

Replication Is More Efficient Than You Think

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Replication Is More Efficient Than You Think*

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Abstract: This paper revisits replication coupled with checkpointing for fail-stop errors. Replication enables the application to survive many fail-stop errors, thereby allowing for longer checkpointing periods. Previously published works use replication with the no-restart strategy, which works as follows: (i) compute the application Mean Time To Interruption (MTTI) M as a function of the number of processor pairs and the individual processor Mean Time Between Failures (MTBF); (ii) use checkpointing period $T_{MTTI}^{\rm no} = \sqrt{2MC}$ à la Young/Daly, where C is the checkpoint duration; and (iii) never restart failed processors until the application crashes. We introduce the restart strategy where failed processors are restarted after each checkpoint. We compute the optimal checkpointing period $T_{opt}^{\rm rs}$ for this strategy, which is much larger than $T_{opt}^{\rm no}$ thereby decreasing I/O pressure. We show through simulations that using $T_{opt}^{\rm rs}$ and the restart strategy, instead of $T_{MTTI}^{\rm no}$ and the usual no-restart strategy, significantly decreases the overhead induced by replication, in terms of both total execution time and energy consumption.

Key-words: replication, checkpoint, optimal checkpointing period, restart strategy.

Note: A shorter version of this work appears in the proceedings of SC'19, the 2019 ACM/IEEE International Conference for High Performance Computing, Networking, Storage, and Analysis.

La réplication est plus efficace que vous ne le pensez

Résumé: Cet article revisite la réplication couplée au checkpoint pour les erreurs fatales. La réplication permet à l'application de survivre à plusieurs erreurs, allongeant de fait les périodes de checkpoint. Les anciens travaux sur la réplication utilisent la stratégie no-restart, qui fonctionne de la facon suivante: (i) calculer le Temps Moyen D'Interruption (MTTI) de l'application M en fonction du nombre de paires de processeurs et du Temps Moyen Entre chaque Erreur (MTBF) individuel de chaque processeur; (ii) utiliser la période de checkpoint $T_{MTTI}^{\text{no}} = \sqrt{2MC}$ à la Young/Daly, où C est la durée du checkpoint; et (iii) ne jamais redémarrer un processeur tombé en panne tant que l'application ne s'interrompt pas totalement. Nous présentons la stratégie restart où les processeurs en panne sont redémarrés à chaque checkpoint, ce qui peut augmenter le coût d'un checkpoint mais permet à la configuration de l'application de ne pas se dégrader au fil des périodes de checkpoint. Nous montrons comment calculer la période de checkpoint optimale T_{opt}^{rs} pour la stratégie restart et nous prouvons qu'elle est d'un ordre de grandeur plus grande que T_{MTTI}^{no} . Nous montrons à travers des simulations qu'utiliser T_{opt}^{rs} et la stratégie restart, au lieu de T_{MTTI}^{no} et la stratégie classique no-restart, décroit significativement le coût additionnel lié à la réplication, à la fois en terme de temps d'exécution et de consommation d'énergie.

Mots-clés : réplication, checkpoint, période optimale de checkpoint, stratégie de redémarrage.

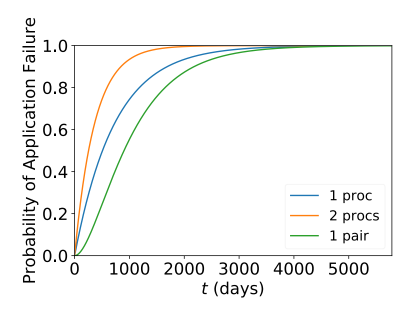
1 Introduction

Current computing platforms have millions of cores: the Summit system at the Oak Ridge National Laboratory (ORNL) is listed at number one in the TOP500 ranking [38], and it has more than two million cores. The Chinese Sunway TaihuLight (ranked as number 3) has even more than 10 million cores. These large-scale computing systems are frequently confronted with failures, also called fail-stop errors. Indeed, even if individual cores are reliable, for instance if the *Mean Time Between Failures* (MTBF) for a core is $\mu = 10$ years, then the MTBF for a platform with a million cores $(N = 10^6)$ becomes $\mu_N = \frac{\mu}{N} \approx 5.2$ minutes, meaning that a failure strikes the platform every five minutes, as shown in [24].

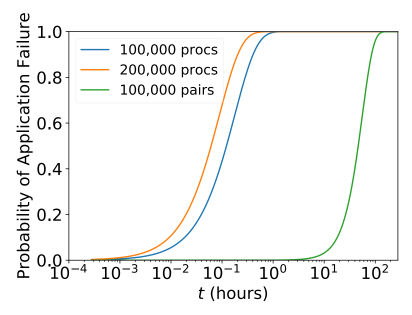
The classical technique to deal with failures consists of using a checkpoint-restart mechanism: the state of the application is periodically checkpointed, and when a failure occurs, we recover from the last valid checkpoint and resume the execution from that point on, rather than starting the execution from scratch. The key for an efficient checkpointing policy is to decide how often to checkpoint. Young [42] and Daly [13] derived the well-known Young/Daly formula $T_{YD} = \sqrt{2\mu_N C}$ for the optimal checkpointing period, where μ_N is the platform MTBF, and C is the checkpointing duration.

Another technique that has been advocated for dealing with failures is process replication, where each process in a parallel MPI (Message Passing Interface) application is duplicated to increase the Mean Time To Interruption (MTTI). The MTTI is the mean time between two application failures. If a process is struck by a failure, the execution can continue until the replica of this process is also struck by a failure. More precisely, processors are arranged by pairs, i.e., each processor has a replica, and the application fails whenever both processors in a same pair have been struck by a failure. With replication, one considers the MTTI rather than the MTBF, because the application can survive many failures before crashing. Given the high rate of failures on large-scale systems, process replication is combined with periodic checkpoint-restart, as proposed for instance in [35, 45, 18] for high-performance computing (HPC) platforms, and in [28, 41] for grid computing. Then, when the application fails, one can recover from the last valid checkpoint, just as was the case without replication. Intuitively, since many failures are needed to interrupt the application, the checkpointing period should be much larger than without replication. Previous works [20, 11, 25] all use $T_{MTTI}^{\text{no}} = \sqrt{2M_NC}$ for the checkpointing period, where M_N is the MTTI with N processors (instead of the MTBF μ_N).

To illustrate the impact of replication on reliability at scale, Figure 1 compares the probability distribution of the time to application failure for: (a) a single processor, two parallel processors and a pair of replicated processors; and (b) a platform of N = 100,000 parallel processors without replication, and b = 100,000 processor pairs



(a) CDFs of the probability distribution of time to app. failure for one processor, two parallel processors and one proc. pair (replication).



(b) CDFs of the proba. distrib. of time to app. failure for 100,000 parallel proc., 200,0000 parallel proc. and 100,000 proc. pairs (replication).

Figure 1: Comparison of CDFs with and without replication.

with replication. In all cases, the individual MTBF of a single processor is $\mu=5$ years. The time to reach 90% chances of having a fatal failure is: (a) 1688 days for one processor, 844 days for two processors and 2178 days for a processor pair; and (b) 24 minutes for 100,000 processors, 12 minutes for 200,000 processors and 5081 minutes (almost 85 hours) for 100,000 processor pairs. We see that replication is key to safe application progress at scale! Again, the cost is that half of the resources are doing redundant work, hence time-to-solution is increased. We compare time-to-solution with and without replication in Section 7.6. We also see that in heavily failure-prone environments (small MTBF values), checkpoint/restart alone cannot ensure full reliability, and must be complemented by replication.

One major contribution of this paper is to introduce a new approach that minimizes the overhead incurred by the checkpoint-restart mechanism when coupled with replication. Previous works [20, 11, 25] use the no-restart strategy: if a processor was struck by a failure (but not its replica), then the processor remains failed (no recovery) until the whole application fails. Hence, there is a recovery only every M_N seconds on average, whenever the application fails. Many periodic checkpoints are taken in between two application crashes, with more and more processors failing on the fly. To the best of our knowledge, analytically computing the optimal period for no-restart is an open problem (see Section 4.2 for more details, where we also show that non-periodic strategies are more efficient for no-restart), but simulations can help assess this approach.

The study of the *no-restart* strategy raises an important question: should failed processors be restarted earlier on in the execution? Instead of waiting for an application crash to rejuvenate the whole platform, a simple approach would be to restart processors immediately after each failure. Let restart-on-failure denote this strategy. It ensures that all processor pairs involve two live processors throughout execution, and would even suppress the notion of checkpointing periods. Instead, after each failure striking a processor, its replica would checkpoint immediately, and the spare processor replacing the failed processor would read that checkpoint to resume execution. There is a small risk of fatal crash if a second failure should strike the replica when writing its checkpoint, but (i) the risk is very small because the probability of such a cascade of two narrowly spaced failures is quite low; and (ii) if the checkpoint protocol is scalable, every other processor can checkpoint in parallel with the replica, and there is no additional time overhead. With tightly coupled applications, the other processors would likely have to wait until the spare is able to restart, and they can checkpoint instead of idling during that wait. While intuitively appealing, the restart-on-failure strategy may lead to too many checkpoints and restarts, especially in scenarios when failures strike frequently. However, frequent failures were exactly the reason to deploy replication in the first place, precisely to avoid having to restart after each failure.

In this work, we introduce the *restart* strategy, which requires any failed processor to recover each time a checkpoint is taken. This ensures that after any checkpoint at the end of a successful period, all processors are alive. This is a middle ground between the *no-restart* and *restart-on-failure* strategies, because failed processors are restarted at the end of each period with *restart*. On the one hand, a given period may well include many failures, hence *restart* restarts processors less frequently than *restart-on-failure*. On the other hand, there will be several periods in between two application crashes, hence *restart* restarts processors more frequently than *no-restart*.

Periodic checkpointing is optimal with the restart strategy: the next period should have same length as the previous one, because we have the same initial conditions at the beginning of each period. Restarting failed processors when checkpointing can introduce additional overhead, but we show that it is very small, and even non-existent when in-memory (a.k.a. buddy) checkpointing is used as the first-level of a hierarchical multi-level checkpointing protocol (such state-of-the-art protocols are routinely deployed on large-scale platforms [3, 29, 10]). A key contribution of this paper is a mathematical analysis of the restart strategy, with a closed-form formula for its optimal checkpointing period. We show that the optimal checkpointing period for the restart strategy has the order $\Theta(\mu^{\frac{1}{3}})$, instead of the $\Theta(\mu^{\frac{1}{2}})$ used in previous works for no-restart as an extension of the Young/Daly formula [20, 11, 25]. Hence, as the error rate increases, the optimal period becomes much longer than the value that has been used in all previous works (with *no-restart*). Consequently, checkpoints are much less frequent, thereby dramatically decreasing the pressure on the I/O system.

The main contributions of this paper are the following:

- We provide the first closed-form expression of the application MTTI M_N with replication;
- We introduce the *restart* strategy for replication, where we recover failed processors during each checkpoint;
- We formally analyze the *restart* strategy, and provide the optimal check-pointing period with this strategy;
- We apply these results to applications following Amdahl's law, i.e., applications that are not fully parallel but have an inherent sequential part, and compare the time-to-solution achieved with and without replication;
- We validate the model through comprehensive simulations, by showing that analytical results, using first-order approximations and making some additional assumptions (no failures during checkpoint and recovery), are quite close to simulation results; for these simulations, we use both randomly generated failures and log traces.
- We compare through simulations the overhead obtained with the optimal strategy introduced in this work (*restart* strategy, optimal checkpointing period) to those used in all previous works (*no-restart* strategy, extension

of the Young/Daly checkpointing period), as well as with strategies that use partial replication or that restart only at some of the checkpoints, and demonstrate that we can significantly decrease both total execution time and utilization of the I/O file system.

• Finally, we show that similarly good results are obtained when aiming at minimizing the *energy consumption* of the application, instead of its total execution time.

The paper is organized as follows. We first describe the model in Section 2. We recall how to compute the optimal checkpointing period when no replication is used in Section 3. The core contribution is presented in Section 4, where we explain how to compute the MTTI with $b = \frac{N}{2}$ processor pairs, detail the restart strategy, and show how to derive the optimal checkpointing period with this restart strategy. Results are applied to applications following Amdahl's law in Section 5. An asymptotic analysis of no-restart and restart is provided in Section 6. The experimental evaluation in Section 7 presents extensive simulation results, demonstrating that replication is indeed more efficient than you think, when enforcing the restart strategy instead of the no-restart strategy. We discuss related work in Section 8, and conclude in Section 9. Finally, results for energy consumption are presented in Section A.

2 Model

This section describes the model, with an emphasis on the cost of a combined checkpoint-restart operation. We differ the description of energy-related parameters to Section A.

Fail-stop errors. Throughout the text, we consider a platform with N identical processors. The platform is subject to fail-stop errors, or failures, that interrupt the application. Similarly to previous work [25, 20, 17], for the mathematical analysis, we assume that errors are independent and identically distributed (IID), and that they strike each processor according to an exponential probability distribution $\exp(\lambda)$ with support $[0, \infty)$, probability density function (PDF) $f(t) = \lambda e^{-\lambda t}$ and cumulative distribution function (CDF) $F(T) = \mathbb{P}(X \leq T) = 1 - e^{-\lambda T}$. We also introduce the reliability function $G(T) = 1 - F(T) = e^{-\lambda T}$. The expected value $\mu = \frac{1}{\lambda}$ of the $\exp(\lambda)$ distribution is the MTBF on one processor. We lift the IID assumption in the performance evaluation section by using trace logs from real platforms.

Checkpointing. To cope with errors, we use periodic coordinated checkpointing. We assume that the divisible application executes for a very long time (asymptotically infinite) and we partition the execution into periods. Each period \mathcal{P} consists of a work segment of duration T followed by a check-point of duration C. After an error, there is a downtime of duration D (corresponding to the time needed to migrate to a spare processor), a recovery of size R, and then one needs to re-execute the period from its beginning.

Replication. We use another fault tolerance technique, namely replication. Each process has a replica, which follows the exact same states in its execution. To ensure this, when a process receives a message, its replica also receives the same message, and messages are delivered in the same order to the application (an approach called *active* replication; see [23, 20]). If a crash hits a process at any time, and its replica is still alive, the replica continues the execution alone until a new process can replace the dead one.

We rely on the traditional process allocation strategy that assigns processes and their replicas on remote parts of the system (typically different racks) [8]. This strategy mitigates the risk that a process and its replica would both fail within a short time interval (much shorter than the expected MTTI). As stated in [16], when failure correlations are observed, their correlation diminishes when the processes are far away from each other in the memory hierarchy, and becomes undistinguishable from the null hypothesis (no correlation) when processes belong to different racks.

Combined checkpoint-restart. In this paper, we propose the restart strategy where failed processes are restarted as soon as the next checkpoint wave happens. When that happens, and processes need to be restarted, the cost of a checkpoint and restart wave, C^R , is then increased: one instance of each surviving process must save their state, then processes for the missing instances of the replicas must be allocated; the new processes must load the current state, which has been checkpointed, and join the system to start acting as a replica. The first part of the restart operation, allocating processes to replace the failed ones, can be managed in parallel with the checkpoint of the surviving processes. Using spare processes, this allocation time can be very small and we will consider it negligible compared to the checkpoint saving and loading times. Similarly, integrating the newly spawned process inside the communication system when using spares is negligible when using mechanisms such as the ones described in [7].

There is a large variety of checkpointing libraries and approaches to help applications save their state. [29, 3, 10] are typically used in HPC systems for coordinated checkpointing, and use the entire memory hierarchy to speed up the checkpointing cost: the checkpoint is first saved on local memory, then uploaded onto local storage (SSD, NVRAM if available), and eventually to the shared file system. As soon as a copy of the state is available on the closest memory, the checkpoint is considered as taken. Loading that checkpoint requires that the application state from the closest memory be

sent to the memory of the new hosting process.

Another efficient approach to checkpoint is to use in-memory checkpoint replication using the memory of a 'buddy' process (see [31, 44]). To manage the risk of losing the checkpoint in case of failure of two buddy processes, the checkpoint must also be saved on reliable media, as is done in the approaches above. Importantly, in-memory checkpointing is particularly fitted for the restart strategy, because the buddy process and the replica are the same process: in that case, the surviving processes upload their checkpoint directly onto the memory of the newly spawned replicas; as soon as this communication is done, the processes can continue working. Contrary to traditional buddy checkpointing, it is not necessary to exchange the checkpoints between a pair of surviving buddies since, per the replication technique, both checkpoints are identical.

In the worst case, if a sequential approach is used, combining checkpointing and restart takes at most twice the time to checkpoint only; in the best case, using buddy checkpointing, the overhead of adding the restart to the checkpoint is negligible. We consider the full spectrum $C \leq C^R \leq 2C$ in the simulations.

As discussed in [20, 32], checkpoint time varies significantly depending upon the target application and the hardware capabilities. We will consider a time to checkpoint within two reasonable limits: $60s \le C \le 600s$, following [25].

First-order approximation. Throughout the paper, we are interested in first-order approximations, because exact formulas are not analytically tractable. We carefully state the underlying hypotheses that are needed to enforce the validity of first-order results. Basically, the first-order approximation will be the first, and most meaningful, term of the Taylor expansion of the overhead occurring every period when the error rate λ tends to zero.

3 Background

In this section, we briefly summarize well-known results on the optimal checkpointing period when replication is not used, starting with a single processor in Section 3.1, and then generalizing to the case with N processors in Section 3.2.

3.1 With a Single Processor

We aim at computing the expected time $\mathbb{E}(T)$ to execute a period of length $\mathcal{P} = T + C$. The optimal period length will be obtained for the value of T, minimizing the overhead

$$\mathbb{H}(T) = \frac{\mathbb{E}(T)}{T} - 1. \tag{1}$$

We temporarily assume that fail-stop errors strike only during work T and not during checkpoint C nor recovery R. In fact, this assumption has no impact on the first-order approximation of the period, as shown below. The following recursive equation is the key to most derivations:

$$\mathbb{E}(T) = (1 - F(T))(T + C) + F(T)(T_{lost}(T) + D + R + \mathbb{E}(T)). \tag{2}$$

Equation (2) reads as follows: with probability 1 - F(T), the execution is successful and lasts T + C seconds; with probability F(T), an error strikes before completion, and we need to account for time lost $T_{\text{lost}}(T)$, downtime D and recovery R before starting the computation anew. The expression for $T_{\text{lost}}(T)$ is the following:

$$T_{\text{lost}}(T) = \int_0^\infty t \mathbb{P}(X = t | X \le T) dt = \frac{1}{F(T)} \int_0^T t f(t) dt.$$

Integrating by parts and re-arranging terms in Equation (2), we derive $\mathbb{E}(T) = T + C + \frac{F(T)}{1-F(T)}(T_{\text{lost}}(T) + D + R)$ and $\mathbb{H}(T) = \frac{C}{T} + \frac{F(T)}{T(1-F(T))}(D + R) + \frac{\int_0^T G(t)dt}{T(1-F(T))} - 1$. Now, if we instantiate the value of $F(T) = 1 - G(T) = 1 - e^{-\lambda T}$, we obtain $\mathbb{H}(T) = \frac{C}{T} + \frac{e^{\lambda T} - 1}{T}(D + R + \frac{1}{\lambda}) - 1$. We can find the value T_{opt} by differentiating and searching for the zero of the derivative, but the solution is complicated as it involves the Lambert function [13, 24]. Instead, we use the Taylor expansion of $e^{-\lambda T} = \sum_{i=0}^{\infty} (-1)^i \frac{(\lambda T)^i}{i!}$ and the approximation $e^{-\lambda T} = 1 - \lambda T + \frac{(\lambda T)^2}{2} + o(\lambda^2 T^2)$. This makes sense only if λT tends to zero. It is reasonable to make this assumption, since the length of the period $\mathcal P$ must be much smaller than the error MTBF $\mu = \frac{1}{\lambda}$. Hence, we look for $T = \Theta(\lambda^{-x})$, where 0 < x < 1. Note that x represents the order of magnitude of T as a function of the error rate λ . We can then safely write

$$\mathbb{H}(T) = \frac{C}{T} + \frac{\lambda T}{2} + o(\lambda T). \tag{3}$$

Now, $\frac{C}{T} = \Theta(\lambda^x)$ and $\frac{\lambda T}{2} = \Theta(\lambda^{1-x})$, hence the order of magnitude of the overhead is $\mathbb{H}(T) = \Theta(\lambda^{\max(x,1-x)})$, which is minimum for $x = \frac{1}{2}$. Differentiating Equation (3), we obtain

$$T_{opt} = \sqrt{\frac{2C}{\lambda}} = \Theta(\lambda^{-\frac{1}{2}}), \text{ and } \mathbb{H}_{opt} = \sqrt{2C\lambda} + o(\lambda^{\frac{1}{2}}) = \Theta(\lambda^{\frac{1}{2}})$$
 (4)

which is the well-known and original Young formula [42].

Variants of Equation (4) have been proposed in the literature, such as $T_{opt} = \sqrt{2(\mu + R)C}$ in [13] or $T_{opt} = \sqrt{2(\mu - D - R)C} - C$ in [24]. All variants are approximations that collapse to Equation (4). This is because the resilience parameters C, D, and R are constants and thus negligible in front of T_{opt} when λ tends to zero. This also explains that assuming that fail-stop errors may strike during checkpoint or recovery has no impact on on

the first-order approximation of the optimal period given in Equation (4). For instance, assuming that fail-stop errors strike during checkpoints, we would modify Equation (2) into

$$\mathbb{E}(T+C) = (1-F(T+C))(T+C) + F(T+C)(T_{lost}(T+C) + D + R + \mathbb{E}(T+C))$$

and derive the same result as in Equation (4). Similarly, assuming that fail-stop errors strike during recovery, we would replace R with $\mathbb{E}(R)$, which can be computed via an equation similar to that for $\mathbb{E}(T)$, again without modifying the final result.

Finally, a very intuitive way to retrieve Equation (4) is the following: consider a period of length $\mathcal{P} = T + C$. There is a failure-free overhead $\frac{C}{T}$, and a failure-induced overhead $\frac{1}{\mu} \times \frac{T}{2}$, because with frequency $\frac{1}{\mu}$ an error strikes, and on average it strikes in the middle of the period and we lose half of it. Adding up both overhead sources gives C

$$\frac{C}{T} + \frac{T}{2\mu},\tag{5}$$

which is minimum when $T = \sqrt{2\mu C}$. While not fully rigorous, this derivation helps understand the tradeoff related to the optimal checkpointing frequency.

3.2 With N Processors

The previous analysis can be directly extended to multiple processors. Indeed, if fail-stop errors strike each processor according to an $\exp(\lambda)$ probability distribution, then these errors strike the whole platform made of Nidentical processors according to an $\exp(N\lambda)$ probability distribution [24]. In other words, the platform MTBF is $\mu_N = \frac{\mu}{N}$, which is intuitive: the number of failures increases linearly with the number of processors N, hence the mean time between two failures is divided by N. All previous derivations apply, and we obtain the optimal checkpointing period and overhead:

$$T_{opt} = \sqrt{\frac{2C}{N\lambda}} = \Theta(\lambda^{-\frac{1}{2}}), \text{ and } \mathbb{H}_{opt} = \sqrt{2CN\lambda} + o(\lambda^{\frac{1}{2}}) = \Theta(\lambda^{\frac{1}{2}})$$
 (6)

This value of T_{opt} can be intuitively retrieved with the same (not fully rigorous) reasoning as before (Equation (5)): in a period of length $\mathcal{P} = T + C$, the failure-free overhead is $\frac{C}{T}$, and the failure-induced overhead becomes $\frac{1}{\mu_N} \times \frac{T}{2}$: we factor in an updated value of the failure frequency, using $\frac{1}{\mu_N} = \frac{N}{\mu}$ $\frac{\overline{\mu_N}}{\overline{\mu_N}} \wedge 2$. We have the instead of $\frac{1}{\mu}$. Both overhead sources add up to $\frac{C}{T} + \frac{T}{2\mu_N} = \frac{C}{T} + \frac{NT}{2\mu},$

$$\frac{C}{T} + \frac{T}{2u_N} = \frac{C}{T} + \frac{NT}{2u},\tag{7}$$

which is minimum when $T = \sqrt{\frac{2\mu C}{N}}$.

4 Replication

This section deals with process replication for fail-stop errors, as introduced in [20] and recently revisited by [25]. We consider a platform with N=2b processors. Exactly as in Section 3, each processor fails according to a probability distribution $\exp(\lambda)$, and the platform MTBF is $\mu_N = \frac{\mu}{N}$. We still assume that checkpoint and recovery are error-free: it simplifies the analysis without modifying the first-order approximation of the optimal checkpointing period.

Processors are arranged by pairs, meaning that each processor has a replica. The application executes as if there were only b available processors, hence with a reduced throughput. However, a single failure does not interrupt the application, because the replica of the failed processor can continue the execution. The application can thus survive many failures, until both replicas of a given pair are struck by a failure. How many failures are needed, in expectation, to interrupt the application? We compute this value in Section 4.1. Then, we proceed to deriving the optimal checkpointing period, first with one processor pair in Section 4.2, before dealing with the general case in Section 4.3.

4.1 Computing the Mean Time To Interruption

Let $n_{\text{fail}}(2b)$ be the expected number of failures to interrupt the application, with b processor pairs. Then, the application MTTI M_{2b} with b processor pairs (hence N=2b processors) is given by

$$M_{2b} = n_{\text{fail}}(2b) \,\mu_{2b} = n_{\text{fail}}(2b) \frac{\mu}{2b} = \frac{n_{\text{fail}}(2b)}{2\lambda b},$$
 (8)

because each failure strikes every μ_{2b} seconds in expectation. Computing the value of $n_{\rm fail}(2b)$ has received considerable attention in previous work. In [34, 20], the authors made an analogy with the birthday problem and use the Ramanujan function [21] to derive the formula $n_{\rm fail}(2b) = 1 + \sum_{k=0}^{b} \frac{b!}{(b-k)!b;k} \approx \sqrt{\frac{\pi b}{2}}$. The analogy is not fully correct, because failures can strike either replica of a pair. A correct recursive formula is provided in [11], albeit without a closed-form expression. Recently, the authors in [25] showed that

$$n_{\text{fail}}(2b) = 2b4^b \int_0^{\frac{1}{2}} x^{b-1} (1-x)^b dx \tag{9}$$

but did not give a closed-form expression either. We provide such an expression below:

$$n_{\text{fail}}(2b) = 1 + 4^b / {2b \choose b}.$$
 (10)

Proof. The integral in Equation (9) is known as the incomplete Beta function $B(\frac{1}{2},b,b+1)$, where $B(z,u,v)=\int_0^z x^{u-1}(1-x)^{v-1}dx$. This incomplete Beta function is also known [40] as the hypergeometric function $B(z,u,v)=\frac{z^u}{u}\times_2F_1\Big[{u,1-v\atop u+1};z\Big]$, where

$${}_{2}F_{1}\left[{u,v\atop w};z\right] = \sum_{n=0}^{\infty} \frac{\langle u\rangle_{n}\langle v\rangle_{n}}{\langle w\rangle_{n}} \frac{z^{n}}{n!} = 1 + \frac{uv}{1!w}z + \frac{u(u+1)v(v+1)}{2!w}z^{2} + \dots$$

We need to compute $B(\frac{1}{2}, b, b+1) = \frac{1}{b2^b} \times {}_2F_1\left[\begin{smallmatrix} b, & -b \\ b+1 \end{smallmatrix}; \frac{1}{2} \right]$, and according to [39], we have

$${}_2F_1\bigg[{b, \ -b \atop b+1}; {1\over 2}\bigg] = {\sqrt{\pi} \ \Gamma(b+1) \over 2^{b+1}} \Big[{1 \over \Gamma(b+1)\Gamma({1\over 2})} + {1 \over \Gamma(b+{1\over 2}))\Gamma(1)}\Big].$$

Here, Γ is the well-known Gamma function extending the factorial over real numbers: $\Gamma(z) = \int_0^\infty x^{z-1} e^{-x} dx$. We have $\Gamma(1) = 1$, $\Gamma(b+1) = b!$, $\Gamma(\frac{1}{2}) = \sqrt{\pi}$, and $\Gamma(b+\frac{1}{2}) = \frac{\sqrt{\pi}(2b)!}{4^b b!}$. Hence,

$$_{2}F_{1}\begin{bmatrix}b, -b \\ b+1\end{bmatrix}; \frac{1}{2} = \frac{1}{2^{b+1}} \left[1 + \frac{4^{b}(b!)^{2}}{(2b)!}\right] = \frac{1}{2^{b+1}} \left[1 + \frac{4^{b}}{\binom{2b}{b}}\right].$$

For the last equality, we observe that $\binom{2b}{b} = \frac{(2b)!}{(b!)^2}$. We derive $B(\frac{1}{2}, b, b+1) = \frac{1}{2b4^b} \left[1 + \frac{4^b}{\binom{2b}{b}}\right]$, and finally $n_{\text{fail}}(2b) = 1 + \frac{4^b}{\binom{2b}{b}}$, which concludes the proof. \square

Using Sterling's formula, we easily derive that $n_{\text{fail}}(2b) \approx \sqrt{\pi b}$, which is 40% more than the value $\sqrt{\frac{\pi b}{2}}$ used in [34, 20].

Plugging the value of $n_{\text{fail}}(2b)$ back in Equation (8) gives the value of the MTTI M_{2b} . As already mentioned, previous works [20, 11, 25] all use the checkpointing period

$$T_{MTTI}^{\text{no}} = \sqrt{2M_{2b}C} \tag{11}$$

to minimize execution time overhead. This value follows from the same derivation as in Equations (5) and (7). Consider a period of length $\mathcal{P}=T+C$. The failure-free overhead is still $\frac{C}{T}$, and the failure-induced overhead becomes $\frac{1}{M_{2b}} \times \frac{T}{2}$: we factor in an updated value of the failure frequency, which now becomes the fatal failure frequency, namely $\frac{1}{M_{2b}}$. Both overhead sources add up to

$$\frac{C}{T} + \frac{T}{2M_{2h}},\tag{12}$$

which is minimum when $T = \sqrt{2M_{2b}C}$.

In the following, we analyze the *restart* strategy. We start with one processor pair (b=1) in Section 4.2, before dealing with the general case in Section 4.3.

4.2 With One Processor Pair

We consider two processors working together as replicas. The failure rate is $\lambda = \frac{1}{\mu}$ for each processor, and the pair MTBF is $\mu_2 = \frac{\mu}{2}$, while the pair MTTI is $M_2 = \frac{3\mu}{2}$ because $n_{\text{fail}}(2) = 3$. We analyze the restart strategy, which restarts a (potentially) failed processor at every checkpoint. Hence, the checkpoint has duration C^R and not C. Consider a period of length $\mathcal{P} = T + C^R$. If one processor fails before the checkpoint but the other survives until reaching it, the period is executed successfully. The period is re-executed only when both processors fail within T seconds. Let $p_1(T)$ denote the probability that both processors fail during T seconds: $p_1(T) = (1 - e^{-\lambda T})^2$. We compute the expected time $\mathbb{E}(T)$ for period of duration $\mathcal{P} = T + C^R$ using the following recursive equation:

$$\mathbb{E}(T) = (1 - p_1(T))(T + C^R) + p_1(T)(T_{lost}(T) + D + R + \mathbb{E}(T)).$$
 (13)

Here, C^R denotes the time to checkpoint, and in addition, to recover whenever one of the two processors had failed during the period. As discussed in Section 2, we have $C \leq C^R \leq C + R$: the value of C^R depends upon the amount of overlap between the checkpoint and the possible recovery of one processor.

Consider the scenario where one processor fails before reaching the end of the period, while the other succeeds and takes the checkpoint. The *no-restart* strategy continues execution, hence pays only for a regular checkpoint of cost C, and when the live processor is struck by a failure (every M_2 seconds on average), we roll back and recover for both processors [20, 11, 25]. However, the new restart strategy requires any failed processor to recover whenever a checkpoint is taken, hence at a cost C^R . This ensures that after any checkpoint at the end of a successful period, we have two live processors, and thus the same initial conditions. Hence, periodic checkpointing is optimal with this strategy. We compare the restart and no-restart strategies through simulations in Section 7.

As before, in Equation (13), $T_{\text{lost}}(T)$ is the average time lost, knowing that both processors have failed before T seconds. While $T_{\text{lost}}(T) \sim \frac{T}{2}$ when considering a single processor, it is no longer the case with a pair of replicas. Indeed, we compute $T_{\text{lost}}(T)$ as follows:

$$T_{\text{lost}}(T) = \int_0^\infty t \mathbb{P}(X = t | t \le T) dt = \frac{1}{p_1(T)} \int_0^T t \frac{d\mathbb{P}(X \le t)}{dt} dt$$
$$= \frac{2\lambda}{(1 - e^{-\lambda T})^2} \int_0^T t (e^{-\lambda t} - e^{-2\lambda t}) dt.$$

After integration, we find that

$$T_{\text{lost}}(T) = \frac{(2e^{-2\lambda T} - 4e^{-\lambda T})\lambda T + e^{-2\lambda T} - 4e^{-\lambda T} + 3}{2\lambda (1 - e^{-\lambda T})^2} = \frac{1}{2\lambda} \frac{u(\lambda T)}{v(\lambda T)},$$

with $u(y) = (2e^{-2y} - 4e^{-y})y + e^{-2y} - 4e^{-y} + 3$ and $v(y) = (1 - e^{-y})^2$.

Assuming that $T=\Theta(\lambda^{-x})$ with 0 < x < 1 as in Section 3.1, then Taylor expansions lead to $u(y)=\frac{4}{3}y^3+o(y^3)$ and $v(y)=y^2+y^3+o(y^3)$ for $y=\lambda T=o(1)$, meaning that $T_{\rm lost}(T)=\frac{1}{2\lambda}\frac{\frac{4\lambda T}{3}+o(\lambda T)}{1+\lambda T+o(\lambda T)}$. Using the division rule, we obtain $T_{\rm lost}(T)=\frac{1}{2\lambda}(\frac{4\lambda T}{3}+o(\lambda T))=\frac{2T}{3}+o(T)$. Note that we lose two thirds of the period with a processor pair rather than one half with a single processor. Plugging back the value of $T_{\rm lost}(T)$ and solving, we obtain:

$$\mathbb{E}(T) = T + C^R + \left(D + R + \frac{(2e^{-2\lambda T} - 4e^{-\lambda T})\lambda T + e^{-2\lambda T} - 4e^{-\lambda T} + 3}{2\lambda(1 - e^{-\lambda T})^2}\right) \cdot \frac{(e^{\lambda T} - 1)^2}{2e^{\lambda T} - 1}.$$
(14)

We then compute the waste $\mathbb{H}^{rs}(T)$ of the restart strategy as follows:

$$\mathbb{H}^{rs}(T) = \frac{\mathbb{E}(T)}{T} - 1 = \frac{C^R}{T} + \frac{2}{3}\lambda^2 T^2 + o(\lambda^2 T^2). \tag{15}$$

Moreover, with $T = \Theta(\lambda^{-x})$, we have $\frac{C^R}{T} = \Theta(\lambda^x)$ and $\frac{2}{3}\lambda^2T^2 = \Theta(\lambda^{2-2x})$, hence $\mathbb{H}^{rs}(T) = \Theta(\lambda^{\max(x,2-2x)})$, which is minimum for $x = \frac{2}{3}$. Differentiating, we readily obtain:

$$T_{opt} = \left(\frac{3C^R}{4\lambda^2}\right)^{\frac{1}{3}} = \Theta(\lambda^{-\frac{2}{3}}),\tag{16}$$

$$\mathbb{H}^{\mathrm{rs}}(T_{opt}) = \left(\frac{3C^R \lambda}{\sqrt{2}}\right)^{\frac{2}{3}} + o(\lambda^{\frac{2}{3}}) = \Theta(\lambda^{\frac{2}{3}}). \tag{17}$$

Note that the optimal period has the order $T_{opt} = \Theta(\lambda^{-\frac{2}{3}}) = \Theta(\mu^{\frac{2}{3}})$, while the extension $\sqrt{2M_2C}$ of the Young/Daly formula has the order $\Theta(\lambda^{-\frac{1}{2}}) = \Theta(\mu^{\frac{1}{2}})$. This means that the optimal period is much longer than the value that has been used in all previous works. This result generalizes to several processor pairs, as shown in Section 4.3. We further discuss asymptotic results in Section 6.

For an intuitive way to retrieve Equation (16), the derivation is similar to that used for Equations (5), (7) and (12). Consider a period of length $\mathcal{P} = T + C^R$. The failure-free overhead is still $\frac{C^R}{T}$, and the failure-induced overhead becomes $\frac{1}{\mu}\frac{T}{\mu} \times \frac{2T}{3}$: we factor in an updated value of the fatal failure frequency $\frac{1}{\mu}\frac{T}{\mu}$: the first failure strikes with frequency $\frac{1}{\mu}$, and then with frequency $\frac{T}{\mu}$, there is another failure before the end of the period. As for the time lost, it becomes $\frac{2T}{3}$, because in average the first error strikes at one third of the period and the second error strikes at two-third of the period: indeed, we know that there are two errors in the period, and they are equally spaced in average. Altogether, both overhead sources add up to

$$\frac{C}{T} + \frac{21}{3\mu^2},$$
 (18)

which is exactly Equation (15).

We conclude this section with a comment on the *no-restart* strategy. The intuitive derivation in Equation (12) leads to $\mathbb{H}^{no}(T) = \frac{C}{T} + \frac{T}{2M_{2b}}$. We now understand that this derivation is accurate if we have $T_{lost}(T) = \frac{T}{2} + o(T)$. While this latter equality is proven true without replication [13], it is unknown whether it still holds with replication. Hence, computing the optimal period for *no-restart* remains an open problem, even with a single processor pair.

Going further, Figure 2 shows that periodic checkpointing is not optimal for no-restart with a single processor pair, which provides another hint of the difficulty of the problem. In the figure, we compare four approaches: in addition to Restart (T_{opt}^{rs}) and NoRestart (T_{MTTI}^{no}) , we use two non-periodic variants of no-restart, Non-Periodic (T_1, T_2) . In both variants, we use a first checkpointing period T_1 while both processors are alive, and then a shorter period T_2 as soon as one processor has been struck by a failure. When an application failure occurs, we start anew with periods of length T_1 . For both variants, we only restart processors after an application failure, just as no-restart does. The first variant uses $T_1 = T_{MTTI}^{\text{no}} = \sqrt{3\mu C}$ (the MTTI is $M_2 = 3\frac{\mu}{2}$) and the second variant uses $T_1 = T_{opt}^{rs} = \left(\frac{3}{4}C\mu^2\right)^{\frac{1}{3}}$. We use the Young/Daly period $T_2 = \sqrt{2\mu C}$ for both variants, because there remains a single live processor when period T_2 is enforced. The figure shows the ratio of the time-to-solution for the two non-periodic approaches over that of periodic no-restart (with period T_{MTTI}^{no}). Note that the application is perfectly parallel, and that the only overhead is for checkpoints and re-executions after failures. Both non-periodic variants are better than no-restart, the first one is within 98.3% of no-restart, and the second one is even better (95% of no-restart) when the MTBF increases. We also see that restart is more than twice better than no-restart with a single processor pair. Note that results are averaged over 100,000 simulations, each lasting for 10,000 periods, so that they are statistically guaranteed to be accurate.

4.3 With b Processor Pairs

For b pairs, the reasoning is the same as with one pair, but the probability of having a fatal error (both processors of a same pair failing) before the end of the period changes. Letting $p_b(T)$ be the probability of failure before time T with b pairs, we have $p_b(T) = 1 - (1 - (1 - e^{-\lambda T})^2)^b$. As a consequence, computing the exact value of $T_{\text{lost}}(T)$ becomes complicated: obtaining a compact closed-form is not easy, because we would need to expand terms using the binomial formula. Instead, we directly use the Taylor expansion of $p_b(T)$ for λT close to 0. Again, this is valid only if $T = \Theta(\lambda^{-x})$ with x < 1. We have $p_b(T) = 1 - (1 - (\lambda T + o(\lambda T))^2)^b = b\lambda^2 T^2 + o(\lambda^2 T^2)$ and compute $T_{\text{lost}}(T)$ with b pairs as $T_{\text{lost}}(T) = \frac{1}{p_b(T)} \int_0^T t \frac{d\mathbb{P}(X \leq t)}{dt} dt = \frac{2T}{3} + o(T)$. As before, $T_{\text{lost}}(T) \sim \frac{2T}{3}$. Also, as in Section 4.2, we analyze the restart strategy,

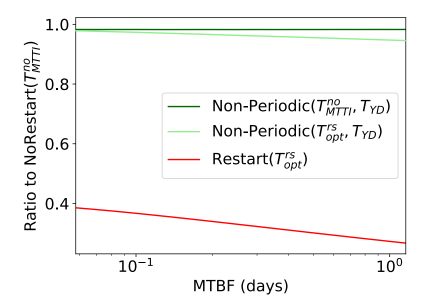


Figure 2: Ratio of time-to-solution of two non-periodic strategies and restart over time-to-solution of no-restart (one processor pair, $C = C^R = 60$).

which requires any failed processor to recover whenever a checkpoint is taken. We come back to the difference with the *no-restart* strategy after deriving the period for the *restart* strategy. We compute the expected execution time of one period: $\mathbb{E}(T) = p_b(T) \left(T_{\text{lost}}(T) + D + R + \mathbb{E}(T)\right) + (1 - p_b(T)) \left(T + C^R\right) = T + \frac{2b\lambda^2 T^3}{3} + o(\lambda^2 T^3)$, and

$$\mathbb{H}^{rs}(T) = \frac{\mathbb{E}(T)}{T} - 1 = \frac{C^R}{T} + \frac{2b\lambda^2 T^2}{3} + o(\lambda^2 T^2). \tag{19}$$

We finally derive the expression of the optimal checkpointing period with b pairs:

$$T_{opt}^{rs} = \left(\frac{3C^R}{4b\lambda^2}\right)^{\frac{1}{3}} = \Theta(\lambda^{-\frac{2}{3}}). \tag{20}$$

When plugging it back in Equation (19), we get

$$\mathbb{H}^{\text{rs}}(T_{opt}^{\text{rs}}) = \left(\frac{3C^R\sqrt{b}\lambda}{\sqrt{2}}\right)^{\frac{2}{3}} + o(\lambda^{\frac{2}{3}}) = \Theta(\lambda^{\frac{2}{3}}). \tag{21}$$

for the optimal overhead when using b pairs of processors.

The derivation is very similar to the case with a single pair, and the result is essentially the same, up to factoring in the number of pairs to account for a higher failure rate. However, the difference between the *no-restart* and the *restart* strategies gets more important. Indeed, with the *no-restart*

strategy, several pairs can be struck once (and even several times if the failures always strike the failed processor) before a pair finally gets both its processors killed. While the *no-restart* strategy spares the cost of several restarts, it runs at risk with periods whose length has been estimated à la Young/Daly, thereby assuming an identical setting at the beginning of each period.

Finally, for the intuitive way to retrieve Equation (20), it goes as for Equation (18), multiplying the frequency of fatal failures $\frac{1}{\mu}\frac{T}{\mu}$ by a factor b to account for each of the b pairs possibly experiencing a fatal failure.

5 Time-To-Solution

So far, we have focused on period length. In this section, we move to actual work achieved by the application. Following [25], we account for two sources of overhead for the application. First, the application is not perfectly parallel and obeys Amdahl's law [1], which limits its parallel speedup. Second, there is an intrinsic slowdown due to active replication related to duplicating every application message [20, 25].

First, for applications following Amdahl's law, the total time spent to compute W units of computation with N processors is $T_{Amdahl} = \gamma W + (1-\gamma)\frac{W}{N} = (\gamma + \frac{1-\gamma}{N})W$, where γ is the proportion of inherently sequential tasks. When replication is used, this time becomes $T_{Amdahl} = (\gamma + \frac{2(1-\gamma)}{N})W$. Following [25], we use $\gamma = 10^{-5}$ in Section 7. Second, as stated in [20, 25], another slowdown related to active replication and its incurred increase of communications writes $T_{rep} = (1+\alpha)T_{Amdahl}$, where α is some parameter depending upon the application and the replication library. Following [25], we use either $\alpha = 0$ or $\alpha = 0.2$ in Section 7.

All in all, once we have derived T_{opt} , the optimal period between two checkpoints without replication (see Equation (6)), and T_{opt}^{rs} , the optimal period between two checkpoints with replication and restart (see Equation (20)), we are able to compute the optimal number of operations to be executed by an application between two checkpoints as $W_{opt} = \frac{T_{opt}}{(\gamma + \frac{1-\gamma}{N})}$ for an application without replication, and $W_{opt}^{rs} = \frac{T_{opt}^{rs}}{(1+\alpha)\left(\gamma + \frac{1-\gamma}{b}\right)} = \frac{T_{opt}^{rs}}{(1+\alpha)\left(\gamma + \frac{2(1-\gamma)}{N}\right)}$ for an application with replication and the restart strategy. Finally, for the no-restart strategy, using T_{MTTI}^{no} (see Equation (11)), the number of operations becomes $W_{MTTI}^{no} = \frac{T_{MTTI}^{no}}{(1+\alpha)\left(\gamma + \frac{2(1-\gamma)}{N}\right)}$.

To compute the actual time-to-solution, assume that we have a total of W_{seq} operations to do. With one processor, the execution time is $T_{seq} = W_{seq}$ (assuming unit execution speed). With N processors working in parallel (no replication), the failure-free execution time is $T_{par} = (\gamma + \frac{1-\gamma}{N})T_{seq}$. Since we partition the execution into periods of length T, meaning that we have $\frac{T_{par}}{T}$

periods overall, the time-to-solution is $T_{final} = \frac{T_{par}}{T} \mathbb{E}(T) = T_{par}(\mathbb{H}(T) + 1)$, hence

$$T_{final} = \left(\gamma + \frac{1 - \gamma}{N}\right) (\mathbb{H}(T) + 1) T_{seq}. \tag{22}$$

If we use replication with b pairs of processors (i.e., $\frac{N}{2}$ pairs) instead, the difference is that $T_{par} = (1 + \alpha) \left(\gamma + \frac{1 - \gamma}{b} \right) T_{seq}$, hence

$$T_{final} = (1 + \alpha) \left(\gamma + \frac{2(1 - \gamma)}{N} \right) \left(\gamma + \frac{2(1 - \gamma)}{N} \right) (\mathbb{H}(T) + 1) T_{seq}. \tag{23}$$

Without replication, we use the optimal period $T = T_{opt}$. For the restart strategy, we use the optimal period $T = T_{opt}^{rs}$, and for no-restart, we use $T = T_{mTT}^{no}$, as stated above.

6 Asymptotic Behavior

In this section, we compare the *restart* and *no-restart* strategies asymptotically. Both approaches (and, as far as we know, all coordinated rollback-recovery approaches) are subject to a design constraint: if the time between two restarts becomes of same magnitude as the time to take a checkpoint, the application cannot progress. Therefore, when evaluating the asymptotic behavior (i.e., when the number of nodes tends to infinity, and hence the MTTI tends to 0), a first consideration is to state that none of these techniques can support infinite growth, under the assumption that the checkpoint time remains constant and that the MTTI decreases with scale. Still, in that case, because the *restart* approach has a much longer checkpointing period than *no-restart*, it will provide progress for lower MTTIs (and same checkpointing cost).

However, we can (optimistically) assume that checkpointing technology will evolve, and that rollback-recovery protocols will be allowed to scale infinitely, because the checkpoint time will remain a fraction of the MTTI. In that case, assume that with any number N of processors, we have $C = xM_N$ for some small constant x < 1 (where M_N is the MTTI with N processors). Consider a parallel and replicated application that would take a time T_{app} to complete without failures (and with no fault-tolerance overheads). We compute the ratio \mathcal{R} , which is the expected time-to-solution using the restart strategy divided by the expected time-to-solution using the no-restart strategy:

$$\mathcal{R} = \frac{(\mathbb{H}^{rs}(T_{opt}^{rs}) + 1)T_{app}}{(\mathbb{H}^{no}(T_{MTTI}^{no}) + 1)T_{app}} = \frac{\sqrt[3]{\frac{9}{8}\pi x^2} + 1}{\sqrt{2x} + 1}.$$

Because of the assumption $C = xM_N$, both the number of nodes N and the MTBF μ simplify out in the above ratio. Under this assumption, the restart

strategy is up to 8.4% faster than the *no-restart* strategy if x is within the range [0, 0.64], i.e., as long as the checkpoint time takes less than 2/3 of the MTTI.

In the next section, we consider realistic parameters to evaluate the performance of various strategies through simulations, and we also provide results when increasing the number of processors N or reducing the MTBF.

7 Experimental Evaluation

In this section, we evaluate the performance of the *no-restart* and *restart* strategies through simulations. Our simulator is publicly available [6] so that interested readers can instantiate their preferred scenarios and repeat the same simulations for reproducibility purpose. The code is written in-house in C++ and does not use any library other than the Standard Template Library (STL).

We compare different instances of the models presented above. We let Restart(T) denote the restart strategy with checkpointing period T, and NoRestart(T) denote the no-restart strategy with checkpointing period T. In most figures, we present the overhead as given by Equation (1): it is a relative time overhead, that represents the time spent tolerating failures divided by the duration of the protected application. Recall previously introduced notations:

- For Restart(T), the overhead $\mathbb{H}^{rs}(T)$ is predicted by the model according to Equation (19);
- For NoRestart(T), the overhead $\mathbb{H}^{no}(T)$ is estimated in the literature according to Equation (12);
- T_{opt}^{rs} denotes the optimal period for minimizing the time overhead for the restart strategy, as computed in Equation (20);
- T_{MTTI}^{no} from Equation (11) is the standard period used in the literature for the *no-restart* strategy, after an analogy with the Young/Daly formula.

The no-restart strategy with overhead $\mathbb{H}^{\text{no}}(T_{MTTI}^{\text{no}})$ represents the state of the art for full replication [20]. For completeness, we also compare the no-restart and restart strategies with several levels of partial replication [17, 25].

We describe the simulation setup in Section 7.1. We assess the accuracy of our model and of first-order approximations in Section 7.2. We compare the performance of restart with restart-on-failure in Section 7.3. In Section 7.4, we show the impact of key parameters on the difference between the checkpointing periods of the no-restart and restart strategies, and on the associated time overheads. Section 7.5 discusses the impact of the different strategies on I/O pressure. Section 7.6 investigates in which scenarios a smaller time-to-solution can be achieved with full or partial replication. Section 7.7 explores strategies that restart after a given number of failures.

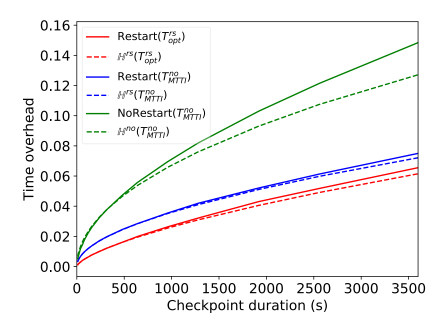


Figure 3: Evaluation of model accuracy for time overhead. $\mu = 5$ years, $b = 10^5$.

7.1 Simulation Setup

To evaluate the performance of the no-restart and restart strategies, we use a publicly available simulator [6] that generates random failures following an exponential probability distribution with a given mean time between individual node failures and number of processor pairs. Then, we set the checkpointing period, and checkpointing cost. Default values are chosen to correspond to the values used in [25], and are defined as follows. For the checkpointing cost, we consider two default values: C = 60 seconds corresponds to buddy checkpointing, and C = 600 seconds corresponds to checkpointing on remote storage. We let the MTBF of an individual node be $\mu = 5$ years, and we use N = 200,000, hence having b = 100,000 pairs when replication is used. We then simulate the execution of an application lasting for 100 periods (total execution time 100T) and we average the results on 1000 runs. We measure two main quantities: time overhead and optimal period length. For simplicity, we always assume that R=C, i.e., read and write operations take (approximately) the same time. We cover the whole range of possible values for C^R , using either C, 1.5C or 2C. This will show the impact of overlapping checkpoint and processor restart.

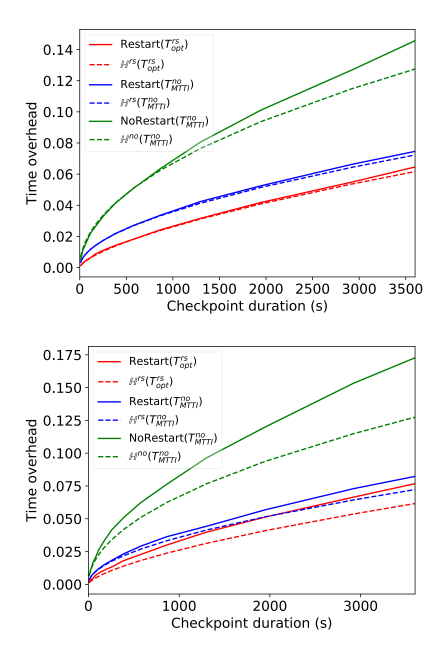


Figure 4: Evaluation of model accuracy for time overhead with two trace logs (LANL#18 on the left, and LANL#2 on the right).

7.2 Model Accuracy

Figure 3 compares three different ways of estimating the time overhead of an application running on $b=10^5$ processor pairs. Solid lines are measurements from the simulations, while dashed lines are theoretical values. The red color is for $Restart(T_{opt}^{rs})$, the blue color is for $Restart(T_{MTTI}^{no})$ and the green color is for $NoRestart(T_{MTTI}^{no})$. For the restart strategy, $C^R = C$ in this figure.

For the restart strategy, the results from simulation match the results from the theory quite accurately. Because our formula is an approximation valid when $T \gg C$, the difference between simulated time overhead and $\mathbb{H}^{\rm rs}(T^{\rm rs}_{opt})$ slightly increases when the checkpointing cost becomes greater than 1500 seconds. We also verify that $Restart(T^{\rm rs}_{opt})$ has smaller overhead than $Restart(T^{\rm no}_{MTTI})$ in the simulations, which nicely corroborates the model.

We also see that $\mathbb{H}^{\text{no}}(T^{\text{no}}_{MTTI})$ is a good estimate of the actual simulated overhead of $NoRestart(T^{\text{no}}_{MTTI})$ only for C < 500. Larger values of C induce a significant deviation between the prediction and the simulation. Values given by $\mathbb{H}^{\text{no}}(T)$ underestimate the overheads for lower values of C more than $\mathbb{H}^{\text{rs}}(T)$, even when using the same T^{no}_{MTTI} period to checkpoint. As described at the end of Section 4.1, the $\mathbb{H}^{\text{no}}(T)$ formula is an approximation whose accuracy is unknown, and when C scales up, some elements that were neglected by the approximation become significant. The formula for T^{rs}_{opt} , on the contrary, remains accurate for higher values of C.

Figure 4 is the exact counterpart of Figure 3 when using log traces from real platforms instead of randomly generated failures with an exponential distribution. We use the two traces featuring the largest number of failures from the LANL archive [27, 26], namely LANL#2 and LANL#18. According to the detailed study in [2], failures in LANL#18 are not correlated while those in LANL#2 are correlated, providing perfect candidates to experimentally study the impact of failure distributions. LANL#2 has an MTBF of 14.1 hours and is composed of 5350 failures, while LANL#18 has an MTBF of 7.5 hours and is composed of 3899 failures. For the sake of comparing with Figure 3 that used a processor MTBF of 5 years (and an exponential distribution), we scale both traces as follows:

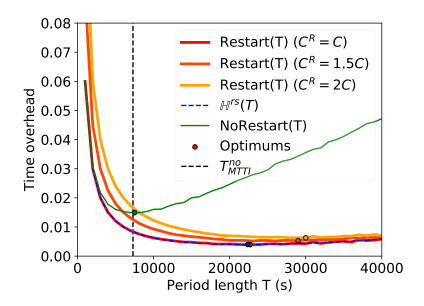
- We target a platform of 200,000 processors with an individual MTBF of 5 years. Thus the global platform MTBF needs to be 64 times smaller than the MTBF of LANL#2, and 32 times smaller than the MTBF of LANL#18. Hence we partition the global platform into 64 groups (of 3,125 processors) for LANL#2, and into 32 groups (of 6,250 processors) for LANL#18;
- Within each group, the trace is rotated around a randomly chosen date, so that each trace starts independently;
- We generate 200 sets of failures for each experiment and report the average time overhead.

We observe similar results in Figure 3 and Figure 4. For LANL#18,

the experimental results are quite close to the model. For LANL#2, the model is slightly less accurate because of some severely degraded intervals with failure cascades. However, the restart strategy still grants lower time overheads than the no-restart strategy. For an exponential distribution, only 15% of the runs where an application failure was experienced did experience two or more failures. This ratio increases to 20% for LANL#18 and reaches 50% for LANL#2; this leads to a higher overhead than estimated for IID failures, but this is true for all strategies, and restart remains the best one.

Next, on both graphs in Figure 5, we present the details of the evolution of the time overhead as a function of the period length for C=60s and C=600s. Here, we compare the overhead of the restart strategy obtained through simulations (solid red, orange and yellow lines for different values of C^R), the overhead of the restart strategy obtained through the theoretical model with $C^R=C$ (dashed blue line), and the overhead of the no-restart strategy obtained through simulations (solid green line). In each case, a circle denotes the optimal period, while $T_{MTTI}^{\rm no}$ (the MTTI extension of the Young/Daly formula for no-restart) is shown with a vertical bar.

 $\mathbb{H}^{rs}(T_{opt}^{rs})$ perfectly matches the behavior of the simulations, and the optimal value is very close to the one found through simulations. The simulated overhead of NoRestart(T) is always larger than for Restart(T), with a significant difference as T increases. Surprisingly, the optimal value for the simulated overhead of NoRestart(T) is obtained for a value of T close to T_{MTTI}^{no} , which shows a posteriori that the approximation worked out pretty well in this scenario. The figure also shows that the restart strategy is much more robust than the *no-restart* one: in all cases, Restart(T) provides a lower overhead than NoRestart(T) throughout the spectrum, even when $C^R = 2C$. More importantly, this overhead remains close to the minimum for a large range of values of T: when $C^R = C = 60s$, for values of T between 21,000s and 25,000s, the overhead remains between 0.39% (the optimal), and 0.41%. If we take the same tolerance (overhead increased by 5%), the checkpointing period must be between 6,000s and 9,000s, thus a range that is 1/3rd larger than for the restart strategy. When considering $C^R = C = 600s$, this range is 18,000s (40,000s to 58,000s) for the restart strategy, and 7,000s (22,000s to 29,000s) for the no-restart one. This means that a user has a much higher chance of obtaining close-to-optimum performance by using the restart strategy than if she was relying on the no-restart one, even if some key parameters that are used to derive T_{opt}^{rs} are mis-evaluated. If $C^R = 1.5C$ or $C^R = 2C$, the same trends are observed: the optimal values are obtained for longer periods, but they remain similar in all cases, and significantly lower than for the *no-restart* strategy. Moreover, the figures show the same plateau effect around the optimal, which makes the restart strategy robust.



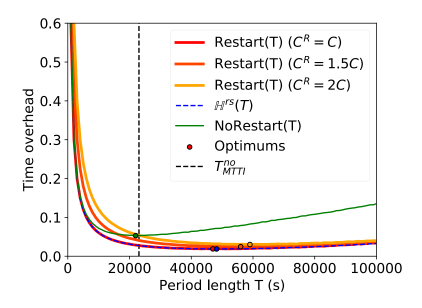


Figure 5: Time overhead as a function of the checkpointing period T for C=60 seconds (left) or C=600 seconds (right), MTBF of 5 years, IID failures and $b=10^5$ processor pairs.

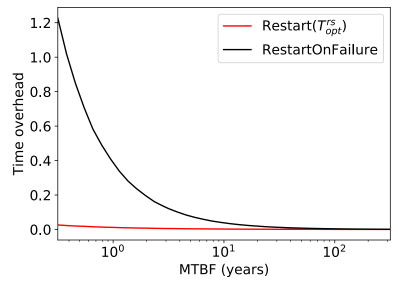


Figure 6: Comparison with restart-on-failure.

7.3 Restart-on-failure

Figures 3 to 5 showed that the restart strategy is more efficient than the no-restart one. Intuitively, this is due to the rejuvenation introduced by the periodical restarts: when reaching the end of a period, failed processes are restarted, even if the application could continue progressing in a more risky configuration. A natural extension would be to consider the restart-on-failure strategy described in Section 1. This is the scenario evaluated in Figure 6: we compare the time overhead of $Restart(T_{opt}^{rs})$ with restart-on-failure, which restarts each processor after each failure.

Compared to $Restart(T_{MTTI}^{no})$, the restart-on-failure strategy grants a significantly higher overhead that quickly grows to high values as the MTBF decreases. The restart-on-failure strategy works as designed: no rollback was ever needed, for any of the simulations (i.e., failures never hit a pair of replicated processors within the time needed to checkpoint). However, the time spent checkpointing after each failure quickly dominates the execution. This reflects the issue with this strategy, and the benefit of combined replication and checkpointing: as failures hit the system, it is necessary for performance to let processors fail and the system absorb most of the failures using the replicates. Combining this result with Figure 5, we see that it is critical for performance to find the optimal rejuvenation period: restarting failed processes too frequently is detrimental to performance, as is restarting them too infrequently.

7.4 Impact of Parameters

The graphs in Figure 7 describe the impact of the individual MTBF of the processors on the time overhead. We compare $Restart(T_{opt}^{rs})$, $Restart(T_{MTTI}^{no})$ (both in the most optimistic case when $C^R = C$ and in the least optimistic case when $C^R = 2C$) and $NoRestart(T_{MTTI}^{no})$. As expected, when C^R increases, the time overhead increases. However, even in the case $C^R = 2C$, both restart strategies outperform the no-restart strategy. As the MTBF increases, the overhead of all strategies tends to be negligible, since a long MTBF has the cumulated effect that the checkpointing period increases and the risk of needing to re-execute decreases. The longer the checkpoint time C, the higher the overheads, which is to be expected; more interestingly, with higher C, the restart strategy needs C^R to remain close to C to keep its advantage against the no-restart strategy. This advocates for a buddy checkpointing approach with restart strategy when considering replication and checkpointing over unreliable platforms.

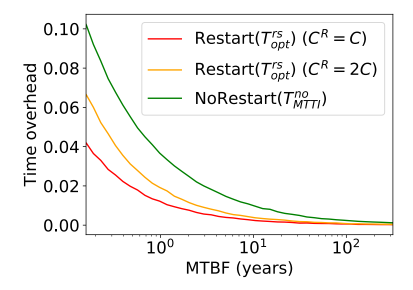
7.5 I/O Pressure

Figure 8 reports the difference between $T_{opt}^{\rm rs}$ and $T_{MTTI}^{\rm no}$. We see that $T_{opt}^{\rm rs}$ increases faster than $T_{MTTI}^{\rm no}$ when the MTBF decreases. This is due to the fact that the processors are restarted at each checkpoint, hence reducing the probability of failure for each period; it mainly means that using the restart strategy (i) decreases the total application time, and (ii) decreases the I/O congestion in the machine, since checkpoints are less frequent. This second property is critical for machines where a large number of applications are running concurrently, and for which, with high probability, the checkpoint times are longer than expected because of I/O congestion.

7.6 Time-To-Solution

Looking at the time overhead is not sufficient to evaluate the efficiency of replication. So far, we only compared different strategies that all use full process replication. We now compare the restart and no-restart strategies to the approach without replication, and also to the approach with partial replication [17, 25]. Figure 9 shows the corresponding time-to-solution for $\gamma = 10^{-5}$ and $\alpha = 0.2$ (values used in [25]), and $C^R = C$ when the individual MTBF varies. Recall that the time-to-solution is computed using Equation (22) without replication (where $\mathbb{H}(T)$ is given by Equation (7)), and using Equation (23) with replication (where $\mathbb{H}(T)$ is given by Equation (12) for no-restart, and by Equation (19) for restart). In the simulations, T_{seq} is set so that the application lasts one week with 100,000 processors (and no replication).

In addition to the previously introduced approaches, we evaluate $Partial90(T_{opt}^{rs})$ and $Partial50(T_{MTTI}^{no})$. Partial90 represents a partial replication approach



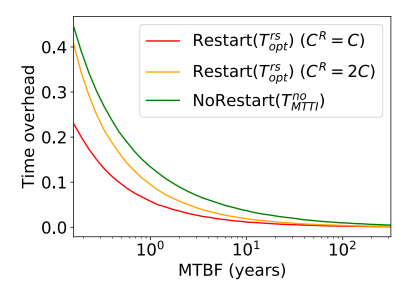
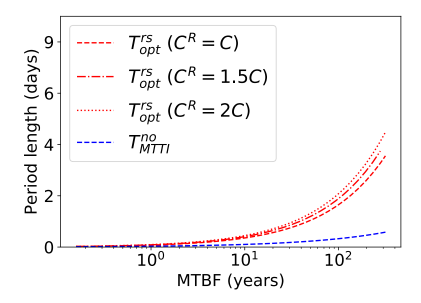


Figure 7: Time overhead as a function of MTBF, with $C=60\mathrm{s}$ (left) or $C=600\mathrm{s}$ (right), $b=10^5$ processor pairs.



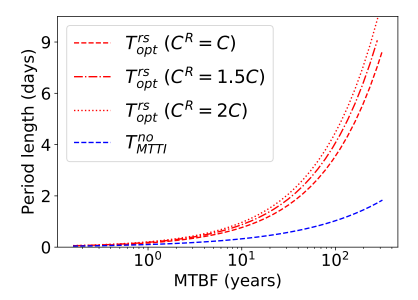


Figure 8: Period length T as function of MTBF, with $C=60\mathrm{s}$ (left) or $C=600\mathrm{s}$ (right), $b=10^5$ processor pairs.

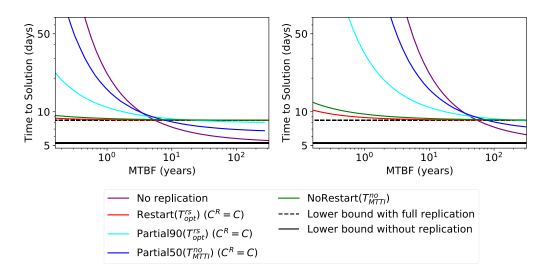


Figure 9: Time-to-solution for $N=2\times 10^5$ standalone proc. against full and partial replication approaches, as a function of MTBF, with $C^R=C=60$ s (left) or $C^R=C=60$ s (right), $\gamma=10^{-5}$, $\alpha=0.2$.

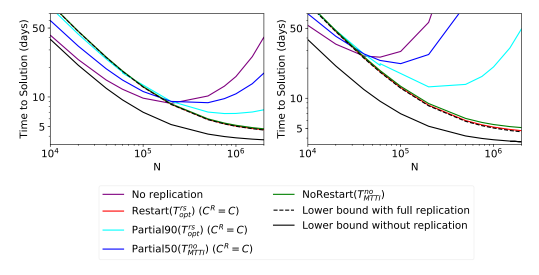


Figure 10: Time-to-solution with MTBF of 5 years against full and partial replication approaches, as a function of N, with $C^R = C = 60$ s (left) or $C^R = C = 60$ os (right), $\gamma = 10^{-5}$, $\alpha = 0.2$.

where 90% of the platform is replicated (there are 90,000 processor pairs and 20,000 standalone processors). Similarly, 50% of the platform is replicated for Partial 50 (there are 50,000 processor pairs and 100,000 standalone processors). Figure 9 illustrates the benefit of full replication: when the MTBF becomes too short, replication becomes mandatory. Indeed, in some cases, simulations without replication or with partial replication would not complete, because one fault was (almost) always striking before a checkpoint, preventing progress. For C = 60s and $N = 2 \times 10^5$, $\gamma = 10^{-5}$ and $\alpha = 0.2$, full replication grants the best time-to-solution for an MTBF shorter than 1.8×10^8 . However, when the checkpointing cost increases, this value climbs up to 1.9×10^9 , i.e., roughly 10 times higher than with 60 seconds. As stated before, T_{opt}^{rs} gives a better overhead, thus a better execution time than T_{MTTI}^{no} . If machines become more unreliable, the restart strategy allows us to maintain the best execution time. Different values of γ and α give the same trend as in our example, with large values of γ making replication more efficient, while large values of α reduce the performance. Similarly to what was observed in [25], for a homogeneous platform (i.e., if all processors have a similar risk of failure), partial replication (at 50% or 90%) exhibits lower performance than no replication for long MTBF, and lower performance than the no-restart strategy (hence even lower performance than the restart strategy) for short MTBF. This confirms that partial replication has potential benefit only for heterogeneous platforms, which is outside the scope of this study.

We now further focus on discussing when replication should be used. Figure 10 shows the execution time of an application when the number of processors N varies. Each processor has an individual MTBF of 5 years. The same general comments can be made: $\mathit{Restart}(T^{\mathrm{rs}}_{opt})$ always grants a slightly lower time-to-solution than $NoRestart(T_{MTTI}^{no})$, because it has a smaller overhead. As before, when N is large, the platform is less reliable and the difference between $Restart(T_{opt}^{rs})$ and $NoRestart(T_{MTTI}^{no})$ is higher compared to small values of N. We see that replication becomes mandatory for large platforms: without replication, or even with 50% of the platform replicated, the time-to-solution is more than 10 times higher than the execution time without failures. With $\gamma = 10^{-5}$ and $\alpha = 0.2$, replication becomes more efficient than no replication for $N \ge 2 \times 10^5$ processors when C = 60s. However, when C = 600s, it starts being more efficient when $N > 2.5 \times 10^4$, i.e., roughly 10 times less processors when C is 10 times longer. This study further confirms that partial replication never proved to be useful throughout our experiments.

7.7 When to Restart

In this section, we consider a natural extension of the *restart* approach: instead of restarting failed processors at each checkpoint, the restart can be

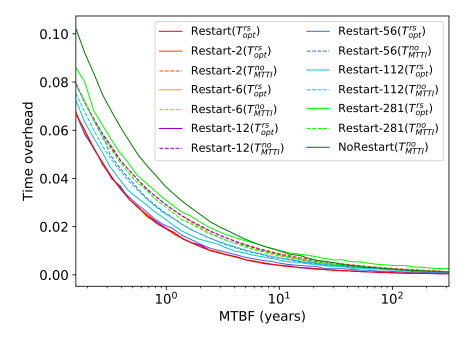


Figure 11: Comparison of restart strategy with restart only every 2, 6, 12, 56, 112, or 281 dead processors, with T_{opt}^{rs} and T_{MTTI}^{no} .

delayed until the next checkpoint where the number of accumulated failures reaches or exceeds a given bound $n_{\rm bound}$, thereby reducing the frequency of the restarts.

The restart strategy assumes that after a checkpoint, the risk of any processor failing is the same as in the initial configuration. For the extension, there is no guarantee that $T_{opt}^{\rm rs}$ remains the optimal interval between checkpoints; worse, there is no guarantee that periodic checkpointing remains optimal. To evaluate the potential gain of reducing the restart frequency, we consider the two proposed intervals: $T_{opt}^{\rm rs}$ and $T_{MTTT}^{\rm no}$. And, since most checkpoints will not incur a restart, we assume $C^R = C$ when computing $T_{opt}^{\rm rs}$. However, checkpoints where processes are restarted have a cost of twice the cost of a simple checkpoint in the simulation: this is the worst case for the restart strategy. We then simulate the execution, including restarts due to reaching $n_{\rm bound}$ failures and due to application crashes. With b=100,000 processor pairs, we expect $n_{\rm fail}(2b)=561$ failures before the application is interrupted; so we will consider a large range of values for $n_{\rm bound}$: from 2, 6, 12, to cover cases where few failures are left to accumulate, to 56, 112, or 281, that represent respectively 10%, 20% and 50% of $n_{\rm fail}(2b)$, to cover cases where many failures can accumulate.

The results are presented in Figure 11, for a variable node MTBF. The time overhead of the extended versions is higher than the time overhead of

the restart approach using T_{opt}^{rs} as a checkpointing (and restarting) interval. The latter is also lower than the overhead of the no-restart strategy, which on average corresponds to restarting after $n_{\text{bound}} = n_{\text{fail}}(2b) = 561$ failures. This shows that restarting the processes after each checkpoint consistently decreases the time overhead. Using the optimal checkpointing period for restart T_{opt}^{rs} , increasing n_{bound} also increases the overhead. Moreover, when using small values (such as 2 and 6) for n_{bound} , we obtain exactly the same results as for the restart strategy. This is due to the fact that between two checkpoints, the restart strategy usually looses around 6 processors, meaning that restart is already the same strategy as accumulating errors up to 6 (or less) before restarting. With $n_{\text{bound}} = 12$, on average the restart happens every two checkpoints, and the performance is close, but slightly slower than the restart strategy.

Finally, an open problem is to determine the optimal checkpointing strategy for the extension of restart tolerating n_{bound} failures before restarting failed processors. This optimal strategy could render the extension more efficient than the baseline restart strategy. Given the results of the simulations, we conjecture this optimal number to be 0, i.e., restart would be the optimal strategy.

Summary. Overall, we have shown that the restart strategy with period T_{opt}^{rs} is indeed optimal and that our model is realistic. We showed that restart decreases time overhead, hence time-to-solution, compared to using no-restart with period T_{MTTI}^{no} . The extended version [6] shows similar gains in energy overheads. The main decision is still to decide whether the application should be replicated or not. However, whenever it should be (which is favored by a large ratio of sequential tasks γ , a large checkpointing cost C, or a short MTBF), we are now able to determine the best strategy: use full replication, restart dead processors at each checkpoint (overlapped if possible), and use T_{opt}^{rs} for the checkpointing period.

8 Related work

Checkpoint-restart is one of the most used strategy to deal with fail-stop errors, and several variants of this policy have been studied, see [24] for a survey. The natural strategy is to checkpoint periodically, and one must then decide how often to checkpoint, hence derive the optimal checkpointing period. For a divisible application, results were first obtained by Young [42] and Daly [13]. This strategy has been extended to deal with a multi-level checkpointing scheme [29, 14, 5], or by using SSD or NVRAM as secondary storage [9].

If the error rate and/or checkpoint cost is too important, and hence the overhead induced by the checkpointing strategy is large, checkpointing can be combined with replication. Hence, some redundant MPI processes are used to execute a replica of the work [19, 20, 11]. For instance, Ferreira et al. [20] used two replicas per MPI process, and they provided a theoretical analysis of parallel efficiency, an MPI implementation that supports transparent process replication (including failure detection, consistent message ordering among replicas, etc.), and a set of experimental and simulation results. Hence, they demonstrate that replication outperforms traditional checkpoint/restart approach in several scenarios.

Partial redundancy is studied in [17, 36, 37] (in combination with coordinated checkpointing) to decrease the overhead of full replication. Recently, Hussain et al. [25] have demonstrated the usefulness of partial redundancy for platforms where individual node failure distributions are not identical. They numerically determine the optimal partial replication degree.

For malleable applications, adaptive redundancy is discussed in [22], where a subset of processes is dynamically selected for replication. Furthermore, the number of processors on which the applications execute is changed at runtime, yielding significant improvement in application performance.

Finally, in contrast to fail-stop errors whose detection is immediate, silent errors are identified only when the corrupted data leads to an unusual application behavior, and several works use replication to detect and/or correct silent errors. For instance, thread-level replication has been investigated in [43, 12, 33], which target process-level replication in order to detect (and correct) silent errors striking in all communication-related operations. Also, Ni et al [30] introduce process duplication to cope with both fail-stop and silent errors. Recently, Benoit et al. [4] extended these work to general applications, and compare traditional process replication with group replication, where the whole application is replicated as a black box. They analyze several scenarios with duplication or triplication.

To the best of our knowledge, all related works use the *no-restart* strategy described in the paper: in a replicated execution, failed processes are not restarted until the application experiences a fatal failure.

9 Conclusion

In this work, we have revisited process replication combined with check-pointing, an approach that has received considerable attention from the HPC community in recent years. Opinion is divided about replication. By definition, its main drawback is that 50% of platform resources will not contribute to execution progress, and such a reduced throughput does not seem acceptable in many scenarios. However, checkpoint/restart alone cannot ensure full reliability in heavily failure-prone environments, and must be complemented by replication in such unreliable environments. Previous approaches all used the *no-restart* strategy. In this work, we have introduced a new rollback/recovery strategy, the *restart* strategy, which consists

of restarting all failed processes at the beginning of each period. Thanks to this rejuvenation, the system remains in the same conditions at the beginning of each checkpointing period, which allowed us to build an accurate performance model and to derive the optimal checkpointing period for this strategy. This period turns out to be much longer than the one used with the no-restart strategy, hence reducing significantly the I/O pressure introduced by checkpoints, and improving the overall time-to-solution. To validate this approach, we have simulated the behavior of realistic large-scale systems, with failures either IID or from log traces. We have compared the performance of restart with the state-of-the-art strategies. Another key advantage of the restart strategy is its robustness: the range of periods in which its performance is close to optimal is much larger than for the no-restart strategy, making it a better practical choice to target unreliable platforms where the key elements (MTBF and checkpoint duration) are hard to estimate. In the future, we plan to evaluate, at least experimentally, non-periodic checkpointing strategies that rejuvenate failed processors after a given number of failures is reached or after a given time interval is exceeded.

Acknowledgement

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A Appendix: Energy consumption

In this appendix, we extend the approach to a different objective function: the goal is now to minimize the energy overhead. If $\mathcal{E}(T)$ is the expected energy consumption of a period of length $\mathcal{P} = T + C$, the energy overhead with a single processor is expressed as:

$$\mathbb{H}^{\text{energy}}(T) = \frac{\mathcal{E}(T)}{T(P_{\text{comp}} + P_{\text{static}})} - 1, \tag{24}$$

where P_{comp} is the dynamic power consumption of a processor when computing, and P_{static} denotes the static power, which is paid when the processor is kept idle, but still turned on.

We also denote by $P_{\text{I/O}}$ the dynamic power when performing I/O operations, which has to be accounted for when checkpointing, hence in the expression of $\mathcal{E}(T)$. We express below $\mathcal{E}(T)$ and $\mathbb{H}^{\text{energy}}(T)$ in the cases without replication (single processor or N processors, Section A.1) and with replication (one pair or b pairs, Section A.2), and derive in each case the optimal period, and the optimal energy overhead. Finally, we present comprehensive simulation results in Section A.3.

A.1 Without replication

A.1.1 With a single processor

In this case, we use the same approach as in Section 3.1 and we write a recursive formula similar to Equation (2):

$$\mathcal{E}(T) = (1 - F(T))(T(P_{\text{comp}} + P_{\text{static}}) + C(P_{\text{I/O}} + P_{\text{static}})) + F(T)(T_{\text{lost}}(T)(P_{\text{comp}} + P_{\text{static}}) + DP_{\text{static}} + R(P_{\text{I/O}} + P_{\text{static}}) + \mathcal{E}(T))$$
(25)

If the computation is successful, then we compute at power $P_{\text{comp}} + P_{\text{static}}$ during a time T and use the power $P_{\text{I/O}} + P_{\text{static}}$ during the checkpoint. However if a failure strikes, the machine is used at power $P_{\text{comp}} + P_{\text{static}}$ for $T_{\text{lost}}(T)$ seconds, then used at power P_{static} for the downtime, and used at power $P_{\text{I/O}} + P_{\text{static}}$ during the recovery before starting the period anew. Finally, we obtain:

$$\begin{split} \mathcal{E}(T) &= \left(C + (e^{\lambda T} - 1)R\right) \left(P_{\text{I/O}} + P_{\text{static}}\right) \\ &+ (e^{\lambda T} - 1)DP_{\text{static}} + \frac{e^{\lambda T} - 1}{\lambda} (P_{\text{comp}} + P_{\text{static}}). \end{split}$$

Using the Taylor expansion as previously, we obtain the overhead

$$\mathbb{H}^{\text{energy}}(T) = \frac{C(P_{\text{I/O}} + P_{\text{static}})}{T(P_{\text{comp}} + P_{\text{static}})} + \frac{\lambda T}{2} + o(\lambda T).$$
 (26)

Again, this overhead is minimized for $T = \Theta(\lambda^{-\frac{1}{2}})$. By differentiating Equation (26), we get the optimal period minimizing energy consumption:

$$T_{opt}^{\text{energy}} = \sqrt{\frac{2C(P_{\text{I/O}} + P_{\text{static}})}{\lambda(P_{\text{comp}} + P_{\text{static}})}} = \Theta(\lambda^{-\frac{1}{2}}).$$
 (27)

Plugging it back into Equation (26), we get the optimal energy overhead:

$$\mathbb{H}_{opt}^{\text{energy}} = \sqrt{\frac{2C\lambda(P_{\text{I/O}} + P_{\text{static}})}{P_{\text{comp}} + P_{\text{static}}}} + o(\lambda^{\frac{1}{2}}) = \Theta(\lambda^{\frac{1}{2}}). \tag{28}$$

A.1.2 With N processors

We can generalize the previous result for the case with N processors, as done in Section 3.2 for the time overhead. We obtain another similar formula:

$$T_{opt}^{\text{energy}} = \sqrt{\frac{2C(P_{\text{I/O}}^{(N)} + NP_{\text{static}})}{N^2\lambda(P_{\text{comp}} + P_{\text{static}})}} = \Theta(\lambda^{-\frac{1}{2}})$$
 (29)

for the optimal checkpointing period, while the overhead becomes:

$$\mathbb{H}_{opt}^{\text{energy}} = \sqrt{\frac{2C\lambda(P_{\text{I/O}}^{(N)} + NP_{\text{static}})}{P_{\text{comp}} + P_{\text{static}}}} + o(\lambda^{\frac{1}{2}}) = \Theta(\lambda^{\frac{1}{2}}). \tag{30}$$

The main difference between Equations (27), (28) and Equations (29), (30) is that the dynamic power and the static power is multiplied by N, the number of processors, as more processors consume more energy. Similarly, $P_{\rm I/O}$ becomes $P_{\rm I/O}^{(N)} = P_{\rm I/O,static} + NP_{\rm I/O,comm}$ to take into account that more nodes are sending data to the external storage.

A.2 With replication

A.2.1 With one processor pair

We now compute the expected energy consumption $\mathcal{E}(T)$ of a period of length $\mathcal{P} = T + C^R$. We use the same approach as in Section A.1.1 and aim at minimizing the energy overhead

$$\mathbb{H}^{\text{energy}}(T) = \frac{\mathcal{E}(T)}{2T(P_{\text{comp}} + P_{\text{static}})} - 1 \tag{31}$$

We write a recursive formula similar to Equation (13):

$$\mathcal{E}(T) = (1 - p_1(T)) \Big(T(2P_{\text{comp}} + 2P_{\text{static}}) + C^R(P_{\text{I/O}}^{(2)} + 2P_{\text{static}}) \Big)$$

$$+ p_1(T) \Big(T_{\text{lost}}(T) (2P_{\text{comp}} + 2P_{\text{static}}) + 2DP_{\text{static}} + R(P_{\text{I/O}}^{(2)} + 2P_{\text{static}}) + \mathcal{E}(T) \Big).$$
(32)

After solving, we obtain:

$$\begin{split} \mathcal{E}(T) &= T(2P_{\text{comp}} + 2P_{\text{static}}) + C^R(P_{\text{I/O}}^{(2)} + 2P_{\text{static}}) \\ &+ \frac{(e^{\lambda T} - 1)^2}{2e^{\lambda T} - 1} \Big(\frac{(2e^{-2\lambda T} - 4e^{-\lambda T})\lambda T + e^{-2\lambda T} - 4e^{-\lambda T} + 3}{2\lambda(1 - e^{-\lambda T})^2} (2P_{\text{comp}} + 2P_{\text{static}}) \\ &+ 2DP_{\text{static}} + R(P_{\text{I/O}}^{(2)} + 2P_{\text{static}}) \Big). \end{split}$$

After Taylor expansion, we derive the overhead $\mathbb{H}^{\text{energy}}(T)$:

$$\mathbb{H}^{\text{energy}}(T) = \frac{C^R(P_{\text{I/O}}^{(2)} + 2P_{\text{static}})}{2T(P_{\text{comp}} + P_{\text{static}})} + \frac{2\lambda^2 T^2}{3} + o(\lambda^2 T^2)$$
(33)

The computations are similar to Section 4.2 and we find the following optimal values for T_{opt}^{energy} and $\mathbb{H}_{opt}^{\text{energy}}$:

$$T_{opt}^{\text{energy}} = \left(\frac{3C^R(P_{\text{I/O}}^{(2)} + 2P_{\text{static}})}{8\lambda^2(P_{\text{comp}} + P_{\text{static}})}\right)^{\frac{1}{3}} = \Theta(\lambda^{\frac{-2}{3}}). \tag{34}$$

$$\mathbb{H}_{opt}^{\text{energy}} = \left(\frac{3C^R \lambda (P_{\text{I/O}}^{(2)} + 2P_{\text{static}})}{2\sqrt{2}(P_{\text{comp}} + P_{\text{static}})}\right)^{\frac{2}{3}} + o(\lambda^{\frac{2}{3}}) = \Theta(\lambda^{\frac{2}{3}}). \tag{35}$$

A.2.2 With b processor pairs

As previously, we compute the energy consumption for the execution of one period of duration $\mathcal{P} = T + C^R$ using the following recursion:

$$\mathcal{E}(T) = p_b(T) \left(T_{\text{lost}}(T) (2bP_{\text{comp}} + 2bP_{\text{static}}) + D2bP_{\text{static}} \right)$$

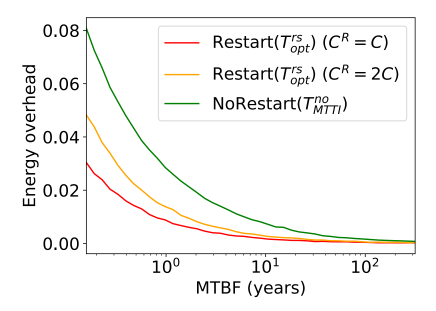
$$+ R(P_{\text{I/O}}^{(N)} + 2bP_{\text{static}}) + \mathcal{E}(T) \right)$$

$$+ (1 - p_b(T)) \left(T(2bP_{\text{comp}} + 2bP_{\text{static}}) + C^R(P_{\text{I/O}}^{(N)} + 2bP_{\text{static}}) \right).$$

With probability $p_b(T)$, the application fails so we account for the energy consumed until the failure $T_{\rm lost}(T)(2bP_{\rm comp}+2bP_{\rm static})$, followed by a downtime and a restart (power consumption of $P_{\rm I/O}^{(N)}+2bP_{\rm static}$). Otherwise, the application is successful, meaning that we computed at power $2b(P_{\rm comp}+P_{\rm static})$ during T seconds and we stored a checkpoint (overlapped with a restart) at power $P_{\rm I/O}^{(N)}+2bP_{\rm static}$. We already computed $T_{\rm lost}(T)$ in the previous subsection so we can directly derive, using a Taylor expansion of the exponential function and solving the previous equation that:

$$\mathbb{H}^{\text{energy}}(T) = \frac{\mathcal{E}(T)}{T \cdot 2b(P_{\text{comp}} + P_{\text{static}})} - 1$$

$$= \frac{C^R(P_{\text{I/O}}^{(N)} + 2bP_{\text{static}})}{2bT(P_{\text{comp}} + P_{\text{static}})} + \frac{2b\lambda^2 T^2}{3} + o(\lambda^2 T^2),$$
(36)



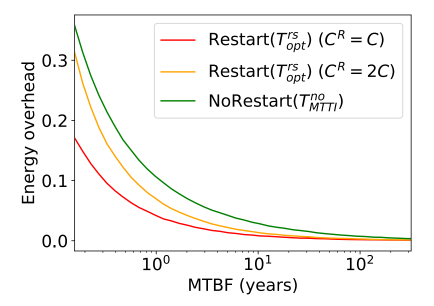


Figure 12: Influence of the MTBF on the energy overhead for the *restart* and *no-restart* strategies. Checkpointing time set to 60 seconds (left) or 600 seconds (right), with 10^5 pairs of processors.

which is very similar to Equation (33), with the only difference being a factor b on the second term and power consumption factors. We then derive a similar optimal period time T_{opt}^{energy} as well as the optimal energy overhead $\mathbb{H}_{ont}^{\text{energy}}$:

$$T_{opt}^{\text{energy}} = \left(\frac{3C^R(P_{\text{I/O}}^{(N)} + 2bP_{\text{static}})}{8b^2\lambda^2(P_{\text{comp}} + P_{\text{static}})}\right)^{\frac{1}{3}} = \Theta(\lambda^{\frac{-2}{3}}).$$
(37)

$$\mathbb{H}_{opt}^{\text{energy}} = \left(\frac{3C^R \lambda (P_{\text{I/O}}^{(N)} + 2bP_{\text{static}})}{2\sqrt{2b}(P_{\text{comp}} + P_{\text{static}})}\right)^{\frac{2}{3}} + o(\lambda^{\frac{2}{3}}) = \Theta(\lambda^{\frac{2}{3}}). \tag{38}$$

A.3 Experiments

For the power consumption, we chose $P_{\rm static}=10{\rm W/node}$ and $P_{\rm comp}=P_{\rm static}$, so that the non-idle power consumption of a node is 20W (i.e., an exascale machine with 10^6 nodes would reach the proposed bound of 20MW). For $P_{\rm I/O}$, as measured in [15], we set it to 15% of the static power, i.e., $P_{\rm I/O}=0.15P_{\rm static}=1.5{\rm W/node}$. With these values, we have $\frac{P_{\rm I/O}+P_{\rm static}}{P_{\rm comp}+P_{\rm static}}=0.575$, meaning that optimizing energy overhead will result in a shorter period than when optimizing time overhead.

Graphs in Figure 12 describe the impact of the individual MTBF of the processors on the energy overhead: they are the counterpart of Figure 7 that focused on execution time. The energy overheads reduce by a factor ranging from 62% to 80%, with the average being 72%.

Figure 13 shows the difference between the two optimal periods $T_{opt}^{\rm rs}$ and $T_{opt,en}^{\rm rs}$. As we can see, optimizing the time overhead or the energy overhead has a negligible impact on their values. When we optimize the energy overhead, our worst increase for the time overhead is around 15% for a MTBF ranging from 5×10^6 to 10^10 , $C^R = C = 60$ seconds and $b = 10^5$. The average increase however is of 3.1% over the whole range. When optimizing the time overhead, we measured a maximum of 23% improvement under the same conditions with the average increase being 4.2%. Overall, with our values we do not need to specifically optimize the energy overhead, except if the ratio between $P_{\rm I/O} + P_{\rm static}$ and $P_{\rm comp} + P_{\rm static}$ is much greater or much smaller than 1, where the difference between the two optimal periods might differ more than that.

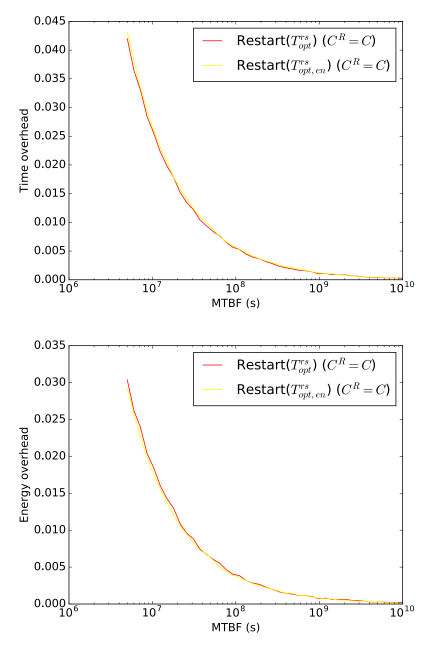


Figure 13: Impact of optimizing the time overhead or the energy overhead on the time overhead (left) or the energy overhead (right) as a function of the MTBF (C=60s, 10^5 pairs of processors).

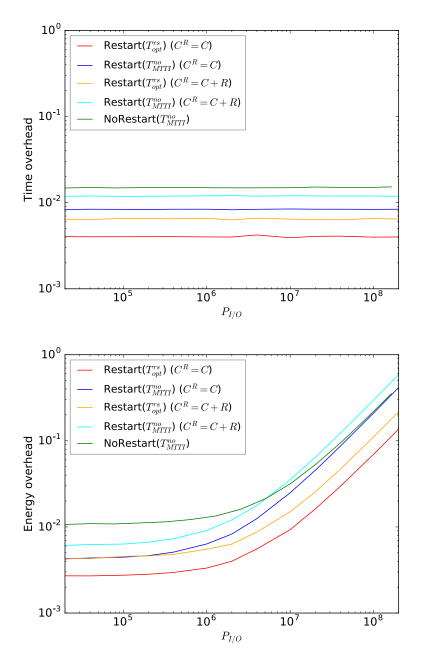


Figure 14: Time and energy overheads when varying $P_{\rm I/O}$ (MTBF 5 years, $C=60{\rm s},\,b=10^5$) when optimizing the time overhead.

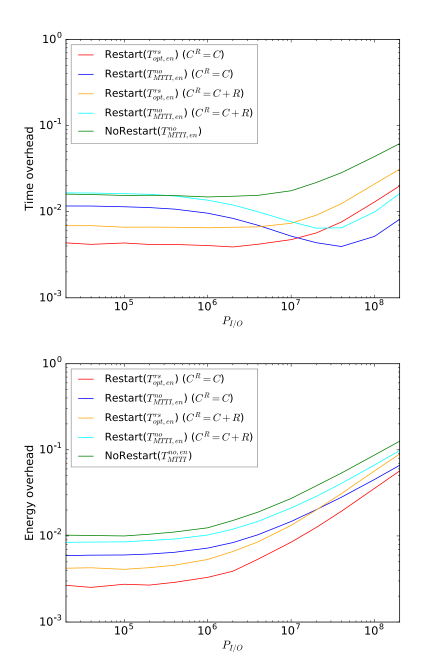


Figure 15: Time and energy overheads when varying $P_{\rm I/O}$ (MTBF 5 years, $C=60{\rm s},\,b=10^5$) when optimizing the energy overhead.



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