the a ect of increasing the ensemble size on the error, it is important to see if the ensemble standard deviation is behaving as it should be. However, this was only looked at for the ETKF. Using the Matlab programme of Livings [2005] it was possible to calculate the fraction of analyses having an ensemble mean within one standard deviation of the truth for each coordinate, averaged over 100 runs. For unbiased, normally distributed analysis errors with standard deviation equal to the ensemble standard deviation, this fraction should be about 0.68 (see Kreyszig, 1999 Appendix 5, table A7). If the fraction is below 0.68 this suggests that the standard deviation could be too small. Table 5.10 shows that this fraction increases signi cantly for larger ensemble sizes and goes from being well below 0.68 in all coordinates to being well above this fraction in all 4 coordinates. However, the very large fractions of 1.00 seen in the fast coordinates may be due to ignorance of the imperfect ETKF in the fast coordinates as the fast coordinates are not observed (Livings, 2005). It is above 0.68 in all coordinates for ensemble sizes of 6 or larger.

Ensemble size		р	r	pr
2	0.1183	0.3464	0.0580	1.8381
4	0.0592	0.1823	0.0411	1.3073
6	0.0584	0.1795	0.0382	1.2170
8	0.0589	0.1818	0.0398	1.2672
10	0.0703	0.2124	0.0348	1.0880
50	0.0604	0.1873	0.0427	1.3580

Table 5.8: Time averaged errors of the ensemble mean for di erent ensemble sizes. Ensemble mean averaged from 100 runs of the ETKF with di erent random initial conditions.

Ensemble size		р	r	p _r
2	0.1156	0.3122	0.0553	1.7489
4	0.0406	0.1103	0.0395	1.2478
6	0.0390	0.1055	0.0355	1.1707
8	0.0391	0.1066	0.0394	1.2484
10	0.0425	0.1201	0.0336	1.0589
50	0.0392	0.1047	0.0426	1.3529

Table 5.9: Time averaged errors of the ensemble mean for di erent ensemble sizes. Ensemble mean averaged from 100 runs of the ETKS2 with di erent random initial conditions.

5.2.4 ETKS2 with varying lags using imperfect observations

So far, the ETKS2 experiments have been of xed-interval type, smoothing the analyses using all available observations in the time window. Now the ETKS2 will be run for varying lags. Figure 5.9 is a graph plotted with one quarter the lag of the xed-interval problem. In other words the analyses for the ETKS2 will be smoothed using a maximum of 4 future observations instead of a maximum of 16 observations (all observations available) in the xed-interval problem. The graph shows that the errors are signi cantly higher in the slow coordinates at the beginning of the time window, but are more even in the fast coordinates. This is expected since the forecast is carrying forward less information from

observations from the past earlier in the time-window. The e ect is less noticeable in the fast coordinates since they are not observed. Table 5.11 shows the absolute value of the error over time using the ensemble mean smoothed analysis averaged over 100 runs for the di erent lags. Halving the lag from 16 to 8 increases the average error, except in the component. Decreasing the lag further increases the error in all four coordinates. This is expected since less observations are being assimilated at the beginning of the time-window for lower lags.

Ensemble size		D	r	Dr
2	0.282	0.281	0.0678	0.0679
4	0.640	0.634	0.530	0.530
6	0.699	0.690	0.785	0.786
8	0.722	0.717	0.906	0.906
10	0.0701	0.700	0.954	0.954
50	0.754	0.764	1.000	1.000

than about 6 could mean the standard deviation is too small for the ETKF. However, the very large fractions of 1.00 seen in the fast coordinates for large ensemble sizes may be due to ignorance of the imperfect ETKF in the fast coordinates as the fast coordinates are not observed (Livings, 2005)

Decreasing the lag for ETKS2 caused the average errors to increase. This is expected since the forecast is carrying forward information from less observations in the analysis step near the beginning of the time window.



Figure 5.1: ETKF, perfect observations, 10 ensemble members. Coordinates are plotted relative to the truth. Lines showing individual ensemble members are superimposed on observations plotted as error bars. Radius of error bars equals standard deviation of Iter. The error bars form a vertical grid centred on zero since the observations are frequent and perfect.



Figure 5.2: ETKF, perfect observations, absolute error trajectory of the ensemble mean (N = 10), averaged over 100 Iter runs. Coordinates are plotted relative to the truth. Lines showing individual ensemble members are superimposed on observations plotted as error bars. Radius of error bars equals standard deviation of Iter. The error bars form a vertical grid centred on zero since the observations are frequent and perfect.


Figure 5.3: ETKS2, perfect observations, 10 ensemble members. Coordinates are plotted relative to the truth. Lines showing individual ensemble members are superimposed on observations plotted as error bars. Radius of error bars equals standard deviation of Iter. The error bars form a vertical grid centred on zero since the observations are frequent and perfect.



Figure 5.4: ETKS2, perfect observations, absolute error trajectory of the ensemble mean (N = 10), averaged over 100 smoothed runs.



Figure 5.5: ETKF, imperfect observations, 10 ensemble members. Coordinates are plotted relative to the truth. Plotting conventions as for 5.1. Note that only the component has error bars since this is the only variable observed.



Figure 5.6: ETKF, imperfect observations, absolute error trajectory of the ensemble mean (N = 10), averaged over 100 smoothed runs. Plotting conventions as in Figure 5.2. Only the rst graph shows error bars since only the rst coordinate is observed.





Figure 5.8: ETKS2, imperfect observations, absolute error trajectory of the ensemble mean (N = 10), averaged over 100 smoothed runs. Plotting conventions as in Figure 5.2. Only the rst graph shows error bars since only the rst coordinate is observed.



Figure 5.9: ETKS2-lag 4, imperfect observations, absolute error trajectory of the ensemble mean (N = 10), averaged over 100 smoothed runs. Plotting conventions as in Figure 5.2. Only the rst graph shows error bars since only the rst coordinate is observed.

Chapter 6

Conclusions and Future work

6.1 Summary and discussion

The goals for this project were to compare two di erent sequential data assimilation methods, the Ensemble Transform Kalman Filter (ETKF) of Bishop et al. [2001] and the Ensemble Transform Kalman Smoother type 2 (ETKS2) introduced in this report (see section 2.9.2). The aim was to compare the errors of the ensemble members for the ETKF and ETKS2 for perfect (noise-free) and imperfect observations. In the imperfect case, di erent ensemble sizes and di erent lags for the smoother are looked at.

Chapter 2 introduces the Kalman Filter (KF) of Kalman [1960] and the formulations of the KF including the ETKF that is used for experiments in this report. In a lter a forecast model is used to produce a forecast state estimate using the analysis (best estimate of the system) of the previous time. This forecast state is updated to produce the current analysis by giving a weighting to the observations according to the ratio of estimated observation covariances and forecast covariances. Two types of ensemble lters are introduced, stochastic and deterministic, which di er in the analysis step. The main advantage of the deterministic lter is that perturbed observations are not used, which introduce additional noise. The Ensemble Kalman Filter (EnKF) of Evensen [1994] (see and the square root smoother of Whitaker and Compo [2002]. The smoothing method used in the experimental part of this report is called ETKS2 (see section 2.9.2) and di ers slightly from the ETKS (section 2.9) in that a xed forecast state is used in the innovation vector and square root matrix.

Chapter 3 discusses the two-dimensional swinging spring model. The concept of initialization is introduced and normal mode initialization is implemented for the swinging spring experiment of Lynch [2002]. Non-linear Normal Mode Initialization (NNMI) and Linear Normal Mode Initialization (LNMI) results are compared. Due to the non-linear nature of the system, NNMI is better than LNMI at removing the high frequency oscillations in the fast variables.

Chapter 4 discusses the implementation of the ETKF, the ETKS and the ETKS2. The formulations are equivalent to the formulations in Chapter 2ionsnutto the implementatioe

that ofvions[th[2502].462(Algorithmsre)-32quaregquin72enrefctor the implementation of the ETKs

e45iskal**fa(pisoft)365fit5a(maninautebas)&37210/556n(n**ation)]TJ0-16.435 Td[(v)28(ector)1334(and)1233(square)133rthronomatrinde ETxke®ivatiodectasmon expted tobr8orexpv-s 100 runs and averaged over the time interval. The statistical signi cance test measured the con dence in the analysis errors for the 100 run average. The mean ensemble absolute errors averaged over the 100 runs were less for the ETKS2 than the ETKF, proving that the smoother has improved accuracy over the Iter. Also since the 100 run average standard deviations were less for the smoother, there was less variation in the ensemble spread. However, the ETKF is known to underestimate ensemble spread so a smaller standard deviation is not necessarally a good thing for either the ETKF or ETKS2. Ensemble mean absolute errors averaged over 100 runs were plotted over the time interval for both the ETKF and ETKS2. The ETKF errors were worse at the beginning of the time window. This was expected since the Iter error decreases as more observations are assimilated. The ETKS2 errors were consistent over the time interval, as expected since the xed-interval smoother assimilates all available observations at each analysis step.

Imperfect observations were tested for both the ETKF and ETKS2 in the same way as the perfect observations. Only the slow coordinate was observed. The analysis errors

xed-lag ETKS2 than the ETKF in most of the analysis states.

6.2 Limitations of the experiments

6.2.1 The Swinging Spring system

There are some limitations in the experiments in this report. Firstly, the swinging spring system has only four parameters. In Numerical Weather Prediction the number of parameters p of the model is of $O(10^7)$. The number of ensemble members $N \ll p$. In order to relate the experiments for the smoother to NWP a much higher dimensional system would be needed. For example, the square root smoother of Whitaker and Compo [2002] gives results using a 40-dimensional model. Using this model it is discovered that sampling errors in the cross-covariance matrix are produced by higher lags. Thus higher ensemble sizes are needed to take advantage of observations further removed from the analysis. The four-dimensional swinging spring does not have a large enough range of ensemble sizes for $N \ll p$ to nd a relationship between lag, sampling errors and ensemble size.

6.2.2 Experimental work

In the experimental work in this report, only two di erent cases were considered for the ETKF anf ETKS2. The perfect case was rst looked at, using noise-free, high frequency and observations in all four parameters. Then the imperfect case looked at noisy observations at a lower frequency and with only the component observed. It would have been better to look at the imperfect case with two and three parameters observed rst to see the impact on changing things one by one. This could 495(c)27(hangegpring)-dae.sen-saa. g435 Tdi(Th7.e)5one. a

accurate than the ETKS2 results since the square root matrix T uses smoothed forecast states for each time step instead of using only the original litered forecast states. However the extra smoothing of the forecast states would make it more expensive to implement.

spacial and time scales. However, in certain situations it may be necessary to predict high frequency variations, such as localised mountain updraughts where the rising condensing air could cause cloud hazardous to aircraft.

The low frequency variations that are of no interest are called noise. If noisy observations are incorporated into the basic equations in numerical prediction the forecast may contain spurious large amplitude high frequency oscillations (Lynch, 2002). This is because the balance between the mass and velocity elds is not fairly re ected in the forecast. In other words, the numerical model cannot cope with the high frequency variables. The elimination of this noise can be achieved by adjusting the initial elds, a process called initialization. Detail on di erent forms of initialization that can be found is in Lynch [2002]. Initialization is an important part of data assimilation. Any linear balances in a system should be preserved with the ETKF since the ETKF takes a linear combination of the ensemble members to create the analysis. However, a smoother would be expected to give a smoother solution that ts the observations better. Thus a smoother might be expected to dampen the high frequency and high amplitude errors in the solution more than the lter. The relationship between initialization, the ETKS/ETKS2 and ETKF could be investigated using the Swinging Spring model or another model by measuring the spread of the errors in the ensemble members in the solutions. This could be done by plotting the ensemble mean and the ensemble mean standard deviation for the ETKS/ETKS2 and comparing the results with the ETKF plots from Livings [2005]. Neef et al. [2005] investigate initialization properties of a stochastic EnKF using a four-dimensional system. Perhaps the deterministic ETKF, ETKS and ETKS2 could be applied using similar experiments.

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