Optimization of a Multilevel Checkpoint Model with Uncertain Execution Scales

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Abstract-Future extreme-scale systems are expected to experience different types of failures affecting applications with different failure scales, from transient uncorrectable memory errors in processes to massive system outages. In this paper, we propose a multilevel checkpoint model by taking into account uncertain execution scales (different numbers of processes/cores). The contribution is threefold: (1) we provide an in-depth analysis on why it is difficult to derive the optimal checkpoint intervals for different checkpoint levels and optimize the number of cores simultaneously; (2) we devise a novel method that can quickly obtain an optimized solution-the first successful attempt in multilevel checkpoint models with uncertain scales; and (3) we perform both largescale real experiments and extreme-scale numerical simulation to validate the effectiveness of our design. The experiments confirm that our optimized solution outperforms other stateof-the-art solutions by 4.3-88% on wall-clock length.

I. INTRODUCTION

Extreme-scale environments with a million cores or more will be used more and more commonly in solving scientific problems [1]. With such numbers of cores, however, such environments become more fragile, and fault tolerance mechanisms are needed in order to protect high-performance computing (HPC) executions.

To solve the fault tolerance issue, many researchers have proposed solutions [2] that adopt a periodic checkpoint/restart model (or checkpoint model for short) [3], [4], because of its ease of use and acceptable overall performance in practice [5], [6], [7]. For an MPI program, one just needs to periodically set checkpoints (i.e., store running processes' memories) via a *parallel file system (PFS)*, and roll back execution to the most recent checkpoint upon a failure. Such a simple PFS-based checkpoint model is called single level, because it uses only one storage technology for storing checkpoints.

For exascale applications, however, the classic singlelevel checkpoint model may result in a huge performance degradation. One reason is that exascale applications [8] are likely to experience different types of failures frequently, because of the number of cores used simultaneously. Another reason is that classic checkpoint/restart models always store checkpoint files on the PFS, whereas exascale applications tend to use very large datasets during the execution, leading to a huge checkpoint/restart overhead (up to 25%) [9] due to the I/O bottleneck. Compared with the single-level checkpoint/restart model, multilevel checkpoint/restart [10], [11], [12], [13] is a promising model to solve the cited problem. The Fault Tolerance Interface (FTI) [13] toolkit, for example, provides four-level checkpointing interfaces, including local-storagedevice, partner-copy [13], [14], Reed-Solomon encoding (RS-encoding) [15], [16], and PFS. Different checkpoint levels handle different types of failure events. For example, partner-copy technology makes each checkpoint file have two copies stored in local storage device and another partnernode, respectively. Thus, upon a failure event with multiple simultaneous hardware crashes,¹ the whole execution can still be recovered via partner-copy as long as no adjacent/partner nodes have crashed.

Since different levels correspond to various checkpoint overheads, such a multilevel checkpoint model creates a new avenue to refine the checkpoint strategies. For example, saving processes' runtime memories in the form of checkpoint files on local storage devices in parallel would be much faster than on the PFS. The difference of checkpoint overheads between local storage devices and the PFS will be larger in the near-future systems, especially because of the rapid development of non-volatile dynamic RAM (NVDRAM) (such as DVDIMM [19]), which is now available with DDR3 DRAM [20] and to be integrated with DDR4 [21].

In our previous work [22], we proposed an efficient method to compute the optimal checkpoint intervals for different levels and to optimize the selection of levels for each HPC application. However, that solution optimizes the checkpoint intervals for fixed execution scales. Our recent experiments indicate that an application's performance with the checkpoint model not only is determined by the checkpoint/recovery overheads and rollback loss but also is related to the execution scale with consideration of checkpoint/recovery overhead and failure events. Hence, we revisit the multilevel checkpoint model in order to optimize the execution scales based on multilevel checkpoint overheads and various failure events.

¹Simultaneous failures means multiple nodes fail in a short period (i.e., correlated failure window), such as a resource allocation period. In [17] and [18], the window lengths are set to 1 minute and 2 minutes, respectively. Simultaneous node failures may also occur because of malfunctioned switches or power boards shared by multiple nodes.

Our main contributions are threefold.

- We first reformulate and analyze the multilevel checkpoint model based on both checkpoint interval variables and the number of cores to be used in execution. Our problem formulation takes into account both linear and nonlinear speedup applications. The key challenge of the problem is twofold. First, the lower-level checkpoint overheads will impact the higher-level rollback loss, so we have to combine all checkpoint levels with different failure rates to optimize the entire performance for each HPC application. Second, the failure event probability likely depends on execution scales; that is, different numbers of cores in execution will definitely impact the failure rates on different checkpoint levels, which leads to a nonconvex optimization problem. Hence, finding the optimal solution based on such a new multilevel checkpoint model is extremely difficult. This contrasts with Young's formula [3] or Daly's work [4], which is much easier to use in order to get approximate checkpoint intervals because of the single checkpoint level and fixed execution scales.
- We propose a method that can quickly optimize the trade-off between speedup and various overheads, with consideration of uncertain execution scales. That is, the new solution can optimize the checkpoint intervals for different levels and optimize the number of cores simultaneously. On the one hand, the larger the number of processes/cores used, the shorter the productive time we obtain in general. On the other hand, different numbers of processes to be launched in the execution will lead to different failure rates and checkpoint overheads. Hence, there would be an optimum scale for a specific application. Based on the multilevel checkpoint model, we comprehensively analyze how to optimize the scales with regard to both the application speedup and various overheads. To the best of our knowledge, this is the first attempt to solve such a problem with this approach.
- We evaluate our multilevel checkpoint model and some related work by using an exascale simulation environment. This environment provides simulation results close to practical situations, since we carefully emulate the execution of real MPI programs and also validate the correctness of its generated results using a real cluster environment with 1000+ cores. Our solution adopts both the optimized checkpoint intervals and optimized execution scales computed by our algorithm. The related work we evaluate includes (1) a singlelevel PFS checkpoint model with Young's formula [3]; (2) a multilevel checkpoint model with optimized checkpoint intervals yet without optimization of scales (i.e., our previous work [22]); and (3) a single-level PFS checkpoint model with both Young's formula and optimized number of processes, which is proposed by

[23]. Experiments show that our approach outperforms other solutions by 4.3–88%.

The rest of the paper is organized as follows. In Section II, we formulate the multilevel periodic checkpoint model to minimize the entire wall-clock time for each HPC application with respect to execution scales and checkpoint intervals. In Section III, we derive the provably optimal solution for different types of HPC applications. We present our experimental results in Section IV, discuss the related works in Section V, and provide concluding remarks and a look at future work in Section VI.

II. PROBLEM FORMULATION

In this section, we formulate our research as an optimization problem based on the multilevel checkpoint/restart model. We focus mainly on the periodic checkpoint model (i.e., equidistant checkpoint model) because it is a de facto standard in the fault tolerance research community.

Here, we propose a generic multilevel checkpoint/restart model with L checkpoint levels. The checkpoint level 1 corresponds to transient/software failures. The remaining higher levels (2, 3, \cdots , L) correspond to different cases of hardware failures. In FTI, for example, there are four checkpoint levels (local storage, partner-copy, RS-encoding, and PFS), which correspond to software error with no hardware failure, nonadjacent node failures, a certain number of node failures with adjacent failure cases, and the situations that lower levels cannot take over, respectively. Upon any type of hardware failure, the system will reallocate a new set of nodes/cores to replace the crashed nodes/cores; and the resource allocation is a constant period, denoted by A, which is far shorter than the application execution time.

The checkpoint overhead and recovery overhead¹ are different from level to level and may also be different with various numbers of processes/cores used. Suppose we are given N processes running on N cores in parallel. Then the checkpoint overhead at the checkpoint level *i* is denoted by $C_i(N)$. In general, $C_1(N) \leq C_2(N) \leq \cdots \leq C_L(N)$. Similarly, the recovery overhead at level *i* is denoted by $R_i(N)$, where $R_1(N) \leq R_2(N) \leq \cdots \leq R_L(N)$ in general.

Considering the various checkpoint/recovery overheads with different numbers of cores, minimizing the wall-clock time for a specific HPC application is actually a tradeoff between its original speedup and increasing failure rates as the execution scales. In general, the speedup² of an HPC application increases more and more slowly with more cores, as shown in the grey curve in Figure 1. On the other hand, the failure rates may increase with the execution scales, shown as dash-dot lines in Figure 1. The figure illustrates

¹*Recovery overhead* means the time cost in restarting a failed application, so it is also known as restart overhead.

 $^{^{2}}Speedup = \frac{single-core length}{parallel execution time}$. For simplicity, we focus on the HPC applications whose speedups can be characterized as a continuous function instead of a piecewise function.

typical performance curves with and without regard to the checkpoint overheads and failure events. As shown by this figure, the optimal number of cores with checkpoint is lower than without checkpoint. The application's real productive time with N cores is denoted as $f(T_e, N)$, where T_e refers to the single-core productive time (or execution length) by excluding any failure-related costs such as checkpoint overhead and roll-back loss. We denote the speedup of the application running with N cores as g(N). Hence, $f(T_e, N) = \frac{T_e}{g(N)}$. For example, if the parallel execution follows a linear speedup, then $g(N) = \kappa N$ and $f(T_e, N) \approx \frac{T_e}{\kappa N}$, where κ is a constant.



Figure 1. Tradeoff between Execution Speedup and Checkpoint Overhead

The optimization of such a multilevel checkpoint model is nontrivial because of the following factors: (1) the entire wall-clock time of any application is synthetically determined by checkpoint/recovery overheads and roll-back losses at different levels; (2) the parallel execution may be struck by different types of failures, and the failure locations are randomly distributed; and (3) the failure probabilities are different from level to level and are related to the number of processes/cores. To make the problem tractable, we introduce a key random variable called the number of failures (denoted by Y) during the application's execution.

Our objective is to minimize the expected wall-clock length $E(T_w)$ for each given application. $E(T_w)$ can be written as Formula (1), where L denotes the total number of checkpoint levels, x_i refers to the number of checkpoint intervals at level i, Γ_{ij} refers to the roll-back loss due to the *j*th failure occurring at level *i* in the execution, and $P_i(Y=K)$ denotes the probability of experiencing K failure events at checkpoint level *i*. We need to determine the optimal values of x_1, x_2, \dots, x_L , and N, with minimized $E(T_w)$ regarding all possible overheads and roll-back losses.

$$E(T_w) = f(T_e, N) + \sum_{i=1}^{L} C_i(N)(x_i - 1) + \sum_{i=1}^{L} \left[\sum_{K=1}^{\infty} \left(P_i(Y = K) \sum_{j=1}^{K} (\Gamma_{ij} + A + R_i(N)) \right) \right]$$
(1)

Key notations are summarized in Table I.

Such a multilevel checkpoint model is generic enough to be suitable for different scenarios. For example, the key difference between the strong-scaling scenario and weakscaling scenario is different speedup functions (either with

Table I SUMMARY OF KEY NOTATIONS

Notation	Description
L	# of checkpoint levels each with various failure types
N	# of processes (or cores) of the focused application
T_e	single-core productive time of the application
$f(T_e, N)$	parallel execution time with N cores
g(N)	speedup function of the parallel application
$C_i(N)$	checkpoint overhead at level i
$R_i(N)$	restart overhead at level i
A	resource allocation period (a constant)
x_i	# of checkpoint intervals of the application at level i
$P_i(Y=K)$	probability of encountering K failures at level i
Γ_{ij}	roll-back loss in execution due to j th failure at level i

consideration of scale increase or not) and checkpoint overhead/recovery functions. Our model is suitable for both cases because of the generic definitions of the these functions in our model.

III. OPTIMIZATION OF MULTILEVEL CHECKPOINT MODEL WITH UNCERTAIN EXECUTION SCALES

In this section, we first theoretically analyze the huge challenges in solving the multilevel checkpoint model and then propose a method to optimize the solution.

A. Difficulty Analysis

One straightforward idea is to leverage convex optimization theory [24]. Note that the preceding problem has L+1 variables, x_1, x_2, \dots, x_L , N. One must prove that the target function $E(T_w)$ is always convex with respect to any of these variables (i.e., $\forall i=1,2,\dots,L, \frac{\partial^2 E(T_w)}{\partial x_i^2} > 0$ and $\frac{\partial^2 E(T_w)}{\partial x_i} > 0$). Then, the optimal solution can be computed based on $\frac{\partial E(T_w)}{\partial x_i} = 0$ and $\frac{\partial E(T_w)}{\partial N} = 0$. Unfortunately, $E(T_w)$ is not always convex with respect to its variables.

Indeed, in the following, we show that $E(T_w)$ is not convex even with respect to the **single-level** checkpoint model and linear-speedup application and, by generalization, to the **multilevel** checkpoint model with more complicated applications. That is, $g(\kappa)=\kappa N$ holds in the following analysis.

In the single-level checkpoint model with only the PFS used to store checkpoint files, the checkpoint overhead and recovery overhead can be represented by Formula (2) and Formula (3), respectively. In the two formulas, ϵ_0 , and η_0 refer to constant costs, α_0 and β_0 denotes two constant coefficients, and $H_c(N)$ and $H_r(N)$ denotes the increasing overhead rate with the execution scale. For instance, if the checkpoint/recovery overheads increase linearly, then $H_c(N)=H_r(N)=N$ holds; if they are constants, then $H_c(N)=H_r(N)=0$ holds.

$$C(N) = \epsilon_0 + \alpha_0 H_c(N) \tag{2}$$

$$R(N) = \eta_0 + \beta_0 H_r(N) \tag{3}$$

We will show that given $H_c(N)=H_r(N)=N$, the problem is already extremely difficult to solve directly. We rewrite the problem formulation as Formula (4), where x is the number of checkpoint intervals and Γ_j refers to the roll-back loss upon the *j*th failure during the execution.

$$E(T_w) = \frac{T_e}{\kappa N} + (\epsilon_0 + N\alpha_0) \cdot (x-1) + \sum_{K=1}^{\infty} \left(P(Y=K) \sum_{j=1}^{K} (\Gamma_j + A + \eta_0 + N\beta_0) \right)$$
(4)

Since the failure events occur randomly during the task execution, the expected Γ_j (i.e., average rollback loss for each failure) can be approximated as $\frac{T_e/(\kappa N)}{2x}$, where $\frac{T_e/(\kappa N)}{x}$ refers to the maximum rollback length¹ upon a failure event. By further leveraging the definition of mathematical expectation (i.e., $\sum_{K=1}^{\infty} [K \cdot P(Y=K)] = E(Y)$), we can convert Formula (4) to Formula (5).

$$E(T_w) = \frac{T_e}{\kappa N} + (\epsilon_0 + N\alpha_0) \cdot (x-1) + E(Y)(\frac{T_e/(\kappa N)}{2x} + \eta_0 + N\beta_0 + A)$$
(5)

In general, the expected number of failures, E(Y), is not a constant but is related to the whole wall-clock length, and the expected wall-clock length is the variable $E(T_w)$ in the formula. If we denote the expected failure rate of the parallel application by $\lambda(N)$, then E(Y) can be approximated as $\lambda(N)E(T_w)$, which leads to the following formula by eliminating E(Y) from Formula (5).

$$E(T_w) = \frac{\frac{T_e}{\kappa N} + (\epsilon_0 + N\alpha_0) \cdot (x-1)}{1 - \lambda(\frac{T_e}{2\pi \kappa N} + \eta_0 + N\beta_0 + A)}$$
(6)

As long as we can derive $\frac{\partial E^2(T_w)}{\partial x^2} \ge 0$ and $\frac{\partial E^2(T_w)}{\partial N^2} \ge 0$, we can compute the optimal solution based on $\frac{\partial E(T_w)}{\partial x} = 0$ and $\frac{\partial E(T_w)}{\partial N} = 0$. However, this is not a viable idea. On the one hand, it is hard to prove that the second-order derivatives $\frac{\partial E^2(T_w)}{\partial x^2}$ and $\frac{\partial E^2(T_w)}{\partial N^2}$ are always greater than 0 because of their extremely complicated representations. We find that they are actually lower than 0 in some situations. On the other hand, the first-order derivatives $\frac{\partial E(T_w)}{\partial x} = 0$ and $\frac{\partial E(T_w)}{\partial N} = 0$ also lead to complicated high-degree equations (such as quartic equations), which are hard to solve [25].

B. Overview of Optimized Algorithm

We explore an efficient algorithm to optimize the solution to the above problem. The basic idea is to introduce an extra condition that assumes that the expected number of failures (denoted by μ_i) at each specific level during the execution is related only to the execution scale (i.e., the number of processes/cores). That is, we can denote μ_i to be $\mu_i(N)$, regardless of the application length. With such a condition, the target $E(T_w)$ can be simplified as a convex optimization problem with respect to all its variables in general, so we can try to resolve a convex-optimization problem instead. The extra condition is removed later through a set of iterative steps: the algorithm alternatively derives optimal solution with the given numbers of failures and computes the new numbers of failures based on the changed expected wallclock length until convergence. The pseudo-code is shown in Algorithm 1.

Algorithm 1 OPTIMIZED ALGORITHM

Input: productive time T_e , estimated speedup function g(N), checkpoint overheads, recovery overheads

1: for (level $i=1 \rightarrow L$) do

- 2: Compute μ_i based on $f(T_e, N)$ /*Initialize expected # of failures*/ 3: end for
- 4: repeat
- 5: Compute optimal solution x_i^* $(i=1,2,\dots,L)$ and N^* , based on convex optimization and μ_i $(i=1,2,\dots,L)$.
- 6: Compute expected $E(T_w)$ based on x_i^* and N^* .
- 7: for (level $i=1 \rightarrow L$) do
- 8: $\mu'_i \leftarrow \mu_i$. /*Save old μ_i */
- 9: Recompute μ_i based on $E(T_w)$.
- 10: end for
- 11: **until** $(\max(\mu'_i \mu_i) \le \delta, \forall i = 1, 2, \cdots, L)$

In this algorithm, we first initialize the expected number of failure events for each checkpoint level i based on the estimated productive time $f(T_e, N) = \frac{T_e}{g(N)}$ (line 1-3). Then, the algorithm goes into a loop that iteratively computes the optimal number of checkpoint intervals x_i^* for each level as well as the optimal number of cores N^* . In each iteration step, after computing the optimal solution with convex-optimization theory (line 5), the algorithm adopts the solution to estimate the wall-clock length (line 6), based on which it can recompute more accurate expected numbers of failure events (lines 7–10). This iteration loop stops when the expected numbers of failure events converge within a specified error threshold δ . Obviously, the key step is line 5, which computes the optimal solution based on the expected numbers of failures on different levels, which are regardless of the wall-clock length.

In the following text, we focus on this key step. We start with the single-level checkpoint model and then present the complete optimal solution for the multilevel checkpoint model.

C. Optimization for the Single-Level Model

In this subsection, we investigate how to optimize checkpoint intervals and the execution scale simultaneously, for the linear-speedup and nonlinear speedup applications, respectively.

1) Optimization Based on Linear Speedup: The productive time of the linear-speedup application can be approximated as $f(T_w, N) \approx \frac{T_e}{\kappa N}$, where T_e denotes the singlecore execution length and κ is a constant. In this case, we set $\mu(N)=bN$ for simplicity. Similarly, for simplicity, we suppose the checkpoint/recovery overheads (C(N)) and R(N) are constants.

The representation of the target function $E(T_w)$ can be simplified as Formula (7).

$$E(T_w) = \frac{T_e}{\kappa N} + \epsilon_0(x-1) + bN(\frac{T_e/(\kappa N)}{2x} + \eta_0 + A)$$
(7)

¹For simplicity, we do not consider the situation in which a new failure event occurs during recovery period. In fact, based on our analysis in previous work [22], the failure-over-recovery situation occurs rarely because the recovery period is usually far shorter than productive time and failure interval.

Since $\frac{\partial E^2(T_w)}{\partial x^2} = \frac{bT_e}{\kappa x^3} > 0$ and $\frac{\partial E^2(T_w)}{\partial N^2} = \frac{2T_e}{\kappa N^3} > 0$, there must be an optimum point for the variables $\{x, N\}$ such that the expected wall-clock length $E(T_w)$ is minimized. We can compute the optimal values of x and N, as long as we solve both Formula (8) and Formula (9) simultaneously.

$$\frac{\partial E(T_w)}{\partial x} = \epsilon_0 + \frac{bT_e}{\kappa} \cdot \frac{-1}{2x^2} = 0$$
(8)

$$\frac{\partial E(T_w)}{\partial N} = \frac{-T_e}{\kappa N^2} + b(\eta_0 + A) = 0 \tag{9}$$

Then, we can derive the following two simple formulas to directly compute the optimal number of checkpoint intervals x^* and the optimal scale N^* .

$$x^* = \sqrt{\frac{bT_e}{2\kappa\epsilon_0}} \tag{10}$$

$$N^* = \sqrt{\frac{T_e}{\kappa b(\eta_0 + A)}} \tag{11}$$

2) Optimization Based on Nonlinear Speedup: In our model, we propose a generic method to optimize the performance over the checkpoint model, which can fit different types or functions of speedup curves. In this work we focus mainly on quadratic nonlinear curves. However, our model can also be easily extended to more complicated speedup functions if needed, because of the generic definition of the speedup function q(N). As an example, let us consider Figure 2(a). The blue points shown in the figure are speedup values computed based on real experiments with up to 1,024 cores on the Argonne Fusion cluster [26]. The speedup of the *Heat Distribution* application¹ increases like a linear curve for the relatively small execution scale range and follows a quadratic curve for the long range of execution scale. In fact, some application speedups may not follow quadratic curve in the whole execution range, as shown in Figure 2(b). The speedup of the Nek5000 eddy uv application² increases quickly in the initial range while decreasing after 100 cores because of the increasing communication cost. Since the optimal scale with regard to the checkpoint model must be no bigger than the original optimal scale without the checkpoint model, we need to focus only on the initial scale range through the point with the maximum original speedup. Then, we can still use the quadratic curve to fit the speedup very well, as shown in Figure 2(b). Specifically, the quadratic curve generated based on the initial scale range (1-100 cores) fits the increasing speedup records best from among all different fitting curves.

The speedup function is defined as Formula (12). We denote the symmetrical axis of the quadratic curve to be located at $N^{(*)}$. Then, since the speedup curve must pass



through the origin (0,0), we can derive the speedup curve as Formula (12), where N is the number of cores and κ is referred to as the curve's slope at (0,0).

$$g(N) = -\frac{\kappa}{2N^{(*)}}N^2 + \kappa N \tag{12}$$

In practice, κ can be estimated by a simple test with a small or middle execution scale. In the Heat Distribution program, for example, the speedup is 77 when using 160 cores, so κ can be approximated as $\frac{77}{160} \approx 0.48$, which is close to the real value κ =0.46. In fact, the coefficients of Formula (12) can also be estimated via a set of laws such as Amdahl's law [31], Gustafson-Barsis's law [32], and Karp-Flatt metric [33], which are used to predict, respectively, the upper limit of the speedup, how long the application would take to run on a single core, and the performance of the application on larger execution scales.

Similar to the derivation of Formula (7), the target function with respect to the nonlinear speedup applications can be written as Formula (13). The only difference is that the execution speedup g(N) in Formula (13) follows a nonlinear curve, as shown in Figure 2.

$$E(T_w) = \frac{T_e}{g(N)} + \epsilon_0(x-1) + bN(\frac{T_e/g(N)}{2x} + \eta_0 + A)$$
(13)

One can easily verify that the target function Formula (13) is always convex with respect to x and N, respectively. Hence, there must be a unique value point for (x,N) with the minimum value of $E(T_w)$. We can compute the optimal solution (x^*,N^*) by making $\frac{\partial E(T_w)}{\partial x}=0$ and $\frac{\partial E(T_w)}{\partial N}=0$. Accordingly, we derive the following equations:

$$\frac{\partial E(T_w)}{\partial x} = \epsilon_0 + \frac{bNT_e}{2g(N)} \left(\frac{-1}{x^2}\right) = 0 \tag{14}$$

$$\frac{\partial E(T_w)}{\partial N} = T_e \frac{b}{2x} \frac{1}{g(N)} - T_e \frac{(1 + \frac{bN}{2x})g'(N)}{g^2(N)} + b(\eta_0 + A) = 0 (15)$$

Directly solving these two equations simultaneously is difficult because the transformed equation after eliminating x using Equation (14) leads to an extremely complicated equation. Instead, we use a fixed-point iteration method to compute the solution to the equation system. More specifically, we construct the following two iterative formulas based on Equation (14) and Equation (15), where (k) refers to the iteration index. Then, we iteratively compute x and N until a convergence with an acceptable tiny error between $x^{(k)}$ and $x^{(k+1)}$.

¹The *Heat Distribution* application computes the distribution of the heat for a room over time given a set of initial heat sources. The MPI program splits a particular space into several blocks and computes the heat distribution for each of them in parallel with communicated messages on the shared edges of the blocks. Such a parallel computation is commonly used in real scientific projects such as ocean simulation [27], [28].

²This Nek5000 eddy_uv application monitors the error for a 2D solution to the Navier-Stokes equations [29].

$$x^{(k+1)} = \sqrt{\frac{bN^{(k)}T_e}{2\epsilon_0 g(N^{(k)})}}$$
(16)

$$\frac{T_e b/x^{(k)}}{2g(N^{(k+1)})} - T_e \frac{(1 + \frac{bN^{(k+1)}}{2x^{(k)}})g'(N^{(k+1)})}{g^2(N^{(k+1)})} + b(\eta_0 + A) = 0 \quad (17)$$

In fact, solving Equation (17) directly is still very challenging, but we can compute the approximate root $N^{(k+1)}$ for Equation (17) efficiently by using a bisection method. Specifically, there must be an ideal number of cores (denoted by $N^{(*)}$ for any application because of the inevitable synchronization/communication among processes. Since $\frac{\partial E^2(T_w)}{\partial N^2}$ must always be greater than or equal to 0 in the range $[0, N^{(*)}]$, and the optimal solution N^* must be no greater than $N^{(*)}$, there must be at most one root in $[0, N^{(*)}]$ to satisfy $\frac{\partial E(T_w)}{\partial N}$ =0. If one root does exist in $[0, N^{(*)}]$, we can use bisection method to approximate it, because the lefthand side of Formula (17) is a monotonically increasing function (note that $\frac{\partial E^2(T_w)}{\partial N^2} > 0$). Since N^* must be an integer, the bisection method can stop whenever the error of N between two adjacent steps (such as step k and step k+1) is smaller than 0.5; thus the convergence speed could be further improved (our simulation shows there are only about 10 iteration steps in general). On the other hand, if no root exists in $[0, N^{(*)}]$, the optimal number of processes N^* must be equal to $N^{(*)}$. This situation occurs with very few failures or small checkpoint overhead on the PFS.

Based on the Heat Distribution application speedup, we perform a numerical simulation to confirm the correctness of our optimal derivation. In the simulation, checkpoint overheads are based on our characterization of FTI on the Fusion cluster. The coefficients of the speedup functions are computed by using a least squares method based on real experimental data as shown in Figure 2. The error threshold is set to 10^{-6} , and *x*'s initial value is set to 100,000. Our iterative method needs just 30–40 iterations to converge, which means a fairly high convergence speed.

Figure 3 presents different wall-clock times when running the Heat Distribution application, comparing the optimal solution computed by our method with other solutions. The workload to process is 4,000 core-days and the original optimal execution scale $N^{(*)}$ is set to 100,000 cores. In addition, b=0.005, and $\kappa=0.46$. As for the two subfigures, the checkpoint overheads are set to constant values (C(N)=R(N)=5 seconds) and linear-increasing values (C(N)=R(N)=5+0.005N), respectively. The optimal number of checkpoint intervals and the optimal number of processes/cores are 797 and 81,746, respectively, for the case with constant checkpoint cost; and their values are 140 and 20,215, respectively, for the case with linear-increasing checkpoint cost. Note that the original optimal execution scale without checkpoints and failure events is 100,000 cores for the Heat Distribution application. Thus, the optimal execution scale changes prominently because of the impact of the failure rate that increases with the execution scale and checkpoint overheads. More experimental evaluation results can be found in Section IV.



Figure 3. Confirming the Optimal Solution via Numerical Simulation

D. Optimization of the Multilevel Model

Since multiple checkpoint levels are involved in the application's execution, we need to combine them in a target function, namely, Formula (1). Compared with the single-level checkpoint model, the multilevel checkpoint model has two critical differences for the derivation of the expected wall-clock length $E(T_w)$.

• The roll-back loss in the multilevel checkpoint model is different from that in the traditional single-level checkpoint model. Since the application is restarted based on a checkpoint at level *i*, the total roll-back loss at this level must include all checkpoint overheads at lower levels in addition to the lost execution time. For example, when the application rolls back to a level-3 checkpoint, both the level-1 checkpoint overheads and level-2 checkpoint overheads must be counted in the roll-back loss. On the other hand, since all failures are unpredictable with random arrival locations and the checkpoints are taken periodically with equal distances, the expected value of Γ_{ij} can be represented as Formula (18), where $\frac{f(T_e,N)/(2x_i)}{f(T_e,N)/x_k}$ refers to the number of checkpoints at level *k* during the roll-back period.

$$E(\Gamma_{ij}) = \frac{f(T_e, N)}{2x_i} + \frac{C_i(N)}{2} + \sum_{\substack{k=1\\k=1}}^{i-1} \left(\frac{\frac{f(T_e, N)}{2x_i}}{\frac{f(T_e, N)}{x_k}} C_k(N) \right)$$
(18)
$$= \frac{f(T_e, N)}{2x_i} + \sum_{\substack{k=1\\k=1}}^{i} \frac{C_k(N)x_k}{2x_i}$$

• In the multilevel model, we need to investigate how to represent checkpoint overhead $C_i(N)$ and recovery overhead $R_i(N)$ on each level *i*. In general, if we save checkpoint files to local storage devices of running processes, the checkpoint overhead is a constant because these is no congestion; if we use the PFS to save checkpoint files, the checkpoint overhead may increase with the execution scale because of more checkpoint files (thus more metadata) to be handled by the PFS and inevitable congestion. Our characterization of the Heat Distribution application's checkpoint overheads on the Argonne Fusion cluster (Table II¹) shows that the

¹The checkpoint overheads characterized in this work are lower than those of our previous work [22] because of the improved FTI.

checkpoint overheads on level 1, 2, and 3 are stable, whereas they increase prominently on the PFS.

Table II

CHECKPOINT OVERHEAD OF FTI (IN SECONDS)

Exe. Scale	Ckpt Cost (level 1–4)				
128 cores	0.9	2.53	3.7	7	
256 cores	0.67	2.54	4.1	8.1	
384 cores	0.67	2.25	3.9	14.3	
512 cores	0.99	3.05	4.12	21.3	
1024 cores	1.1	2.56	3.61	25.15	

Based on the characterization of checkpoint overheads shown in this table, we define the checkpoint overhead $C_i(N)$ and recovery overhead $R_i(N)$ as Formula (19) and Formula (20), respectively.

$$C_i(N) = \epsilon_i + \alpha_i H_c(N) \tag{19}$$

$$R_i(N) = \eta_i + \beta_i H_r(N) \tag{20}$$

Here $i=1,2,\cdots,L$ refers to checkpoint level; N refers to the number of cores; ϵ_i , α_i , η_i , and β_i are four constant coefficients with respect to level i and can be derived by a least squares fitting method; and $H_c(N)$ and $H_r(N)$ are two baseline functions that always pass through (0,0). For example, $H_c(N)=H_r(N)=0$ means constant checkpoint/recovery overhead, and $H_c(N)=H_r(N)=N$ implies linear-increasing checkpoint/recovery overhead. All the coefficients and baseline functions in Formula (19) and (20) are supposed to be determined based on characterization of checkpoint/recovery overheads with different execution scales. With respect to Table II, since the checkpoint overheads for the first three levels look like constants, $\alpha_1 = \alpha_2 = \alpha_3 = 0$ approximately holds.

Now, let us derive the optimal solution (to determine the optimal checkpoint intervals for each level and optimize the execution scale) to our multilevel checkpoint model. Combining Formula (1) and Formula (18), we derive Formula (21), where μ_i is the expected number of failures belonging to level *i*, as presented by Formula (22) in which $P_i(Y=K)$ denotes the probability of the number of failure events during the execution. Note that μ_i is actually a variate of the execution scale N because the failure probability may change with execution scales.

$$E(T_w) = \frac{T_e}{g(N)} + \sum_{i=1}^{L} C_i(N)(x_i - 1) + \sum_{i=1}^{L} \left[\mu_i \left(\frac{T_e/g(N)}{2x_i} + \sum_{k=1}^{i} \frac{C_k(N) \cdot x_k}{2x_i} + (A + R_i(N)) \right) \right]^{(21)} \\ \mu_i = E_i(Y) = \sum_{K=1}^{\infty} K \cdot P_i(Y = K)$$
(22)

Similar to the single-level model presented in Section III-C, we can derive $\frac{\partial E^2(T_w)}{\partial x_i^2} > 0$ and $\frac{\partial E^2(T_w)}{\partial N^2} > 0$, so there must be a unique optimum point for $\{x_1, x_2, \cdots, x_L, N\}$ with minimized $E(T_w)$. Similar to the derivation of the optimal solution for single-level checkpoint model, we can get the optimal solution to such a multilevel checkpoint model as

follows (based on first-order necessary conditions):

$$\frac{\partial E(T_w)}{\partial x_i} = C_i - \frac{\mu_i}{2x_i^2} \left(\frac{T_e}{g(N)} + \sum_{j=1}^{i-1} C_j x_j \right) + \frac{C_i}{2} \sum_{j=i+1}^{L} \frac{\mu_j}{x_j} = 0$$

$$\frac{\partial E}{\partial N} = \frac{T_e}{g^2(N)} \left(\sum_{i=1}^{L} \frac{\mu_i'}{2x_i} g(N) - (1 + \sum_{i=1}^{L} \frac{\mu_i}{2x_i}) g'(N) \right)$$
(23)

$$+ \sum_{i=1}^{L} C'_{i}(x_{i} - 1)$$

$$+ \sum_{i=1}^{L} \left[\mu'_{i}(\sum_{k=1}^{i} \frac{C_{k}x_{k}}{2x_{i}} + A + R_{i}) + \mu_{i}(\sum_{k=1}^{i} \frac{C'_{k}x_{k}}{2x_{i}} + R'_{i}) \right] = 0$$

$$(24)$$

Note that Formula (23) represents a set of equations, where $i=1,2,\dots,L$. For simple representation, we use C_i and R_i to denote $C_i(N)$ and $R_i(N)$, respectively, which are computed by Formula (19) and Formula (20); C'_i and R'_i denote their derivatives, respectively.

In principle, as long as we find the solution to the above system of L+1 simultaneous equations, we find the optimal solution to the multilevel checkpoint problem that expected numbers of failure events at different levels are assumed to be related only to the execution scale N (i.e., $\mu_i = \mu_i(N)$). A direct solution is extremely difficult, however, because of the high degrees in the equations and multiple variables (including x_1, \dots, x_L and N). Instead we can use fixedpoint iteration to obtain the solution. Specifically, based on Formula (23) and Formula (24), we can construct their corresponding iterative functions and alternatively compute x_i and N based on Formula (23) and Formula (24), until each equation approximately holds with little error. The initial values of $\{x_1, x_2, \dots, x_L\}$ (as shown in Formula (25)) can be obtained by Young's formula [3], in that it leads to the suboptimal checkpoint interval result for a particular level *i* without taking into account the impact of checkpoint overheads at other levels. Based on our model, Young's formula can be approximately presented as Formula (25), where $\frac{T_e}{q(N)}$ denotes the expected productive time without checkpoint model and $C_i(N)$ refers to the checkpoint overhead at level i when running with N processes/cores.

$$x_i = \sqrt{\frac{\mu_i(N)T_e/g(N)}{2C_i(N)}} \tag{25}$$

As analyzed in Section III-B, the estimated wall-clock time may change with changed numbers of failures, which will change the expected numbers of failures at each level in turn. Hence, our complete solution iteratively computes the new wall-clock time based on changing expected numbers of failures for the application until convergence, as presented in Algorithm 1. The experimental results are presented in Section IV.

One critical question is whether our key algorithm (Algorithm 1) can always converge eventually or in what situations it cannot converge. In fact, this algorithm cannot converge in one situation only, namely, when failure rates are extremely high, which would not happen in reality. If failure rates (i.e., μ_i) are fairly high, the new value of $E(T_w)$ in each iteration would become much larger than its old value computed in last iteration, such that the new expected failure rates μ'_i may become unexpectedly larger in turn. In our evaluation (shown in next section), the failure rate is set up to 16+12+8+4=40 failures per day, which is already very high. Algorithm 1 can still converge quickly in this situation, which means that convergence issue is not a concern here.

IV. PERFORMANCE EVALUATION

In this section we frist describe the experimental setup. We then present our results.

A. Experimental Setting

Since our fault-tolerance research is designed for exascale applications, it is best to perform the evaluation with hundreds of thousands of real cores. However, there are at most 128 physical nodes (with totally 1,024 cores) available in the Argonne Fusion cluster [26] with a limited resource usage quota, so we have to use exascale simulation to evaluate our multilevel checkpoint model. We perform practical experiments deployed with FTI and real MPI programs on Fusion to validate and confirm the accuracy of our simulation environment.

The application used in our experiment is a well-known MPI program, called *Heat Distribution*, whose MPI communication methods (such as the ghost array between adjacent blocks) are commonly adopted in real scientific projects such as parallel ocean simulation [27]. The key reason we adopted this application in our evaluation is that it is a real MPI program and it is also suitable to be executed with a large number of cores, as shown in Figure 2. It involves many MPI functions, including MPI_Bcast, MPI_Barrier, MPI_Recv, MPI_Send, MPI_Irecv, MPI_Isend, MPI_Waitall, and MPI_Allreduce. The checkpoint overhead and recovery overhead are both dependent on two factors: the program's memory sizes, which are determined by the problem size, and the execution scale (i.e., the number of processes)

The exascale simulation is described as follows. Each test is performed by running the MPI program for processing an amount of workload (T_e) , which corresponds to the singlecore productive time. Each test is driven by ticks (one tick is equal to one second in the simulation), simulating the whole procedure of running an MPI program with a nonlinear speedup that follows Formula (12). We adopt the real checkpoint/restart overhead characterized on Fusion [26], and we also take into account the possible jittering of checkpoint/restart overheads (with random error ratio up to 30%). The checkpoint overheads are shown in Table II, where the least-squares-fitting coefficients (ϵ_i, α_i) are computed as (0.866,0), (2.586,0), (3.886,0), and (5.5,0.0212), respectively. The simulator comprehensively takes into account possible overlapping of different operations/events under the checkpoint model, such as simultaneous occurrence of taking checkpoint/recovery operations and failure events. The whole simulation closely reproduces real experiments, as shown in Figure 4(a) and (b) with various checkpoint intervals on the four different levels. With the same setting, the simulation results are similar to those on real cluster environments, with the difference being less than 4%.



We perform the evaluation for six cases each with different failure rates at the four checkpoint levels. Given a certain number of cores simultaneously used as the baseline (denoted by N_b , where the subscript *b* refers to baseline), r_1 - r_2 - r_3 - r_4 denotes that $r_1/r_2/r_3/r_4$ failure events occur per day at level 1/2/3/4, respectively. For example, 8-4-2-1 means there are 8 failure events occurring each day at level 1, 4 failure events occurring per day at level 2, and so forth. The real failure rates experienced actually increase with the number of cores proportionally, as compared with the baseline number of cores N_b which is always set to $N^{(*)}=10^6$. Each failure may occur randomly at any time in the whole wall-clock period, including productive time and checkpoint/recovery period. The failure intervals follow exponential distribution, because this is the behavior of the system for most of its lifetime [37]. All results shown in the following text are mean values based on 100 runs for each case with random failure events.

In our evaluation we compare four solutions.

- *ML(opt-scale)*: multilevel model with optimized checkpoint intervals and optimized scale N*.
 the proposed solution in this paper.
- *SL(opt-scale)*: **single-level** model with **optimized** checkpoint intervals and **optimized** scale *N*^{*} simultaneously.
 - improved Young's formula based on [23].
- *ML(ori-scale)*: multilevel model with optimized checkpoint intervals and original optimal scale N^(*).
 the previous work proposed in [22].
- SL(ori-scale): single-level model with optimized checkpoint intervals and original optimal scale N^(*).
 classic Young's formula [3].

The two key indicators are *wall-clock time* and *efficiency* [30]. The wall-clock time includes productive time and all overheads. We also present different time portions,

including checkpoint overhead, restart overhead, and rollback workload time. The efficiency is also called processor utilization, which is defined as the ratio of the wall-clocktime based speedup¹ to the number of processes/cores used.

B. Experimental Results

Figure 5 presents the four different portions of the running time, based on different checkpoint solutions; wall-clock time is the sum of the four portions. The original optimal execution scale $(N^{(*)})$ is 1 million cores, and the workload amount to process (i.e., total single-core productive time) is 3 million core-days. We zoom in on a particular area in Figure 5 for clear observation of the results. We also present the number of processes/cores used by the optimized-scale solutions (including ML(opt-scale) and SL(opt-scale)) in Table III. The other two solutions, ML(ori-scale) and SL(ori-scale), adopt the original optimal scale, namely, 1 million cores, in the evaluation.



16-12-8-4 8-6-4-2 4-3-2-1 16-8-4-2 8-4-2-1 4-2-1-0.5

Figure 5. Time Analysis (T_e =3m core-days, $N^{(*)}$ =1m cores) Table III

OPTIMIZED EXECUTION SCALES IN MULTILEVEL MODEL AND
Single-Level Model ($T_e = 3 \text{ m core-days}, N^{(*)} = 1 \text{ m cores}$)

Solution	16-12-8-4	8-6-4-2	4-3-2-1	16-8-4-2	8-4-2-1	4-2-1-0.5
ML(opt-scale)	472k	564k	658k	563k	657k	734k
SL(opt-scale)	41k	78.6k	36.7k	53.6k	325k	399k

Based on Figure 5 and Table III, we have two important findings. First, the total wall-clock time decreases with decreasing number of failure events (e.g., from 16-8-4-2 to 4-2-1-0.5), which is reasonable because of reduced total checkpoint/restart overheads and roll-back losses with less failures. Second, our solution ML(opt-scale) always outperforms other approaches significantly. The wall-clock times of our multilevel model with optimized execution scale (N^*) can be reduced by 58–84%, 7–26%, and 79–88% compared with the other three solutions, respectively.

¹The speedup here refers to the ratio of the failure-free single-core productive time to the wall-clock time (including parallel productive time and all overheads). It is different from the original speedup, which does not consider failures and checkpoint/recovery overheads.

We analyze as follows the key reasons why our solution significantly outperforms other approaches, based on the different time portions. With regard to SL(ori-scale), since it adopts the PFS only to store checkpoint files and uses all the 1 million processes/cores to perform the execution, the checkpoint/recovery overheads would be higher than those in multilevel model. On the other hand, the failure rates would also be fairly high because of too many cores being used simultaneously, which will cause an extremely large amount of roll-back loss. In comparison with SL(oriscale), SL(opt-scale) optimizes the execution scales to avoid too many failure events during the execution, so it suffers from very low roll-back loss, as shown in the figure, while its productive time is inevitably fairly long because of significantly reduced execution scales (e.g., only 41k cores are used for 16-12-8-4 use case).

Comparing our new solution ML(opt-scale) to our previous work on ML(ori-scale) [22], we observe that (1) the productive time is always extended because of the smaller execution scales used in the execution and (2) the remaining portions of times are all significantly reduced in our new solution, leading to performance improvements.

Arguably, however, our solution ML(opt-scale) may not always lead to huge performance gains over our previous approach ML(ori-scale), which uses all of the available cores. For example, when running the application with 10 million core-days, the wall-clock length under ML(opt-scale) is less than that of SL(ori-scale) by 4.3–42.3%. The details are shown in Figure 6. The relatively degraded performance gains (compared with the test with T_e =3m core-days) is due mainly to significantly longer productive time, which takes relatively larger portions of the total wall-clock length.



Figure 6. Time Analysis (T_e =10m core-days, $N^{(*)}$ =1m cores)

Figure 7 shows the efficiencies of the above two tests with different workloads. We see that the single-level model with the optimization of execution scales (i.e., SL(opt-scale)) leads to the highest efficiency, in particular because too few cores are used in execution, as shown in Table III. This

solution could save a large amount of available resources and energies like electricity power, however, it is definitely not preferred by users because of over-long wall-clock length induced (as shown in Figure 5 and Figure 6). In comparison to SL(opt-scale), our ML(opt-scale) solution results in much shorter wall-clock time and also keeps a relatively higher efficiency than other solutions, which can satisfy both users and system managers. Specifically, Table III shows that our solution just uses 40-79% cores from all 1 million cores, which significantly saves the resources and improves the system availability.



We also evaluate the checkpoint model, provided that the PFS checkpoint overhead is a constant as the storage scale increases, which occurs when using some special file system such as the Blue Waters File System [38]. Although the PFS can perform parallel I/O to improve its read/write rate, the total checkpoint overhead on the PFS will still be much larger than that on local storage devices, because of the much larger total checkpoint file size handled by the PFS. Accordingly, we suppose the problem size is huge such that checkpoint overheads on the four checkpoint levels are relatively large: 50, 100, 200, and 2,000 seconds, respectively. Table IV presents the evaluation results based on the setting with $T_e = 2$ million core-days and $N^{(*)} = 1$ million cores. We can see that ML(opt-scale) always leads to the highest performance among all solutions. In particular, its wall-clock time is shorter than that of ML(ori-scale) by 3.6-6.5%, and its efficiency is higher than that of ML(oriscale) by 12.9-22.1%. The execution scales optimized by ML(opt-scale) are 860k-940k cores, which improves the system availability by 6-16% in comparison with using up all the available resources.

Sol.	16-12-8-4		8-6-	-4-2	4-3-2-1	
WCT	WCT	Eff	WCT	Eff	WCT	Eff
ML(opt-scale)	14.6	0.158	12.8	0.173	11.1	0.193
SL(opt-scale)	37.3	0.092	23.2	0.123	17.2	0.146
ML(ori-scale)	15.4	0.13	13.4	0.15	11.7	0.171
SL(ori-scale)	890	0.002	892	0.002	890	0.002
ML(opt-scale)	13.1	0.171	11.7	0.186	10.6	0.2
SL(opt-scale)	30.6	0.10	20.4	0.133	16	0.153
ML(ori-scale)	14.2	0.14	12.2	0.164	11.4	0.176
SL(ori-scale)	893	0.002	890	0.002	896	0.002

Table IV EVALUATION OF THE MULTILEVEL CKPT MODEL WITH CONSTANT CKPT COST ON THE PFS: WALL-CLOCK TIME (WCT) AND EFFICIENCY

We alsocheck the number of iterations used to converge the estimated failure rate with changing wall-clock length in Algorithm 1. The error threshold is set to 10^{-12} . For the three evaluation cases, our algorithm just costs 8 iterations, 7 iterations, and 15 iterations, respectively, which confirms fairly fast convergence speed.

V. RELATED WORK

Optimization of exascale parallel computing performance based on the checkpoint/restart model is a fundamental and challenging issue, which has been studied extensively in recent years [5], [6], [7], [14], especially with ever-increasing demand on exascale execution environments [1]. Some basic ideas (e.g., diskless checkpoint [11], [14]) are trying to reduce the checkpoint overheads as much as possible in the exascale environment, such that the checkpoint/restart model can still be kept effective with respect to the entire performance. Many researchers have proposed different models to address the exascale resilience issue.

The multilevel checkpoint/restart model with different levels of checkpoint overheads has been proposed to provide an elastic response to tolerate different types of failures. SCR [12] was the first library that can be leveraged to checkpoint and recover HPC applications based on four storage levels (RAM, Flash, disk, and PFS). It also explores a Markov model to optimize the checkpoint strategies. However, the developers did not take into account the impact of the number of processes/cores on the execution performance; thus the proposed Markov model is not optimized with respect to the execution scales. FTI [13] is another outstanding library that also supports RS-encoding technology [15], [16]. It allows the recovery of the application from lightweight checkpoint files in case of multiple simultaneous hardware failures. However, FTI does not help optimize the checkpoint intervals. In our previous work [22], we proposed a multilevel checkpoint/restart model based on FTI, in which the checkpoint intervals are optimized for different checkpoint levels. It took into consideration many factors such as various checkpoint/recovery overheads and failure rates on different levels; however, the application execution scale (i.e., the number of processes) was not considered in the optimization problem. In comparison with these efforts, in this paper we propose a more comprehensive multilevel checkpoint model that can optimize the checkpoint intervals and the execution scales simultaneously.

In addition, other papers have discussed how to simultaneously optimize execution scales and checkpoint intervals for HPC applications. A typical method was proposed by Jin et al. [23]. We note many key differences between their work and our worsk: (1) their optimization research is based on single-level checkpont/restart model, which is much simpler than the multilevel checkpoint model tackled by our work; (2) their work is not based on a real checkpoint toolkit and MPI applications, whereas ours stems from real-world characterization over multilevel checkpoint toolkits; (3) their optimal checkpoint interval is derived by assuming failure's exponential distribution and using first-order approximation, whereas ours is without such an assumption and approximation; and (4) their model defines 30+ notations, whereas our model is more concise: with only about 10 necessary notations (as shown in Table I). Perhaps most significant, they derived the target function $E(T_w)$, but they did not prove that it is a convex function with respect to the variables before using Newton's method to approximate the optimal solution. In that situation, the converged result may not be globally optimized; and even worse, the algorithm may not converge given inappropriate initial values. In contrast, we comprehensively analyzed the multilevel checkpoint model with uncertain execution scales, and we conducted a set of exascale simulations to validate its performance using real-world MPI programs and real checkpoint/recovery overheads.

VI. CONCLUSION AND FUTURE WORK

In this paper, we improved our multilevel checkpoint/restart model by optimizing the checkpoint intervals for different checkpoint levels and the number of processes simultaneously. Such a new problem is extremely difficult to solve, not only because of synthetic performance related to different types of failures, but also because of the varied failure probability with uncertain number of processes/cores. However, we have successfully devised an optimized algorithm that works very efficiently. Some key findings are listed below.

- Our algorithm requires just 7–15 iterations to converge.
- The optimized execution scale is smaller than the original optimal scale by 40–95%, which can significantly improve system availability.
- Our solution outperforms other solutions by 4.3– 88% on wall-clock length and improves efficiency by 12.9+%.

In the future, we will explore how to optimize the checkpoint strategies for more complicated nonlinear applications.

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