Subgrid-Scale Parameterization with Conditional Markov Chains

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ABSTRACT

A new approach is proposed for stochastic parameterization of subgrid-scale processes in models of atmospheric or oceanic circulation. The new approach relies on two key ingredients: first, the unresolved processes are represented by a Markov chain whose properties depend on the state of the resolved model variables; second, the properties of this conditional Markov chain are inferred from data. The parameterization approach is tested by implementing it in the framework of the Lorenz '96 model. Performance of the parameterization scheme is assessed by inspecting probability distributions, correlation functions, and wave properties, and by carrying out ensemble forecasts. For the Lorenz '96 model, the parameterization algorithm is shown to give good results with a Markov chain with a few states only and to outperform several other parameterization schemes.

1. Introduction: Stochastic parameterization of subgrid-scale processes

The parameterization of subgrid-scale processes in models of atmospheric flow has drawn a lot of research attention recently. To get beyond the limitations of parameterizations with deterministic functions, the focus of various recent investigations has been on the potential of stochastic methods for the parameterization of processes that cannot be resolved because they fall below the model grid scale (e.g., Majda et al. 1999; Buizza et al. 1999; Lin and Neelin 2000; Palmer 2001; Lin and Neelin 2002; Majda and Khouider 2002; Majda et al. 2003; Khouider et al. 2003; Wilks 2005; Plant and Craig 2008).

When trying to parameterize unresolved processes stochastically, one is faced with two main issues. The first is to determine the class of models one wants to use for the subgrid processes. Several directions have been proposed and used in the literature, ranging from stochastic differential equations (Majda et al. 1999, 2003; Wilks 2005) to cellular automata (Shutts 2005; see also Palmer 2001), multiplicative randomization of deterministic parameterization schemes (Buizza et al. 1999), and Markov chain models on a discrete state space (Majda and Khouider 2002; Khouider et al. 2003). This last class of models using Markov chains is the one that we used for the present paper, as explained below in more detail.

The second issue one faces is how to choose the parameters in the model for the subgrid processes. This can be done either based on physical intuition (e.g., in Lin and Neelin 2000; Majda and Khouider 2002) or by directly using the data from the time series for the subgrid processes (e.g., Wilks 2005). The second approach has the advantage of typically allowing one to make less ad hoc assumptions about the subgrid processes. This may be less transparent from a physical perspective, but it is also potentially more accurate. The approach proposed in this paper uses the data of the subgrid processes to obtain a stochastic model. The related, more general problem of inferring stochastic models from data is considered in a wide variety of papers; for applications in atmosphere–ocean science, see, for example, Penland and Matrosova (1994), Egger (2001), Sura (2003), Crommelin (2004), and Berner (2005).

Specifically, we propose a strategy for stochastic parameterization that uses Markov processes that are conditional on the state of the resolved variables. As all stochastic parameterizations do, this strategy accounts for the possibility to have a multiplicity of possible states of the unresolved processes given one fixed state.

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of the resolved variables. It also accounts for the possibility that the properties of the unresolved processes (e.g., variance, skewness, and decorrelation time) will vary with the state of the resolved variables. We give an algorithm that translates this dependency into a computationally feasible parameterization scheme. The conditional stochastic processes are represented as Markov chains with a small number of states, making the practical implementation easy. The properties (such as transition probabilities) of the Markov chains are estimated from data in a simple, straightforward way. The algorithm is applied to the Lorenz ’96 (L96) model (Lorenz 1995), a frequently used test bed for parameterization strategies (Palmer 2001; Fatkullin and Vanden-Eijnden 2004; Wilks 2005).

We note that the types of models resulting from the stochastic parameterization developed in this paper have a structure resembling the coupled models proposed by Majda and collaborators in the context of material science (Katsoulakis et al. 2003, 2005, 2006) and focusing on tropical convection (Majda and Khouider 2002; Khouider et al. 2003). In those studies, a systematic subgrid-scale parameterization is proposed in which deterministic equations for macroscopic variables are coupled to a stochastic Ising (spin flip) system that represents microscopic phenomena. The spin-flip system is a Markov chain that can be coarse grained using a systematic closure procedure. Similar to our approach, the resulting models consist of deterministic differential equations coupled to Markov chains that are conditional on the state of the macroscopic variables. The main difference is that in the present paper, the parameterization (or closure) is entirely inferred from data without using any knowledge of the physics or equations that drive the subgrid scales. In the approach by Majda and collaborators, on the other hand, one starts with an explicitly known microscopic model that can be coarse grained because the equilibrium distribution of the microscopic model is known exactly (under the assumption of detailed balance); there is no attempt to determine the parameters in the model from actual data.

The outline of the remainder of this paper is as follows: in section 2, we give a brief description of the Lorenz ’96 model. The stochastic parameterization strategy with data-inferred conditional Markov processes is introduced in section 3. In section 4, practical aspects of the parameterization scheme are discussed, such as data inference and integration scheme. We present numerical results in section 5, where we compare our scheme with other parameterization schemes by inspecting probability distributions, correlation functions, wave statistics, and results from ensemble integrations (both ensemble mean and dispersion). We conclude in section 6 with a discussion, where we also address the question of how the stochastic parameterization scheme could be used in more realistic models.

2. The Lorenz ’96 model: A test bed for parameterization algorithms

The two-layer Lorenz ’96 model, proposed by Lorenz in 1995 (Lorenz 1995), has become a popular toy model of the atmosphere to test various concepts and ideas relating to predictability, model error, and parameterization (Boffetta et al. 1998; Palmer 2001; Orrell 2003; Fatkullin and Vanden-Eijnden 2004; Wilks 2005). The model equations read

\[ \dot{X}_k = X_{k-1}(X_{k+1} - X_{k-2}) - X_k + F + B_k \]  
\[ \dot{Y}_{j,k} = \frac{1}{\varepsilon} \left[ Y_{j+1,k}(Y_{j-1,k} - Y_{j+2,k}) - Y_{j,k} + h_k X_k \right], \]

in which

\[ B_k = \frac{h_k}{\varepsilon} \sum_{j=1}^{J} Y_{j,k}, \]

and \( k = 1, \ldots, K \); \( j = 1, \ldots, J \). The \( X_k \) and \( Y_{j,k} \) are interpreted as variables on a circle of constant latitude, where the \( X_k \) are “large-scale” variables, each coupled to a collection of \( J \) “small-scale” variables \( Y_{j,k} \). The indices \( k \) and \( j \) can be regarded as spatial indices. The periodicity of the spatial domain is reflected in the periodicity of the variables

\[ X_k = X_{k+K}, \]
\[ Y_{j,k} = Y_{j,K+k}, \]
\[ Y_{j+J,k} = Y_{j,k+1}. \]

In Lorenz (1995), the model is formulated slightly different from (1), with parameter settings equivalent to \( \varepsilon = 0.1, K = 36, J = 10, F = 10, h_k = -1, \) and \( h_j = 1 \). We use the formulation above (from Fatkullin and Vanden-Eijnden 2004) because it makes the time scale separation (measured by \( \varepsilon \)) between the \( X_k \) and the \( Y_{j,k} \) explicit. If \( \varepsilon \ll 1 \), the \( X_k \) are slow variables and the \( Y_{j,k} \) are fast; if \( \varepsilon \approx 1 \), there is no time scale separation. Studies using the L96 system are often carried out using parameter settings that amount to such time scale separation (Lorenz 1995; Fatkullin and Vanden-Eijnden 2004; Wilks 2005). In this paper, we use \( \varepsilon = 0.5 \) because (near) absence of time scale separation between resolved and unresolved processes is both more realistic and more difficult to handle for parameterizations. See Fatkullin and Vanden-Eijnden (2004) and references therein for an overview (and implementation for L96).
of recently developed computational strategies to handle the case of \( \varepsilon \ll 1 \).}

Using data from a numerical simulation of the full L96 system (1), a scatterplot of \( B_k \) versus \( X_k \) is shown in Fig. 1. The parameters used in the simulation are \( (\varepsilon, K, J, F, h_x, h_y) = (0.5, 18, 20, 10, -1, 1) \). Because of the translation invariance of the system, this scatterplot has the same statistical properties for all values of \( k \). From the figure, it is clear that for a fixed value of \( X_k \), \( B_k \) can take on a range of values. The properties of \( \rho(B_k|X_k) \), the probability density function (PDF) of \( B_k \) conditional on the value of \( X_k \), are obviously highly dependent on \( X_k \). In Fig. 2, various PDFs of \( B_k \) are shown, estimated from data points with \( X_k \) in different intervals.

3. Parameterization with conditional Markov processes

Within the context of the L96 system (1), the aim of a parameterization scheme is to formulate a model for the large-scale variables \( X_k \) alone, from which the variables \( Y_{k,k'} \) have disappeared entirely. The key element is a suitable representation of the quantities \( B_k \). In the full L96 model (1), they depend on the \( Y_{k,k'} \); in a reduced model for the \( X_k \) variables alone, they must be parameterized in terms of the \( X_k \).

One way of parameterizing is deterministic, with a function \( B = G(X) \). In its most general form, every element \( B_k \) of the vector \( B \) is determined by all elements \( X_k \) of the state vector \( X \). Because the function \( G \) can be nonlinear, this type of parameterization is often too complicated to be of practical use for systems with more than a few degrees of freedom [see, however, Fatkullin and Vanden-Eijnden (2004) for an on-the-fly computational strategy to handle this problem when \( \varepsilon \ll 1 \)]. For complex systems, a simplified function such as \( B_k = g(X_k) \) is typically considered. The assumption that \( B_k \) is determined by \( X_k \), not by variables at other grid points \( k' \neq k \), can be seen as a “locality” assumption.

A more fundamental problem of deterministic parameterizations stems from the chaotic nature of the underlying systems: a function such as \( B_k = g(X_k) \) cannot account for the possibility that in the full system, \( B_k \) can take on a variety of states given a fixed \( X_k \), rather than one unique state. Put differently, the probability distribution of \( B_k \) with \( X_k \) fixed is often not a Dirac delta distribution. It is therefore natural to consider stochastic parameterizations, in which the \( B_k \) are modeled as stochastic processes. They allow for different realizations of \( B_k \) for a fixed value of \( X_k \), consistent with the scatterplot of Fig. 1.

We propose here a new approach to stochastic parameterization, in which we make the following two key assumptions about the stochastic process that replaces \( B_k \): (i) it is a Markov process; (ii) the process is conditional on \( X \). We infer the properties of the stochastic process from the \( (X_k, B_k) \) data of the full L96 model. To make this parameterization scheme computationally as simple as possible, we restrict the conditionality in practice to \( X_k \). Adding conditionality on other grid points (e.g., the nearest neighbor grid points \( X_{k-1} \) and \( X_{k+1} \)) can be expected to improve the performance of the reduced model; however, it will also
make the model parameterization more complicated. We leave exploration of this possibility to another study.

In a nutshell, we model $B_k$ by numerical (Monte Carlo) simulation of the stochastic process with the conditional transition probability

$$P[B_k(t_2)|X_k(t_2), B_k(t_1), X_k(t_1)] \quad \text{with} \quad t_2 \geq t_1, \quad (4)$$

which will be estimated from the $(X_k, B_k)$ data (see section 4 for details). We let $X_k(t_2)$ be determined by integration of (1a), starting from $X(t_1)$ and with $B_k$ fixed at $B_k(t_1)$. Thus, given $X(t_1)$ and $B(t_1)$, first we integrate (1a) to obtain $X(t_2)$, then we obtain $B(t_2)$ by Monte Carlo simulations of (4) for each $k$. More practical aspects of this parameterization scheme are discussed in the next section.

Adding $X_k(t_2)$ to the conditions $X_k(t_1)$ and $B_k(t_1)$ in (4) is natural because the distribution for $B_k(t_2)$ depends strongly on the direction in which $X_k$ is moving. Note that the value of $X_k(t_2)$ information is readily available via integration of (1a) on $t \in [t_1, t_2]$ with $B_k(t) = B_k(t_1)$. As an illustration, we calculated PDFs for $B_k$ on the interval $1.5 < X_k < 2.5$, depending on whether $X_k$ is growing or decreasing in value. More precisely, we calculated the PDFs $\rho[B_k(t)|1.5 < X_k(t) < 2.5, X_k(t) > X_k(t - \Delta t)]$ and $\rho[B_k(t)|1.5 < X_k(t) < 2.5, X_k(t) < X_k(t - \Delta t)]$. They are shown, together with the total PDF $\rho[B_k(t)|1.5 < X_k(t) < 2.5]$, in Fig. 3. The significant differences between these PDFs make clear why a parameterization with the conditional transition probability $P[B_k(t_2)|B_k(t_1), X_k(t_1)]$ instead of (4) would result in a less accurate model in the case of L96. We expect the inclusion of $X_k(t_2)$ as a condition to be less important for models that are more strongly mixing, so that $P[B_k(t_2)|B_k(t_1), X_k(t_1)]$ is roughly the same when $X_k$ is growing as it is when $X_k$ is decreasing.

The parameterization we propose has some similarities with the stochastic parameterization for L96 studied in Wilks (2005). There, the $B_k$ are modeled by a deterministic term [a function $g(X_k)$ found by fitting a polynomial to the $(X_k, B_k)$ data] plus a stochastic term coming from an autoregressive [AR(1)] process. The AR(1) process itself is not conditional on $X_k$. However, because the stochastic term is added to a deterministic one, the scheme studied in Wilks (2005) is equivalent to parameterization with an AR(1) process with $X_k$-dependent mean. The parameterization approach proposed here is more general, because the stochastic process we use need not be of the AR(1) type and its conditionality on $X_k$ is not restricted to the mean but extends to, for example, variance, skewness and decorrelation time. In section 5, we will show that our approach outperforms the one proposed in Wilks (2005), at least for the example of L96 in the parameter regime that we consider.

4. A practical algorithm using Markov chains

In this paper, we choose to model the conditional Markov process that replaces $B_k$ as a collection of Markov chains. As a result, $B_k$ becomes a discrete random variable. One can also choose to model the $B_k$ with Markov processes that are continuous in space (e.g., diffusion processes); we use Markov chains here because of their computational ease. Even with a small number of Markov chain states, statistical properties such as variance, skewness, and decorrelation time can be captured. Moreover, data inference of Markov chains is much easier to carry out than inference of (possibly non-Gaussian) continuous stochastic processes from data. The conditionality on $X_k(t_1)$ and $X_k(t_2)$ is implemented by letting the properties of the Markov chain change stepwise (not continuously), depending on the intervals in which $X_k(t_1)$ and $X_k(t_2)$ reside. It must be stressed that although the $B_k$ have become discrete variables, and although the Markov chain properties change discretely rather than continuously with $X_k$, the resulting (reduced) model for the resolved variables (the $X_k$) is still continuous.

a. Estimation of the Markov chains

For the construction of the Markov chains, all data points $(X_k, B_k)$ are assigned to discrete states $(i, n)$ by
partitioning the \((X_k, B_k)\) plane into bins. First, the range of possible values of \(X_k\) is divided into \(N_X\) non-overlapping intervals \(I_i (i = 1, \ldots, N_X)\). Within each interval \(I_i\), the range of values of \(B_k\) is divided into \(N_B\) non-overlapping intervals \(J_{nm} (n = 1, \ldots, N_B)\). A set of stochastic matrices is constructed by estimating a spatially discrete version of the conditional transition probability \((4)\) from the data:

\[
\mathbf{P}^{(ij)}_{nm} = \mathbf{P}[B_k(t + \Delta t) \in J_{nm} | X_k(t) \in I_i, B_k(t) \in J_{in}, X_k(t + \Delta t) \in I_j],
\]

\[
= \frac{T_{in,jm}}{\sum_{m=1}^{N_B} T_{in,jm}}.
\]

The object \(T_{in,jm}\) counts the transitions from \((i, n)\) to \((j, m)\) in the data:

\[
T_{in,jm} = \sum_t \mathbf{I}[B_k(t) \in J_{in}] \mathbf{I}[X_k(t) \in I_i] \times \mathbf{I}[B_k(t + \Delta t) \in J_{nm}] \mathbf{I}[X_k(t + \Delta t) \in I_j],
\]

where \(\sum_t\) denotes the sum over all discrete times \(t = 0, \Delta t, 2\Delta t, \ldots\) in the dataset and \(\mathbf{I}[]\) denotes the indicator function \(\mathbf{I}[B_k(t) \in J_{in}] = 1 \text{ if } B_k(t) \in J_{in}\) and \(\mathbf{I}[B_k(t) \in J_{in}] = 0 \text{ if } B_k(t) \notin J_{in}\), and so forth. For simplicity, we assume that the time interval \(\Delta t\) between two data points is constant throughout the dataset.

For any \(i\) and \(j\) fixed, the matrix \(\mathbf{P}^{(ij)}\) satisfies

\[
\forall n, m: \mathbf{P}^{(ij)}_{nm} \geq 0, \quad \forall n: \sum_{m=1}^{N_B} \mathbf{P}^{(ij)}_{nm} = 1,
\]

so \(\mathbf{P}^{(ij)}\) is an \((N_B \times N_B)\) stochastic matrix (note that \(i\) and \(j\) themselves run from 1 to \(N_X\)). The discrete states \((i, n)\) are assigned a value \(b_{in}\) for \(B_k\) that is the average of all data points falling in bin \(J_{in}\):

\[
b_{in} = \frac{1}{\sum_t \mathbf{I}[B_k(t) \in J_{in}] \mathbf{I}[X_k(t) \in I_i]} \sum_t \mathbf{I}[B_k(t) \in J_{in}] \mathbf{I}[X_k(t) \in I_i] t.
\]

Construction of the transition probability matrices \(\mathbf{P}^{(ij)}\) is not possible if \(|i - j|\) is too large. The \((X_k, B_k)\) data do not contain pairs of consecutive points with \(X_k(t) \in I_i, X_k(t + \Delta t) \in I_j\) if \(i\) and \(j\) are far apart and the time step \(\Delta t\) is small. However, this is not a problem, because transitions between distant intervals \(I_i\) and \(I_j\) are also unlikely to occur during an integration of the reduced L96 model. In practice, the only matrices \(\mathbf{P}^{(ij)}\) that are really needed are those with \(|i - j| \leq 1\). If the \((X_k, B_k)\) data does not contain any transition out of \((i, n)\) into \(j\) for some combination of \(i, j, n\), then \(\Sigma_n T_{in,jm} = 0\). In that case, we set \(\mathbf{P}^{(ij)}_{nm} = \delta_{nm}\) for that particular triplet of \(i, j, n\) (where \(\delta_{nm}\) is the Kronecker delta). This avoids the numerical integration breaking off on the rare occasion that a transition occurs for which we have no data. A more sophisticated way of dealing with this problem involves summation over all intermediate steps \((i, n) \rightarrow (i', n') \rightarrow (j, m)\) with known transition probabilities. We will not do this here.

b. Time step of the Markov chains

The stochastic matrices \(\mathbf{P}^{(ij)}\) are made up of the transition probabilities \((5)\) over a fixed time interval \(\Delta t\), at which the \((X_k, B_k)\) data is sampled. However, in practical situations it is possible that the data available for the construction of a parameterization scheme is sampled with a time step \(\Delta t\) that is different from the time step \(\Delta t\) at which the numerical model will be integrated. For example, \(\Delta t\) can be so large that an integration scheme with \(\Delta t = \Delta t\) would be numerically unstable.

There are two ways of dealing with this problem of nonmatching time steps. One possibility is to use a split-integration scheme. Assume for simplicity that \(\Delta t\) is an integer multiple of \(\delta t\), say, \(\Delta t = N \delta t\). We integrate \(X_k\) with time step \(\Delta t\) and update \(B_k\) once every \(N\) time steps. Of course, the transition probability matrix \(\mathbf{P}^{(ij)}\) used to evolve \(B_k\) in time must have \(i\) and \(j\) such that \(X_k(t_1) \in I_i\) and \(X_k(t_2) \in I_j\) with \(t_2 - t_1 = N \delta t\), not \(t_2 - t_1 = \Delta t\).

Another possibility to tackle the problem is by calculating transition probability matrices \(\mathbf{P}^{(ij)}\) with an associated time step \(\delta t\), starting from matrices \(\mathbf{P}^{(ij)}\) with time step \(\Delta t\). Unfortunately, the straightforward but naive choice \(\mathbf{P}^{(ij)} = [\mathbf{P}^{(ij)}]^{\delta t / \Delta t}\) does not, in general, result in a new stochastic matrix (some matrix elements may be negative or complex) because of the Markov chain embedding problem (for a discussion, see Bladt and Sørensen 2005; Crommelin and Vanden-Eijnden 2006). To find a matrix \(\mathbf{P}^{(ij)}\) that is both a true stochastic matrix and approximately equal to \([\mathbf{P}^{(ij)}]^{\delta t / \Delta t}\), one should first construct a so-called generator matrix \(\mathbf{L}\) from \(\mathbf{P}^{(ij)}\). The \(\mathbf{L}\) must be such that (i) \(\exp(\Delta t \mathbf{L})\) resembles \(\mathbf{P}^{(ij)}\) as closely as possible and (ii) \(\exp(\delta t \mathbf{L})\) qualifies as a stochastic matrix for all \(\delta t \geq 0\). Two different methods for the numerical construction of generators can be found in Bladt and Sørensen (2005) and Crommelin and Vanden-Eijnden (2006). After \(\mathbf{L}\) is constructed, the matrix \(\mathbf{P}^{(ij)} = \exp(\delta t \mathbf{L})\) is a true stochastic matrix that approximates \([\mathbf{P}^{(ij)}]^{\delta t / \Delta t}\) as best as possible. The degree of proximity depends on how close \(\exp(\Delta t \mathbf{L})\) is to \(\mathbf{P}^{(ij)}\). The difference between \(\exp(\Delta t \mathbf{L})\) and \(\mathbf{P}^{(ij)}\) can be non-negligible for some \(\mathbf{P}^{(ij)}\), creating a source of error for the parameterization scheme. For these reasons, and
because the split-integration scheme described before gave good results for L96 (see section 5c), we will not investigate the generator method any further here.

c. Numerical integration

Having the transition matrices \( P^{(i)} \) [or \( \tilde{P}^{(i)} \)] for time interval \( \delta t \) available, time integration of the reduced L96 model with conditional Markov chain (CMC) parameterization is done as follows: given the vectors \( X(t) \) and \( B(t) \), the next time iterate \( X(t + \delta t) \) is calculated using the derivative of \( X \) determined by Eq. (1a), with \( B \) fixed at \( B(t) \). Then the time step for \( B_k \) is made by Monte Carlo simulation (independent simulations for different \( k \) of the Markov chain with the transition probability matrix \( P^{(i)} \), with \( i \) and \( j \) such that \( X_k(t) \in I_i \) and \( X_k(t + \delta t) \in I_j \). As mentioned before, the \( B_k \) have become discrete variables, because they can only take on a finite number of values (the \( g_{ij}^k \)). If \( B_k \) is sent from state \( (i, n) \) to state \( (j, m) \) by the Monte Carlo simulation, it means that the value of \( B_k \) to be used in the integration of \( X_k \) changes from \( g_{in}^k \) to \( g_{jm}^k \).

5. Numerical results

We compare the results generated by the reduced L96 model [Eq. (1a) without (1b)] using three different parameterization schemes for \( B_k \): deterministic (DTM), the AR(1) scheme as presented in Wilks (2005), and the CMC scheme described in the previous sections. We generate data for \( X_k \) and \( B_k \) by integrating the full L96 model (1) with parameters \( (\varepsilon, K, J, F, h_x, h_y) = (0.5, 18, 20, 10, -1, 1) \). Thus, there is hardly any time scale separation between the resolved and the unresolved variables, making the situation more realistic (and more difficult) than would be the case with the more common choice of \( \varepsilon = 0.1 \). We integrate the full L96 model for \( 10^5 \) time units with time step \( 10^{-3} \) and store the resulting \( (X_k, B_k) \) every \( 0.01 \) time unit (i.e., \( \Delta t = 0.01 \)). Thus, the dataset contains \( 10^5 \) points. The reduced models will be integrated with the same time step, \( \delta t = 0.01 \). In the last part of this section, we consider the case where \( \Delta t > \delta t \), which adds a complication as described in section 4b.

For the DTM parameterization scheme, a fifth-order polynomial is fitted to the \( (X_k, B_k) \) data. The resulting function \( B_k = g(X_k) \), shown in Fig. 4, is used as a parameterization of \( B_k \).

The AR(1) scheme from Wilks (2005) consists of representing \( B_k(t) \) by a deterministic term plus a stochastic term:

\[
B_k(t) \approx g[X_k(t)] + \xi(t). \tag{9}
\]

For \( g(X_k) \), we take the polynomial function also used for the DTM scheme. The stochastic term \( \xi \) is generated by an AR(1) process whose parameters are obtained by fitting the process to the time series of \( B_k(t) - g[X_k(t)] \). The data in this time series are sampled at a time interval of \( \Delta t = 0.01 \) and lead to an AR(1) process with mean zero, standard deviation of 0.88 and \( e \)-folding (decorrelation) time of 4.3. Thus, the AR(1) process can be regarded as red noise. For the CMC scheme, we have to choose the intervals \( I_i \) and \( g_{in}^k \). Once this is done, we construct the transition matrices \( P^{(i)} \) and the discrete values \( g_{in}^k \) for \( B_k \) from the \( (X_k, B_k) \) data, using expressions (5b), (6), and (8). For \( X_k \), we use 16 intervals of width 1, centered at

![Figure 4](image-url)

**Fig. 4.** (top) The solid curve is the fifth-order polynomial fit used for the DTM parameterization of \( B_k \). (bottom) The black squares denote the values of the \( g_{in}^k \) used for the CMC parameterization with \( N_a = 4 \) (see text). In both panels, the dots are the scatterplot of \( B_k \) vs \( X_k \) for the full L96 model (1).
integer values of $X_k$. At the beginning and end of the domain, we use unbounded intervals. Thus, $I_1 = (-\infty, -4.5]$, $I_2 = (-4.5, -3.5]$, $I_3 = (-3.5, -2.5]$, \ldots, and $I_{16} = (9.5, +\infty)$. Within each interval $I_i$, the bins $\mathcal{B}_n^i$ are determined such that each bin contains (approximately) an equal number of data points. We set the number of bins to $N_B = 4$ for all calculations. The resulting values of $\mathcal{B}_n^i$ are plotted in the lower panel of Fig. 4. Using $N_B = 8$ instead of $N_B = 4$ gave no significant improvement in the results and are therefore not shown.

**a. PDFs, correlations, and wave statistics**

We integrate the reduced L96 models with the various parameterization schemes and calculate several quantities from the data of the reduced models as well as from the data of the full L96 model. Each time series is $10^5$ data points long with sampling interval $\Delta \tau = 0.01$. We calculate the following quantities:

- the PDF for $X_k$;
- the autocorrelation function (ACF) for $X_k$;
- the cross-correlation function (CCF) for $X_k$ and $X_{k+1}$;
- by Fourier transforming the state vector $\mathbf{X}$ at every data point, a time series for the wavenumber vector $\mathbf{u}$ is obtained, from which we calculate the wave variance $\langle (u_m - \langle u_m \rangle)^2 \rangle$ for every wavenumber $0 \leq m \leq K/2$, where $\langle \rangle$ denotes time average; and
- the mean wave amplitude $\langle |u_m| \rangle$.

For calculation of the PDFs, ACFs, and CCFs, we take the average over all values of $k$. The PDF for $X_k$ of the full L96 model is well reproduced by all parameterization schemes (Fig. 5). Differences among the schemes are quite small here. The ACF (Fig. 6) and CCF (Fig. 7), both strongly oscillatory, are more accurately reproduced by the CMC scheme than by the others. All schemes give oscillatory ACFs and CCFs, but with the DTM and AR(1) schemes, the amplitude of the oscillation is too low and a phase shift can be seen. The wave variances and mean wave amplitudes of the full L96 model are also better reproduced with the CMC scheme than with the others (Fig. 8). The CMC scheme gives a correct peak in the variance at wavenumber $m = 3$; the other schemes show a lower, broader peak spread out over wavenumbers $m = 3$ and $m = 4$. A similar effect can be seen in the mean wave amplitudes.

In Wilks (2005), the PDF of $X_k$ is also rather well reproduced by various parameterization schemes (DTM, AR(1), with white noise and red noise), similar to what we find. Correlation functions and wave statistics, similar to our Figs. 6–8, are not calculated in Wilks (2005).

**b. Ensemble tests**

We carry out ensemble integrations for further testing of the CMC parameterization. For each ensemble, we calculate the mean trajectory and compare it with the true trajectory from the full L96 model. This is done by calculating the root-mean-square error (RMSE) and the anomaly correlation (ANCR) using results from many different ensembles. The number of ensemble members in each ensemble is 1, 5, or 20. For the 20-member ensembles, we also calculate rank histograms to show the ensemble dispersion.

The setup is as follows. Let $\mathbf{X}_{\text{full}}^t$ be a time series for
X of the full L96 model. A number of $N_{\text{init}}$ initial states from this time series is selected:

$$X_{\text{init},s} = X_{ts}^{\text{full}}, \; s = 1, \ldots, N_{\text{init}}.$$  \hspace{1cm} (10)

We take one data point every 10 time units from $X_{ts}^{\text{full}}$, so $t_{s+1} - t_{s} = 10$. For every initial state, we do $N_{\text{ens}}$ integrations of the reduced model over $T$ time units [using the CMC, DTM, or AR(1) parameterization], starting from $X_{\text{init},s}$ plus a small random perturbation $\xi^{s,n}$ ($n = 1, \ldots, N_{\text{ens}}$). We use random perturbations that are drawn from a Gaussian distribution with mean 0 and standard deviation 0.15 (for comparison: the standard deviation of $X_{k}$ is about 3.5 with the parameter settings we use for the full L96 model). We make no attempt to sample the unstable directions of phase space more heavily than other directions nor to generate ensembles using fastest-growing perturbations from singular-value decomposition. For the purpose of comparing different parameterization approaches, the simple generation of ensemble members described above is sufficient.

The different integrations from one ensemble are averaged, resulting in a mean trajectory $X_{t}^{\text{mean},s}$ [where $t \in (0, T)$ and $s = 1, \ldots, N_{\text{init}}$]. The RMSE measures the average difference between the mean ensemble trajectory and the trajectory from the full L96 model:

$$\text{RMSE}(\tau) = \left( \frac{1}{N_{\text{init}}} \sum_{s} |X_{t}^{\text{mean},s} - X_{t+\tau}^{\text{full}}|^2 \right)^{1/2}.$$  \hspace{1cm} (11)

For the anomaly correlation, we need the anomalies of the full L96 model and the ensemble mean:

$$a_{t}^{\text{full},s} = X_{t+\tau}^{\text{full}} - \langle X^{\text{full}} \rangle,$$  \hspace{1cm} (12)

$$a_{t}^{\text{mean},s} = X_{t}^{\text{mean},s} - \langle X \rangle.$$  \hspace{1cm} (13)
integrations at lead time \( \tau = 2 \). For each grid point \( k \), we rank the \( N_{\text{ens}} + 1 \) values for \( X_k^* \) from the ensemble members and the full L96 model. Ideally, the distribution of the position of the truth (the value from the full L96 model) in this ranking should approach a uniform distribution (Hamill 2001); in that case, the rank histogram is (nearly) flat. If the ensemble is underdispersed, the truth occupies the extremes (locations at or near 1 and \( N_{\text{ens}} + 1 \)) too often, leading to a U-shape rank histogram. In the reverse situation (overdispersion), the extremes are occupied too rarely, which gives a histogram with the shape of an inverted U. For the rank histograms, we combine the results from all grid points \( k \).

The ensemble integrations were carried out using reduced models with CMC, DTM, and AR(1) parameterization schemes, each starting from \( N_{\text{init}} = 1000 \) different initial states. For the calculations, we took ensemble sizes \( N_{\text{ens}} = 1, 5, \) and \( N_{\text{ens}} = 20 \). Results for RMSE and ANCR are shown in Figs. 9–11. The CMC scheme clearly performs better than the other two. The forecast lead time \( \tau \) at which the anomaly correlation drops below 0.6 is extended, with about 20% when changing from the DTM to the CMC schemes in case \( N_{\text{ens}} = 1 \), and with about 40% if \( N_{\text{ens}} = 20 \). From the DTM scheme with \( N_{\text{ens}} = 1 \) to the CMC scheme with \( N_{\text{ens}} = 20 \), this extension is about 65%. The CMC scheme with \( N_{\text{ens}} = 5 \) performs better than the DTM or AR(1) scheme with \( N_{\text{ens}} = 20 \).

Figure 12 shows the rank histograms for \( N_{\text{ens}} = 20 \). The CMC scheme has a positive effect on the ensemble spread; the corresponding rank histogram is nearly flat. The AR(1) scheme gives underdispersed ensembles; the DTM scheme leads to strong underdispersion. The rank histograms of the ensembles with AR(1) and DTM schemes can be made flatter by increasing the amplitude of the initial-state perturbations \( \xi^{\tau,0} \); the ensembles with CMC scheme become overdispersed if the perturbation amplitude is increased, as expected.

It is interesting to note that the AR(1) scheme does not perform better than the DTM scheme in our tests. As mentioned, the properties of the AR(1) stochastic process for the AR(1) scheme were estimated using the data at the shortest available time interval, \( \Delta t = 0.01 \). If we carry out the estimation at longer time intervals, resulting in AR(1) processes with shorter \( e \)-folding times, the performance of the AR(1) parameterization scheme does not improve (results not shown). In Wilks (2005), results are reported where the AR(1) scheme performs better than the DTM scheme in ensemble tests. Two important differences with the tests in our study are (i) the parameter settings of the full L96

\[
\text{ANCR}(\tau) = \frac{1}{N_{\text{init}}} \sum_s \frac{\mathbf{a}_{s,\text{mean}} \cdot \mathbf{a}_{s,\text{mean}}}{\sqrt{|\mathbf{a}_{s,\text{full}}|^2 |\mathbf{a}_{s,\text{mean}}|^2}},
\]

where \( \mathbf{a} \cdot \mathbf{b} = \sum \lambda_i a_i b_i \) and \( |\mathbf{a}|^2 = \mathbf{a} \cdot \mathbf{a} \). The anomaly correlation shows the average correlation after time \( \tau \) between the “true” trajectories (those of the full L96 model) and the mean “forecast” trajectories (the mean ensemble trajectories of the reduced model).

The rank histograms are calculated from ensemble

![Graph showing wave variances and mean wave amplitudes](image)
model used in Wilks (2005) (equivalent to ε = 0.1, K = 8, J = 32, h_x = −3.2, h_y = 1, and F = 18 or 20) and (ii) the setup of the ensemble tests (different ways to generate ensemble members). Rerunning our own tests using the parameter settings from Wilks (2005) gave quite similar results for the DTM and AR(1) schemes, in both the nonensemble tests (PDFs, correlation functions, wave statistics) and the ensemble tests. The difference of our findings with those reported by Wilks (2005) can be due to differences in the ensemble test setup (but note the consistency of our own ensemble and nonensemble test results) or to small yet apparently important differences in the way the schemes are implemented.

c. The case of Δt > δt

As already discussed in section 4b, practical circumstances may be such that the integration time step δt of the numerical model is different from the sampling time step Δt of the available data. Two ways of dealing with this problem were described in section 4b: (i) using a split-integration scheme (update X every δt time units, but B only every Δt time units) and (ii) calculating transition probability matrices \( \hat{P}^{(ij)} \) with time step δt, starting from matrices \( P^{(ij)} \) with time step Δt (the “generator method”). The latter possibility is more complicated to implement and gives less accurate results (for reasons also discussed in section 4b) than the split-integration
scheme in our testing setup. Therefore, we only present results here using the split-integration scheme. However, we note that the generator method can still be of practical use if model output is desired at a shorter sampling interval than that of the available data. Moreover, we speculate that the performance using the generator method will improve for models that are more strongly mixing than the L96 model.

The results obtained by using the split-integration scheme are shown in Figs. 13–16 (RMSE is omitted here, because it gives a similar picture as the anomaly correlation). For the construction of the stochastic matrices $P^{(a)}$, we used data from the full L96 model with sampling interval $\Delta t = 0.1$. The integration time step of the reduced model with CMC parameterization is $\delta t = 0.01$. Thus, during the integration, the $B_k$ are updated once every 10 time steps. As can be seen, this version of the CMC scheme for the case $\Delta t \gg \delta t$ performs well, although it remains slightly less accurate than the CMC method using $\Delta t = \delta t$ presented in sections 5a,b (in the figures, those results are added for comparison). The fact that the $B_k$ are updated only once every 10 time steps speeds up the computation significantly. This computational advantage can be such that using the split scheme may be attractive even if data with $\Delta t = \delta t$...
are available. The increase in speed can outweigh the decrease in performance (which is rather small in our results) in some circumstances, in particular, if $B_k$ does not evolve fast but on a similar time scale as $X_k$.

6. Discussion

The purpose of this study was to present a new approach to stochastic parameterization and to test this approach by implementing it in the framework of the Lorenz ’96 model. The new parameterization scheme we have proposed represents unresolved processes as stochastic (Markov) processes whose properties are conditional on the state of the resolved variables. In our numerical algorithm, these conditional Markov processes take the form of a collection of Markov chains, making practical implementation easy. The Markov chains are inferred from data in a very simple way, using only binning and counting.

We have compared the conditional Markov chain (CMC) scheme with two other parameterization schemes: a standard deterministic one (DTM) and the stochastic schemes [AR(1)] proposed in Wilks [2005]. Several tests were carried out to assess the performance of the various schemes, comparing probability distributions (PDFs), correlation functions (ACFs and CCFs), wave statistics, and ensemble forecasts. The CMC scheme performed better than the DTM and AR(1).
schemes, even though the number of states of the Markov chains was small ($N_g = 4$). To test the robustness of these results, we made a brief exploration of other parameter settings for the L96 model. With $\epsilon = 0.5$, the CMC scheme performed substantially better than the other two schemes. In the presence of clear time scale separation between the resolved and the unresolved variables ($\epsilon = 0.1$), differences in performance between the schemes were rather minor (results not shown).

For a better understanding of the good performance of the CMC scheme, it should be pointed out that with the parameter settings for L96 used in this study, the motion of phase points in the ($X_k$, $B_k$) plane tends to follow a roughly clockwise path. When $X_k$ increases, $B_k$ typically takes a (noisy) path through the upper part of the cloud of points shown in Fig. 1. During a decrease of $X_k$, $B_k$ is more likely to be in the lower part of the cloud. The imprint of this “loop” (somewhat reminiscent of a hysteresis loop) in otherwise noisy behavior can be captured with the CMC scheme but not with the DTM or AR(1) scheme. With the AR(1) scheme driven by red noise, sustained trajectories through the lower or upper part of the data cloud are also possible; however, those trajectories are equally likely to follow the loop as they are to go against it. The CMC scheme is better equipped, by design, to capture such structures.

Altogether, the results from the proposed CMC parameterization scheme are encouraging. Clearly, the L96 model is an idealized toy model, making it a suitable testing ground for a new parameterization approach. However, care should be taken when extrapolating results obtained within the L96 framework to other model situations. As a next step, the CMC scheme will have to be implemented and tested in a more realistic modeling environment.

In our testing, we already took into account two issues that can be expected to be of importance in more comprehensive model setups. First, we used parameter settings for the full L96 model that give little-to-no time scale separation between the resolved and unresolved variables. Previous studies of the L96 system were often carried out with parameters such that the unresolved variables (the $Y_{j,k}$) evolve on a faster time scale than the resolved variables. This time scale separation allows for the use of designated mathematical results and techniques (see, e.g., Fatkullin and Vanden-Eijnden 2004; Vanden-Eijnden 2003) but can be unrealistic.

A second issue is the sampling interval of the data that is available to base the parameterization on. The sampling interval can be different (longer) than the time step at which the numerical model with the parameterization scheme is integrated. This issue may arise when dealing, for example, with data stemming from observations that do not have a high temporal resolution. We discussed two potential solutions to this problem (see sections 4b and 5c). The split-integration scheme solution gave the best results here, with only a minor decrease in performance (and significant reduction in computation time) compared to the case in which both time steps were equal.

We conclude by discussing a number of further issues that will be of relevance for the application of the CMC scheme in more realistic models.

- The $X_k$ variables of the L96 model are usually interpreted as lying on a circle of constant latitude (see section 2). Thus, the L96 model can be regarded, somewhat loosely, as a model with one spatial dimension (the $x$ or E–W direction), discretized on a number of grid points. The $y$ and $z$ directions (N–S and vertical) are absent in L96. In more realistic models, all three spatial directions are present; gridpoint discretization leads to a number of grid points that is orders of magnitude higher than in the L96 model. From a computational point of view, this increase need not be problematic. The CMC scheme runs independently for each grid point and is therefore suitable for parallelization. The computational cost at each grid point is very small: only a small Markov chain must be evolved in time.

- The conditionality of the Markov chain at each grid point (the $X_k$ dependence) was limited by using a
locality” assumption: the Markov chain at grid point \( k \) depends only on the resolved variable \( (X_k) \) at that same grid point, not on variables at other grid points \( (X_{k'} \) with \( k' \neq k \). If the number of grid points increases because one considers models with two or three spatial dimensions, this locality assumption will keep the Markov chain conditionality equivalently simple. Thus, the complexity of the CMC scheme at each grid point is the same for a model with \( \alpha(10^4) \) grid points as it is for a model with \( \alpha(10^6) \) grid points.

- The main challenge for applying the CMC scheme to realistic models concerns the number of different variables at each grid point. In the L96 model, there is one quantity to be parameterized \( (B_k) \) and one resolved variable \( (X_k) \) at each grid point \( k \). However, the adiabatic core of an atmosphere model based on the primitive equations, for example, uses five resolved variables at each grid point. Making the Markov chain conditional on several resolved variables can lead to an intractable scheme if done naively. For example, dividing the range of each resolved variable into 16 intervals (similar to the 16 possible bins in which the vector of the five resolved variables at a single grid point can be at any moment. Keeping the number of these bins limited is necessary, both to limit computer memory demands of the CMC scheme and to keep estimation of the Markov chains from data feasible. Limiting the Markov chain conditionality to one or two judiciously chosen resolved variables and/or making the chain conditional on a (linear) combination of variables are possible solutions.

The number of quantities for which a parameterization is needed at each grid point is usually also larger than one. This can be dealt with by using separate Markov chains for separate quantities. Correlations between these quantities are partly accounted for through the dependence on resolved variables. If strong correlations or physical requirements (e.g., conservation properties) do not permit the use of separate Markov chains at a single grid point, one collective Markov chain must be constructed in which each Markov chain state corresponds to a particular combination of values for the different parameterized quantities at the grid point.

- The properties of the conditional Markov chain were inferred from data. For the L96 model, such data were easily generated by integrating the full L96 model. For more realistic models, there is no “meta-model” that can be run easily and cheaply to produce the necessary data. Instead, one can use data from two sources: 1) data observations (or reanalysis data) or 2) data produced by high-resolution models of limited spatial extent (e.g., a single GCM gridbox) and sufficient physical sophistication.

Using the same conditional Markov chain (i.e., the same set of transition matrices) at every grid point, as we did for the L96 model, will be too simple for some purposes (depending on the parameterized quantity). The geographical location of grid points can be expected to play a role in realistic models. It may be impractical to construct separate Markov chains for each grid point; instead, one can define a few groups of grid points (e.g., tropics or midlatitudes, boundary layer or above, grid points over land or sea) and construct Markov chains for each group separately.

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