Algorithms for Parallelizing AE9/AP9

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Algorithms for Parallelizing AE9/AP9

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Abstract

This report describes algorithms for parallelization of AE9/AP9. Especially for Monte Carlo scenarios, in which the entire mission must be simulated many times, parallelization should afford significant improvements in speed. The simulation can be broken up in the time domain and in the scenario domain. Breaking up the calculation in the time domain requires some care since the “worst-case running average” aggregators (e.g., worst-case one-day average flux) require stitching between time chunks. Parallelizing in the scenario domain requires only that the master processes combine results from the various threads and compute summary statistics (e.g., percentiles). There are additional minor parallelization opportunities as well. This report describes the concept of the various types of parallelization and provides specific algorithms and equations for doing the time-domain stitching of the aggregators.
Acknowledgments

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1. Introduction

The AE9/AP9 computer code is far more computationally intensive than any of its predecessors [Ginet et al., 2013]. First, AE9/AP9 represents the environment in terms of directional fluxes, which, for most applications, it must integrate numerically to compute an omnidirectional flux. In comparison, AE8/AP8 only had to interpolate omnidirectional fluxes to the spacecraft location. This adds a factor of ~10 to the number of fluxes that must be calculated in AE9/AP9. Second, AE9/AP9 uses a more sophisticated drift shell magnetic coordinate system to represent the fluxes, and those coordinates take longer to compute than the field-line coordinates used by AE8/AP8. Third, in order to provide confidence intervals, AE9/AP9 provides a set of scenarios, with each scenario having a perturbed statistical map from which the fluxes are drawn, and there being 40 or more scenarios. Finally, in order to represent dynamics for worst-case calculations, it is necessary to simulate the entire mission under consideration, rather than just a few representative orbits [O’Brien, 2007]. Together, all of these features of AE9/AP9 suggest the need for parallelization.

Individual AE9/AP9 scenarios are independent, thus the scenario is a natural dimension along which to parallelize. Also, AE9/AP9 was designed to permit parallelization in the time domain through the use of an “epoch time” input that initializes the random number generator for a specified start date and then “winds-forward” the state variables to the start of the time chunk to ensure that all chunks computed separately are identical to what they would have been had they been computed serially. Additionally, the random number seed is specific to each scenario, so that a scenario will be the same whether it is run in isolation or with many other scenarios. Finally, a single “run” of AE9/AP9 may be asked to compute a number of useful quantities, which we call aggregators. An example aggregator is the total dose behind a range of shielding depths, or the worst-case 24-hour average electron flux. Since the aggregators are independent of each other, they afford another possibility for parallelization.
2. Algorithm overview

Figure 1 show an overview of a nearly maximal parallelization strategy, i.e., one that minimizes redundant calculations while assuming a low cost for inter-thread communication (e.g., on a shared-memory system). We assume a master thread that launches several slave threads in a sequence of fork and join operations, until the final results are written to the file system.

2.1 Steps 1 and 2—Setup
In step 1 in Figure 1, the master thread ingests user input. The user specifies which model to run, what kind of run to do, the ephemeris for the run, and what aggregators to use. The master thread then breaks up the ephemeris into time chunks that are passed to slave threads for the computation of the projection weights. Projection weights require magnetic coordinates for each time point; then, those coordinates are used to compute the matrices that project the global model flux map onto the spacecraft trajectory. The calculation of coordinates is independent for each time point, so it is natural to split up the calculation along the time domain. The coordinates are re-used for every scenario, so there is no need (yet) to split along the scenario dimension. Step 2 has each slave thread computing coordinates and weights for one time chunk, then passing those results back to the master thread. The magnetic coordinate calculation takes longer depending on the location of the ephemeris point. Therefore, step 2 could benefit substantially from load balancing (although the value of load balancing diminishes if the time chunk is longer than a few orbits).

2.2 Steps 3 and 4—Computing Flux
In step 3 in Figure 1, the master thread again breaks the problem up in the time domain (not necessarily in the same time chunks used by steps 1 and 2) in preparation for the flux calculation. Because at this point in the processing the aggregators require the raw time series of flux at the spacecraft, the master thread must also prepare the aggregators to operate on each time chunk. Finally, the master thread can set up the calculation for each scenario (with its own instances of the aggregators) to run independently. In step 4, each slave thread computes the flux and runs the aggregator for one time chunk and one scenario. The aggregator partial results and ghost data (but not necessarily the full flux time series) are reported back to the master thread.

Because the magnetic coordinate data is very large, it is much more efficient on non-shared-memory machines to send the run parameters for steps 3 and 4 along with the dispatch from step 1 to 2 so that there is no need to return information to the master thread between steps 2 and 4. In this approach, step 3 would occur on each of the slave threads from step 2, and each slave thread would fork multiple subthreads (up to one per scenario) for step 4.

Is it even useful to split in time and scenario? The extra overhead associated with splitting along two dimensions simultaneously may not be worth the trouble if one can split arbitrarily finely in either of the dimensions. In fact, once one has committed to splitting in the time dimension, there is not necessarily anything to be gained by also splitting along the scenario dimension in step 4. Conversely,
Master Thread:
• Ingest user input
• Split ephemeris on time
• Dispatch time chunks

Slave Threads:
(Split on time domain)
• Compute coordinates and weights for each time chunk

Master Thread:
• Merge weights
• Split on time
• Dispatch scenarios for each time chunk

Slave Threads:
(Split by time and scenario)
• Compute flux for each time, scenario
• Run aggs on flux

Master Thread:
• Collect agg results along time dimension
• Dispatch agg merge for each agg, scenario

Slave Threads:
(Split by scenario and aggregator)
• Merge along time dimension for each agg, scenario

Master Thread:
• Collect agg results among scenarios
• Dispatch stats calculator for each agg and report time

Slave Threads:
(Split by aggregator, report time)
• Compute statistics across scenarios
• Write output

Master Thread:
• Collect agg results by report time
• Write stats reports for each agg

Figure 1. Overview strategy for maximal parallelization
splitting only in the scenario dimension is less useful because one cannot split to more threads than there are scenarios. On a shared-memory system with only 4–16 processors, and a typical set of 40 or more scenarios to run, this is not an issue. However, on a cluster, with potentially hundreds of threads and little shared memory, this distinction is very important. We will see that splitting in the time domain does have its costs, but avoiding them to split only in the scenario domain would lock us into a parallelization scheme that cannot scale up to use the full available computational resources.

2.3 Steps 5 and 6—Aggregation
In step 5, the master thread collects the aggregator results along the time dimension in preparation for merging them together. Then, the master dispatches one aggregator and one scenario to each slave thread for merging the aggregators’ results along the time domain. Section 3 describes the aggregator partial results, ghost data, and merging algorithms. Step 6 is broken up by scenario and aggregator. Again, a 2-D split is depicted, but this time neither dimension of the 2-D split can be divided into arbitrarily small pieces: the user defines the number of scenarios and aggregators, so splitting in two dimensions increases the utilization of available processors. It is noteworthy that different aggregators involve different levels of complexity in their merge step (step 6). So steps 5 and 6 are an important place for load balancing. At the completion of step 6, the slave threads return final results for each aggregator for each scenario.

2.4 Steps 7, 8, and 9—Summary Statistics
In step 7, the master thread collects the aggregator results and groups all the scenario results for a particular aggregator. It then dispatches each aggregator to a slave thread for summary statistics (such as percentiles), which are computed by analyzing the distribution of results across scenarios. Because the user may have requested several intermediate reporting times as well as the final report from each aggregator, it is possible to split step 8 by both the aggregator and the report time. In step 8, each slave thread computes the summary statistics for each report time and returns them to the master thread.

Finally, in step 9, the master thread collects the reports by aggregator, and produces one time-ordered file of summary statistics for each aggregator.

2.5 Prioritizing Parallelization
We have a mixed experience with regards to whether it takes longer to do step 4 or step 5. For a fluence/dose type run, step 4 is clearly the longest step because the aggregators are trivially simple. However, when worst-case aggregators are included, step 6 can actually exceed step 4. When a very long time interval is to be simulated, step 2 can become significant. When frequent (e.g., daily) reporting is requested, step 8 can become significant. However, for most of the cases we have run, it is steps 4 and 6 that dominate the calculation.
3. Algorithms for Parallelizing Aggregators

AE9/AP9 supports several different aggregators. They are listed in Table 1. All the aggregators are based on the time series \( j(E,t) \), which is the flux versus energy at a specific time. The dose and doserate aggregators rely on ShielDose2 [Selzter, 1994], \( S(J(E,t);d) \), and so work only with omnidirectional differential (per unit energy) flux. The other aggregators can work with any kind of flux.

We will assume that the time domain is divided into \( N_c \) time chunks, each of which spans a duration \( T_c \), and covers times from \( t_{c-1} \) to \( t_c \), or discrete time steps \( t_{1+k_{c-1}} \) to \( t_{k_c} \). Time tags are elapsed time, so continuous time starts at \( t=0 \) and the first discrete time tag is \( t_0=0 \). The subscript \( c \) will be used to represent information specific to time chunk \( c \). Reports are generated at times designated \( t_r \) or discrete time steps \( t_{k_r} \). Both \( c \) and \( r \) start at 1. In some of the equations below, we will see references to \( k_{c-1} \); for the edge case when \( c=1 \), we take \( k_{c-1}=k_0=0 \); thus, a reference to \( t_{1+k_{c-1}} \) for \( c=1 \) is actually a reference to \( t_1 \).

We will work through each of the aggregators, in turn, explaining how it can be parallelized over time chunks. Because the flux time series itself can be enormous (e.g., 20 floating-point numbers every 10 seconds of simulated time, for many scenarios and many simulated years), we have striven to parallelize the aggregators in a way that requires minimal passing of raw flux time series data between the slaves and the master thread. Thus, the aggregator may return partial results for the time chunk as well as ghost data that is needed to merge the partial results from multiple time chunks, but it is assumed the flux time series itself will never be passed between threads, except as contained in aggregator ghost data.

<table>
<thead>
<tr>
<th>Name</th>
<th>Equation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fluence</td>
<td>( J(E,t) = \int_{0}^{t} j(E,t')dt' )</td>
<td>Time integral of flux along orbit versus energy since start of mission</td>
</tr>
<tr>
<td>Mean</td>
<td>( \bar{j}(E,t) = \frac{J(E,t)}{t} )</td>
<td>Time average of flux along orbit versus energy since start of mission</td>
</tr>
<tr>
<td>Dose</td>
<td>( D(d,t) = \int_{0}^{\infty} G(d,E)J(E,t)dE = S(J(E,t);d) )</td>
<td>Total dose since start of mission</td>
</tr>
<tr>
<td>Boxcar</td>
<td>( \bar{j}<em>{\text{max}}(E,t;\tau) = \max</em>{\frac{t}{1} \leq t' \leq t} \bar{j}(E,t';\tau) ) ( \bar{j}(E,t;\tau) = \frac{1}{\tau} \int_{t-\tau}^{t} j(E,t')dt' )</td>
<td>Worst case box-car average flux (time window ( \tau ) since start of mission</td>
</tr>
<tr>
<td>Name</td>
<td>Equation</td>
<td>Description</td>
</tr>
<tr>
<td>------------</td>
<td>-------------------------------------------------------------------------------------------</td>
<td>-------------------------------------------------------------------------------------------------------------------------------------------</td>
</tr>
</tbody>
</table>
| Expavg     | \[
\hat{j}_{\text{max}}(E, t; \tau) = \max_t \left[ \frac{1}{\tau} \int_0^t \exp \left( \frac{t - t'}{\tau} \right) \hat{j}(E, t') dt' \right] \]
|            | Worst case exponential average flux (time window \(\tau\)) since start of mission (simulates RC circuit for internal charging analysis) |                                                                                                                                            |
| Doserate   | \[
\hat{D}(d, t; \tau) = \int_0^\infty G(d, E)\hat{j}(E, t; \tau) dE = S(\hat{j}(E, t; \tau); d) \]
|            | Dose rate averaged over preceding time interval \((\tau)\)                              |                                                                                                                                            |

### 3.1 The Fluence Aggregator

We begin with the *fluence* aggregator, and recast it as a discrete sum:

\[
J(E_i, t_k) = \sum_{k'=1}^{k} j(E_i, t_{k'}) \Delta t_{k'}
\]  
(1)

\[
\Delta t_k = \begin{cases} 
  t_2 - t_1 & k = 1 \\
  t_{N_k} - t_{N_k-1} & k = N_k \\
  \frac{t_{k+1} - t_{k-1}}{2} & \text{otherwise} 
\end{cases}
\]  
(2)

We have defined \(\Delta t_k\) under the assumption that each time point represents a time bin, rather than an entry in a time grid. Thus, to compute fluence for a day, the first time point would be at midnight, and the last time point would be just before midnight on the next day.

The fluence calculation can be parallelized by means of the partial fluence \(J_c(E_i, t_k)\) accumulated within time chunk \(c\), and the final chunk partial fluence \(J_c(E_i, t_{k_c})\):

\[
J_c(E_i, t_k) = \sum_{k'=1+k_{c-1}}^{k_c} j(E_i, t_k) \Delta t_{k'}
\]  
(3)

We note that to compute \(\Delta t_k\) the aggregator must know the time tags immediately before and after the time chunk. That is, it would need to know \(t_{k_c-1}\) and \(t_{1+k_c}\). The aggregator would also need to know \(N_k\).

Ghost data and partial results for the *fluence* aggregator are thus:

- **Input**
  - \(t_k\) for \(k\) in \([k_{c-1}, 1+k_c]\)
  - \(k_c, N_k\)

- **Output**
  - \(J_c(E_i, t_{k_c})\) for \(k_c\) in \([1+k_{c-1}, k_c]\)
  - \(J_c(E_i, t_{k_c})\)
Stitching together fluence for time chunks is simple:

\[ J(E_i, t_{kr}) = J_c(E_i, t_{kr}) + \sum_{c'=1}^{c-1} J_c(E_i, t_{k_{cr}}). \]  

(4)

That is, the reported fluence at \( t_{kr} \) is the partial fluence \( J_c(E_i, t_{kr}) \) for the time chunk containing \( k_r \) plus the sum of the final partial fluences of the previous time chunks.

### 3.2 The Mean Aggregator

The mean aggregator in discrete form is:

\[ j(E_i, t_{kr}) = \begin{cases} J(E_i, t_{kr})/t_2 & k_r = 1 \\ J(E_i, t_{kr})/t_{kr} & \text{otherwise} \end{cases} \]

(5)

Parallelizing the mean aggregator builds on the fluence aggregator, using the same ghost data and partial results, with Eq. (5) applied after Eq. (4).

### 3.3 The Dose Aggregator

The dose aggregator in discrete form is:

\[ D(d, t_{kr}) = S(j(E_i, t_{kr}); d). \]

(6)

As with the mean aggregator, parallelizing the dose aggregator builds on the fluence aggregator, using the same ghost data and partial results, with Eq. (6) applied after Eq. (4).

### 3.4 The Boxcar Aggregator

The boxcar aggregator is the maximum-to-date of a windowed running average, with window length \( \tau \). The discrete representation of the boxcar aggregator is:

\[ j_{\max}(E_i, t_{k}; \tau) = \max_{\tau \leq t_{kr} \leq t_k} j(E_i, t_{kr}; \tau) \]

(7)

\[ j(E_i, t_{k}; \tau) = \frac{1}{\tau} \left( J(E_i, t_k) - J(E_i, t_{k-\tau}) \right). \]

(8)

We note that the boxcar worst case is not defined for \( t<\tau \), because averages over intervals shorter than \( \tau \) can easily exceed the meaningful worst case for averages of length \( \tau \). The boxcar and doserate aggregators both rely on a box-car running average, \( j(E, t; \tau) \) with time window \( \tau \). That running average is, in turn, computed from the fluence at time \( t_k \) minus the fluence at time \( t_{k-\tau} \) in the past. It is best to compute the past fluence \( J(E_i, t_{k-\tau}) \) by interpolation in time.

Ghost data and partial results for the boxcar aggregator are thus:

- Input
\[ t_k \text{ for } k \in [k_{c,i}, 1+k_c] \]

\[ k_c, N_k \]

- Output

\[ \bar{j}_{\text{max},c}(E_i, t_{k_r}; \tau) \text{ for } t_{k_r} \text{ in } [t_{1+k_{c-1}} + \tau, t_{k_c}] \]

\[ \bar{j}_{\text{max},c}(E_i, t_{k_c}; \tau) \]

\[ J_c(E_i, t_k) \text{ for } t_k \in [t_{1+k_{c-1}}, t_{k_{c-1}} + \tau] \text{ or } [t_{k_c} - \tau, t_{k_c}] \]

The partial result \( \bar{j}_{\text{max},c} \) is given by:

\[ \bar{j}_{\text{max},c}(E_i, t_{k_r}; \tau) = \max_{t_{1+k_{c-1}} + \tau \leq t_{k'} \leq t_{k_r}} \bar{j}(E_i, t_{k'}; \tau). \] (9)

That is, it is the maximum starting \( \tau \) after the beginning of time chunk \( c \) rather than at \( t=0 \).

First, we stitch \( \bar{j} \) for the edge window cases, i.e., when \( t_{1+k_{c-1}} \leq t_{k_r} < t_{1+k_{c-1}} + \tau \) for any \( c>1 \) using Eq. (4) then Eq. (8). Equation (4) is not needed when \( T_c > \tau \), and one can simply replace \( \bar{j}(E_i, t_k) \) in Eq. (8) with \( J_c(E_i, t_k) \). We can then compute \( \bar{j}_{\text{max},c}(E_i, t_{k_r}; \tau) \) for all \( t_{k_r} \) in the edge window:

\[ \bar{j}_{\text{max},c}(E_i, t_{k_r}; \tau) = \max_{t_{1+k_{c-1}} - \tau \leq t_{k'} \leq t_{k_r}} \bar{j}(E_i, t_{k'}; \tau). \] (10)

Finally, we revise all the partial maxima \( \bar{j}_{\text{max},c}(E_i, t_{k_r}; \tau) \) into \( \bar{j}_{\text{max}}(E_i, t_{k_r}; \tau) \) by taking the maximum over the previous partial maxima and also the previous end-of-chunk maxima:

\[ \bar{j}_{\text{max}}(E_i, t_{k_r}; \tau) = \max_c \left[ \max_{c' < c} \bar{j}_{\text{max},c}(E_i, t_{k_{c'}}; \tau), \max_{r' \leq r} \bar{j}_{\text{max},c}(E_i, t_{k_{r'}}; \tau) \right]. \] (11)

### 3.5 The Doserate Aggregator

The *doserate* aggregator has a very simple special case when the reporting period exactly matches the averaging window (i.e., in continuous time \( t_r = t_{r-1} + \tau \)), the time step is constant, and the reporting time is an integer multiple of the time step. This special case can be addressed by simple post-processing of the *dose* aggregator:

\[ \dot{D}(d, t_{k_r}; \tau) = \begin{cases} \frac{D_d(d, t_{k_r})}{\tau} & r = 1 \\ \frac{D_d(d, t_{k_r}) - D_d(d, t_{k_{r-1}})}{\tau} & r > 1 \end{cases} \] (12)

In a serial implementation of the *doserate* aggregator, if the time window \( \tau \) exactly matches the reporting period, then there is no need for ghost data. The accumulator for \( \bar{j}(E_i, t; \tau) \) can simply be reset after each report is stored. However, in a parallel implementation, where time windows may involve consecutive time chunks, at least some ghost data is necessary anyway; thus, the parallel
implementation also enables the unlikely case of a time window that is not the same as the reporting window; it also addresses the likelihood that the time step is uneven, or that the reporting time is not an integer multiple of the time step.

For parallel implementation, the doserate aggregator in discrete form is:

$$\dot{D}(d, t_{kr}; \tau) = S(\bar{f}(E_i, t_{kr}; \tau); d)$$  \hspace{1cm} (13)$$

The parallel implementation of the doserate aggregator is similar to the boxcar aggregator, except no running maximum of \(\bar{f}(E_i, t_{kr}; \tau)\) is computed, and there is the added post-processing through \(S\).

Ghost data and partial results for the doserate aggregator are thus:

- **Input**
  - \(t_k\) for \(k\) in \([k_{e-1}, 1+k_c]\)
  - \(k_e, N_k\)

- **Output**
  - \(\dot{D}(d, t_{kr}; d)\) for \(t_{kr}\) in \([t_1+k_{e-1} + \tau, t_k]\)
  - \(J_c(E_i, t_k)\) \(t_k\) in \([t_1+k_{e-1}, t_{k_e-1} + \tau]\) or \([t_{k_e} - \tau, t_{k_e}]\)

As in the boxcar, we stitch \(\bar{f}\) for the edge window cases, i.e., when \(|t_{kr} - t_{k_e}| < \tau\) for any \(1 \leq e < N_e\) using Eq. (4) then Eq. (8) as for the boxcar. Then we compute \(\dot{D}(d, t_{kr}; \tau)\) for the edge cases using Eq. (13).

### 3.6 The Expavg Aggregator

The expavg aggregator is potentially the most challenging to break up in time. This is because the running average is an infinite impulse response moving average filter; at every time step, it responds to all prior fluxes.

The expavg aggregator in discrete form is:

$$\bar{f}_{\text{max}}(E_i, t_k; \tau) = \max_{t_{kr}} \bar{f}(E_i, t_{kr}; \tau)$$ \hspace{1cm} (14)$$

$$\bar{f}(E_i, t_k; \tau) = \left[1 - \exp\left(\frac{t_{k-1} - t_k}{\tau}\right)\right]j(E_i, t_k) + \exp\left(\frac{t_{k-1} - t_k}{\tau}\right)j(E_i, t_{k-1}; \tau)$$

$$= \sum_{k' = 1}^{k} j(E_i, t_{kr}) \left[1 - \exp\left(\frac{t_{k'-1} - t_{k'}}{\tau}\right)\right] \exp\left(\frac{t_{k'-1} - t_{k'}}{\tau}\right)$$

$$= \exp\left(\frac{t_{k-1} - t_k}{\tau}\right)j(E_i, t_{k-1}; \tau) + \sum_{k' = 1+k_{e-1}}^{k} j(E_i, t_{kr}) \left[1 - \exp\left(\frac{t_{k'-1} - t_{k'}}{\tau}\right)\right] \exp\left(\frac{t_{k'-1} - t_{k'}}{\tau}\right)$$  \hspace{1cm} (15)
The third form of Eq. (15) allows us to stitch together consecutive chunks. We could define a partial running average as

$$ j_c(E_i, t_k; \tau) = \sum_{k'=1+k_{c-1}}^k j(E_i, t_k') \left[ 1 - \exp \left( \frac{t_k' - t_k}{\tau} \right) \right] \exp \left( \frac{t_k - t_{k-1}}{\tau} \right) $$

(16)

We could have each slave thread return the entire partial moving average time series $j_c(E_i, t_k; \tau)$, and then compute the whole series of moving averages:

$$ j(E_i, t_k; \tau) = j_c(E_i, t_k; \tau) + \sum_{c' < c} j_c(E_i, t_{k_{c-1}}; \tau) \exp \left( \frac{t_{k_{c-1}} - t_k}{\tau} \right) $$

(17)

We could then apply Eq. (14) to compute $\bar{\max}(E_i, t_k; \tau)$.

However, we would like to avoid the need to pass the entire time series $j_c(E_i, t_k; \tau)$ between threads because it can amount to a lot of data. Instead, we will have the slave pass back partial results $\bar{j}_{\max,c}(E_i, t_{k_r}; \tau)$ at the reporting times and ghost data $j(E_i, t_k; \tau)$ for a window that is several (e.g., $N\tau = 10$) $\tau$ long. For raw fluxes that vary over 5 orders of magnitude, a ghost window of 10 $\tau$ ensures that the contribution of anything prior to the ghost window is negligible. Such a strategy degenerates to passing the entire series of $\bar{j}_{\max,c}(E_i, t_{k}; \tau)$ when $N\tau$ is longer than the time chunk $T_c$. A typical internal charging specification employs a $\tau$ of 1 day, but special cases may employ a $\tau$ of 6 months or longer.

Ghost data and partial results for the expavg aggregator are thus:

- **Input**
  - $t_k$ for $k$ in $[k_{c-1}, k_c]$
  - $k_c$, $N_k$

- **Output**
  - $\bar{j}_{\max,c}(E_i, t_{k_r}; \tau)$ for $t_{k_r}$ in $[t_{1+k_{c-1}} + N\tau, t_{k_c}]$
  - $j_c(E_i, t_{k_c}; \tau)$
  - $j_c(E_i, t_k; \tau)$ for $t_k$ in $[t_{1+k_{c-1}}, \min (t_{k_c}, t_{1+k_{c-1}} + N\tau)]$

The partial result $\bar{j}_{\max,c}(E_i, t_{k_r}; \tau)$ is given by:

$$ \bar{j}_{\max,c}(E_i, t_{k_r}; \tau) = \max_{t_{1+k_{c-1}} \leq t_{k'} \leq t_{k_c}} j(E_i, t_{k'}; \tau) $$

(18)

We can compute $j(E_i, t_k; \tau)$ for $t_k$ in the edge window $[t_{1+k_{c-1}}, \min (t_{k_c}, t_{1+k_{c-1}} + N\tau)]$ by adding in the last value from the prior time chunks according to Eq. (17). From this, we can compute $\bar{j}_{\max}(E_i, t_{k_r}; \tau)$ for all $t_{k_r}$ in the edge window. Then, for all $t_{k_r}$ outside the edge window, we simply add on the prior end-of-chunk values $j_c(E_i, t_{k_c}; \tau)$ with a proper exponential decay factor:
\[
\bar{j}_{\text{max}}(E_i, t_{k_r}; \tau) = \max [\bar{j}_{\text{max}}(E_i, t_{k_{r-1}}; \tau), \bar{j}'_{\text{max}}(E_i, t_{k_r}; \tau)]
\]

\[
\bar{j}'_{\text{max}}(E_i, t_{k_r}; \tau) \approx \begin{cases} 
\bar{j}_{\text{max}}(E_i, t_{k_r}; \tau) & 0 \leq t_{k_r} - t_{1+k_{c-1}} \leq N_{\tau} \tau \\
\bar{j}_{\text{max}, c}(E_i, t_{k_r}; \tau) + \sum_{c'=1}^{c-1} j_{c'}(E_i, t_{k_{c'-1}}; \tau) \exp \left( \frac{t_{k_{c'-1}} - t_k}{\tau} \right) & \text{otherwise}
\end{cases}
\]

We are thus assuming that by the time we get to \( t_{k_r} > t_{1+k_{c-1}} + N_{\tau} \tau \), all contributions to \( \bar{j}(E_i, t_{k}; \tau) \) from prior time chunks are negligible. In Eq. (19), we take the maximum-to-date to ensure that a maximum from a prior time chunk or a reconstructed maximum earlier in the same time chunk appropriately replaces the partial result maximum.

We have conducted several numerical experiments using the algorithms described here and have confirmed that for a log-normally varying time series that spans 5 orders of magnitude, this approach works to 10^{-13} maximum relative error (a couple orders of magnitude larger than the floating-point error because we are doing sums of many variables, but well below the needs of satellite design, for which relative error 10^{-2} is more than adequate). We have also confirmed that adding in serial correlation (lag correlation) to the underlying data does not cause the algorithm to fail.
4. Summary

We have described the overall approach to parallelizing large AE9/AP9 calculations. We have discussed different strategies for shared-memory and non-shared-memory systems (i.e., clusters). We have also defined the various aggregators built into AE9/AP9 and how to break those aggregators up over time.

In only one case, namely, the `expavg` aggregator, which approximates an RC circuit, do we potentially lose information due to not passing the full flux time series from slave thread back to master. In that case, the estimated relative error (~$10^{-13}$) is many orders of magnitude smaller than what is of consequence for satellite design.

Implementation of these algorithms will enable the large orbit surveys, long mission simulations, and numerous Monte Carlo scenarios that enable sophisticated decision making by modern satellite designers.
References

