Adding Multiple Time Lags to AE9/AP9 V1.0

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Paul O’Brien
Space Science Applications Laboratory
Physical Sciences Laboratories

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Space and Missile Systems Center
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[Signature]
James L. Roeder, Director
Space Sciences Department
Space Science Applications Laboratory
Physical Sciences Laboratories


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Abstract

The new AE9/AP9 models of the natural radiation belts at Earth capture belt dynamics via Monte Carlo scenarios. Because the radiation belts exhibit dynamics on timescales from hours to years, multiple time lags are needed in the dynamic Monte Carlo scenario generators. In the predecessor models (alpha and beta versions), time evolution carried information from only one prior state, that is, only one time lag. This document describes the calculations and challenges associated with adding multiple time lags to the autoregressive part of the V1.0 AE9/AP9 Monte Carlo scenario generator.
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1 Introduction

O'Brien and Guild [2010] (hereafter TEM2) provides a first-order autoregressive (AR-1) equation for the time evolution of the principal components ($\vec{q}$) of the spatial variation of the radiation belts:

$$\vec{q}_{t+\delta t} = G\vec{q}_t + C\vec{\eta}_t + \delta t,$$

(1)

In this document, we extend that equation to an arbitrary set of $N_G$ time lags, and we offset the time notation by $\delta t$ to make the meaning of the time lags more clear:

$$\vec{q}_t = \sum_{i=1}^{N_G} G_i \vec{q}_{t-\tau_i} + C_i \vec{\eta}_t,$$

(2)

This is an AR-N process [for discussion, see, e.g., Neumaier and Schneider, 2001]. The AR-1 is given by $N_G = 1$ and $\tau_1 = 0$. We assume that the time lags $\vec{\tau}$ will not necessarily be uniformly spaced. For example, $\vec{\tau}$ might be [6 hours, 1 day, 9 days, 27 days, 6 months, 1 year]. We note that $\tau_1 \equiv \delta t$ becomes the basic unit of time stepping, and that we will enforce that all other $\tau_i = T_i \delta t$, where $T_i$ is an integer, and $T_1 \equiv 1$. In our example $\vec{\tau}$, $T$ is given by [1, 4, 36, 108, 730, 1460]. We will also define an unlisted value $T_0 \equiv 0$.

The addition of arbitrary time lags presents three problems which will be addressed below: how to compute the $G_i$ and $C_i$ matrices, how to initialize $\vec{q}$ to begin a scenario, and how to store the history of $\vec{\eta}_t$ to support evaluation of (2). This document concludes with some estimates of the size of the matrices that must be computed and manipulated.
2 Computing AR-N Matrices

In TEM2, the AR-1 equations are derived from spatial ($\Sigma$) and one-day-lag ($R$) covariance matrices, transformed into corresponding matrices for the principal components via $Q$ and $Q^\dagger$:

$$QQ^T \approx \Sigma,$$

$$Q^\dagger Q = I,$$  \hspace{1cm} (3)  \hspace{1cm} (4)

$$G = \left[Q^\dagger R^T \left(Q^\dagger\right)^T\right]^{\delta t/T},$$  \hspace{1cm} (5)

$$C = (I - GG^T)^{1/2}.$$  \hspace{1cm} (6)

(Note: the dagger ($\dagger$) denotes a matrix pseudo-inverse, which is typically computed via singular value decomposition).

For the AR-N model, we will need a set of lag covariance matrices. Also, we will not be able to use 1-day time lag covariance matrix and adjust to $\delta t$, but rather we will have to use lag covariance matrices that exactly match certain time lags derived from $\vec{r}$. We define a generic Gaussian covariance matrix for the fluxes (i.e., a linear covariance matrix for the $\vec{z}$'s, which represent particle fluxes transformed to Gaussian variables) and its counterpart for the principal components ($\vec{q}$'s) as:

$$\hat{R}_M = \langle \vec{z}_t \vec{z}_t^T - \delta t\rangle = Q \langle \vec{q}_t \vec{q}_t^T - \delta t\rangle Q^T,$$  \hspace{1cm} (7)

$$R_M = \langle \vec{q}_t \vec{q}_t^T - \delta t\rangle = Q^\dagger \hat{R}_M \left(Q^\dagger\right)^T.$$  \hspace{1cm} (8)

We note that $\hat{R}_0 = \Sigma$ and $R_0 = I$ in TEM2, and that $R_{-M} = R_M^T$. The needed $\hat{R}_M$ matrices will have to be determined from data or from a reanalysis.

We create a series of matrix equations by taking the expected value of $\vec{q}_t \vec{q}_t^T - \delta t$:

$$\hat{R}_{T_j} = \sum_{i=1}^{N_G} G_i R_{T_j-T_i} + \begin{cases} C C^T & T_j = 0 \\ 0 & \text{otherwise} \end{cases}$$  \hspace{1cm} (9)

where the $CC^T$ term arises from $\langle \vec{\eta}_t \vec{q}_t^T \rangle$ because $\vec{\eta}_t$ is uncorrelated with all prior $\vec{q}_t$.

To put the equations in the usual form where we left-multiply the unknowns, we will use:

$$\hat{R}_{T_j} = \sum_{i=1}^{N_G} R_{T_i-T_j} C_i^T + \begin{cases} C C^T & T_j = 0 \\ 0 & \text{otherwise} \end{cases}$$  \hspace{1cm} (10)

We will solve for all the $G_i$ simultaneously, and then solve for $C$. 

2
The number of unique $\hat{R}_M$ matrices required is less than or equal to $1 + (N_G + 1)N_G/2$, of which all but one ($\hat{R}_0 = \Sigma$) are converted to covariances in the principal components ($\hat{R}_M$). Equation (11) is square and the matrix to be inverted is symmetric. After solving for the $G_i$, we will solve for $C$ using the $T_0$ case of (9):

$$CC^T = I - \sum_{i=1}^{N_G} \hat{R}_i G_i^T$$  \hspace{1cm} (12)

A few additional manipulations can show that this expression necessarily yields a symmetric $CC^T$. To obtain $C$, we take a Cholesky or eigenvalue square root of $CC^T$. 

$$C = \sqrt{CC^T}$$
3 Accuracy and Stability Criteria

We must ensure that $CC^T$ is symmetric and does not have any eigenvalues with magnitude larger than 1. Also, because (2) is only an approximation of the time series dynamics, it is possible that the solution $G_i$ is unstable. The stability condition is that none of the eigenvalues of the total time evolution matrix have magnitude 1 or greater.

The first (accuracy) condition is:
\[
\left| (CC^T)_{ij} - (CC^T)_{ji} \right| < \epsilon, \tag{13}
\]
where $\epsilon << 1$ is some small error tolerance, e.g., 0.01.

The second (stability) condition is:
\[
|\Lambda_{jj}| < 1, \tag{14}
\]
\[
CC^T = V\Lambda V^T, \tag{15}
\]
where $\Lambda$ is diagonal, and $VV^T = I$.

The final (stability) condition is:
\[
\left| \tilde{\Lambda}_{jj} \right| < 1, \tag{16}
\]
\[
\tilde{G} \tilde{G}^T = \tilde{V}\tilde{\Lambda}\tilde{V}^T, \tag{17}
\]
\[
\tilde{G} = \begin{pmatrix}
\tilde{G}_1 & 0 & \cdots & 0 \\
0 & \tilde{G}_2 & \cdots & 0 \\
\vdots & \vdots & \ddots & \vdots \\
0 & 0 & \cdots & \tilde{G}_{N_{G}}
\end{pmatrix}, \tag{18}
\]
\[
\tilde{G}_i = \begin{cases}
G_i, & t = T_i \\
0, & t \notin \{T_i\}
\end{cases}, \tag{19}
\]
where $\tilde{\Lambda}$ is diagonal, and $\tilde{V}\tilde{V}^T = I$. (Implementation note: $\tilde{G}$ is sparse, and can be handled with sparse matrix algorithms. In particular, Matlab’s “eigs” function can be used to compute only the largest magnitude eigenvalue of a sparse matrix).

In practice, empirically determined lag covariance matrices sometimes lead to time evolution solutions that do not meet the stability criteria. In order to ensure that the stability criteria are met, a tapering factor is used. A given lag correlation coefficient is multiplied by a value $\gamma^T_i$, where $\gamma \leq 1$ such that covariances at longer lags are suppressed relative to those at shorter lags. This effectively suppresses spurious empirical correlations. In the final version of the V1.0 runtime tables, the stability conditions were met for $\gamma = 1$, thus requiring no tapering.
4 Computing the lagged covariances on a reduced grid

Computing covariance matrices is one of the most computationally intensive parts of the back-end processing necessary to generate the AE9/AP9 runtime tables. One way to reduce the computing power needed is to reduce the size of the grid on which the covariance matrices are computed.

We already implement a version of this for the spatial covariance matrix $\Sigma$, from which we compute $Q$. In that process, we decimate the grid, compute a reduced $\tilde{\Sigma}$ and a reduced $\tilde{Q}$, and then interpolate the columns of $\tilde{Q}$ onto the full grid to obtain $Q$ (with some normalization corrections along the way).

Similarly, we can compute $\tilde{R}_{T_j}$ on a reduced grid: $\tilde{R}_{T_j}$, and, using only the corresponding rows of $Q$, i.e., $\tilde{Q}$, we can obtain $\tilde{R}_{T_j}$:

$$R_{T_j} = \tilde{Q}^\dagger \tilde{R}_{T_j} \left( \tilde{Q}^\dagger \right)^T .$$  \hspace{1cm} (20)

As it turns out, we only need slightly more rows in $\tilde{R}$ than $N_q$. For example, for the AE9 beta runtime tables (AE9VB/rev d), with $N_q = 9$, 9 or more randomly selected rows of $Q$ typically lead to a well-defined $\tilde{Q}^\dagger$. Therefore, the lag covariances can be computed on a dramatically decimated grid, and, to some extent, the points that make up that decimated grid can be selected based on sample size. Also, there is no need to interpolate, as we only need the covariance matrices for the principal components for $T_j \neq 0$. 

5 Initializing a Scenario

The “brute force” approach to initializing a scenario is to seed a series of $\vec{q}_{-\delta t}$ to $\vec{q}_{-T_{NG}\delta t}$ with Gaussian white noise and then step forward (2) many ($>> T_{NG}$) times. As a rough guide, the initialization will have to evolve for several times the longest lived eigenmode of $\tilde{G}$. This approach could conceivably leave out long-lag perturbations. Initializing for $N/\left(1 - \max \left|\Lambda_{jj}\right|\right)$ time steps should shrink long-lag perturbations to have magnitude $e^{-N}$; so, $N \sim 5$, would suppress such effects to be smaller than 0.01. Initial estimates suggest that for typical values of $N_q$ and $T_{NG}$, this initialization approach would take no more than 1-2 seconds. We can verify the validity of this scheme by initializing multiple instances of the $\vec{q}_t$ history to distinct samples of Gaussian white noise, but then evolving each instance with the same series $\vec{\eta}_t$. By determining when the different instances converge, we can determine how long to iterate (2) to initialize the data.

A more rigorous approach would be to initialize the meta-state vector $\hat{\vec{q}}^T = [\vec{q}_{-\delta t}^T, \vec{q}_{-2\delta t}^T, \ldots, \vec{q}_{T_{NG}\delta t}^T]$ with a properly-conditioned multivariate Gaussian so that $\hat{\vec{q}}$ preserves the covariance $\langle \hat{\vec{q}}\hat{\vec{q}}^T \rangle$. This would involve building and factoring a square matrix $N_q \times T_{NG}$ on a side. Such a matrix would include $R_M$ for all $M$ from 0 to $T_{NG}$, which is far beyond what is needed to compute $G_i$ and $C_i$ because $T_{NG} >> 1 + (N_G + 1)N_G/2$. It may be possible to solve this problem by only building and factoring the sub-matrices that make up the large covariance matrix, but so far my attempts to divine such an approach have failed.
6 Maintaining History of $\vec{q}$

To evaluate (2), we need to store a history of $\vec{q}_t$ for at least $T_{NG} + 1$ times. The natural memory structure for this is a loop queue: data does not shift each time step, rather, a pointer that defines $\vec{q}_t$ moves through the loop. After a time value falls off the end of the list, its memory address is reused for the next $\vec{q}_t$. 
7 Estimated Sizes of Matrices

Typical values are $N_q = 10$, $\delta t = 6$ hours, $N_G = 6$, $\tau_{NG} = 1$ year, $T_{NG} = 1460$. Therefore, a typical $R_{T_i}$, $G$, or $C$ matrix is $10 \times 10$, $22$ $R_{T_i}$ matrices are needed, and there are 600 unknowns in (11). The large matrices are all sparse, and can be solved efficiently by solvers that account for sparseness.

The loop queue for $\vec{q}_t$ must hold 14,600 entries, or $\sim 115$ kilobytes at double precision (64-bit). The large matrix in (11) is only $60 \times 60$.

The slowest part of building the tables for the AR-N model is computing the $22$ $R_{T_i}$. Recent code changes to the back-end processing have enable us to limit the size of $R_{T_i}$ computed from the data. Currently, $R_{T_i}$ is computed on a limited grid that has at most about 14,000 grid points. To keep development time under control, the limit could be further reduced without significant loss of precision (14,000 points is still far larger than the $N_q \sim 10$ principal components that are retained).
References
