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Vitamin C Maintenance against Cell Growth Arrest and Reactive Oxygen Species Accumulation in the Presence of Redox Molecular Chaperone hslO Gene

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Abstract: Chromosome damage combined with defective recombinase activity renders cells inviable, owing to deficient double-strand break repair. Despite this, recA polA cells grow well under either DNA damage response (SOS) conditions or catalase medium supplementation. Catalase treatments reduce intracellular reactive oxygen species (ROS) levels, suggesting that recA polA cells are susceptible to not only chronic chromosome damage but also ROS. In this study, we used a reducing agent, vitamin C, to confirm whether cell growth could be improved. Vitamin C reduced ROS levels and rescued colony formation in recAts polA cells under restrictive temperatures in the presence of hslO, the gene encoding a redox molecular chaperone. Subsequently, we investigated the role of hslO in the cell growth failure of recAts polA cells. The effects of vitamin C were observed in hslO+ cells; simultaneously, cells converged along several ploidies likely through a completion of replication, with the addition of vitamin C at restrictive temperatures. These results suggest that HslO could manage oxidative stress to an acceptable level, allowing for cell division as well as rescuing cell growth. Overall, ROS may regulate several processes, from damage response to cell division. Our results provide a basis for understanding the unsolved regulatory interplay of cellular processes.

Keywords: vitamin C; redox chaperone; reactive oxygen species metabolism; hslO

1. Introduction

In combination with recombinase defects, chromosome breakage and double-strand break repair deficiencies render cells inviable. However, in Escherichia coli recAts polA cells, viability was retained when DNA damage response was evoked by constitutive expression of a lexA (Def) mutation [1,2]. We had previously shown that recA polA cell growth arrest was partially due to the accumulation of reactive oxygen species (ROS) and hydrogen peroxide under restrictive temperatures and rich medium conditions [2,3].

E. coli frequently encounter oxygen radicals produced as metabolic byproducts. These oxygen radicals are metabolized into hydrogen peroxide, which, in turn, is detoxified into water by catalase. Oxygen radicals also serve as a stress marker and as an effector owing to their high reactivity. Oxidative damage by superoxide anions directly suppresses metabolic activities that depend on enzymes that conjugate with iron and flavin. Oxygen radical accumulation causes stress and cell death following lethal treatments [4]. Yet, controversy remains as to whether the accumulation of oxygen radicals causes cell death or bacteriostasis [5]. Under thymine deprivation, single-strand regions and cellular oxygen radicals accumulate [6]. Furthermore, Hong et al. reported that lethal effects on the cells resulted from the accelerated accumulation of oxygen radicals in a self-replicating manner, overwhelming primary damage reparation [4]. Vitamin C plays an important role in maintaining proper oxidation–reduction balance through the neutralization of ROS and...
reactive nitrogen species (RNS) formed during cellular metabolism [7]. The ascorbate–
glutathione cycle is involved in the aforementioned process; however, it is unknown
whether this cycle is also functional in E. coli cells.

Redox signaling functions through redox chaperones and transcription factors, which
use sensor regions to bind or react with molecules such as ROS and hydrogen peroxide.
These proteins regulate the transcription or enzymatic activity of other molecules through
structural changes caused by their respective oxidation or reduction [8–11]. Redox molecu-
lar chaperones include HslO [12] and AhpC [13]. Interestingly, AhpC and vitamin C are
associated at an expression level [14]; they regulate other molecules and control biological
reactions via molecular state adjustments, oligomeric aggregation, and dissociation.
Regarding transcription factors, OxyR is oxidized and transcriptionally activated by hydrogen
peroxide [15], while soxRS is activated via ROS-induced intracellular state changes [16,17].

The HslO protein, encoded by hslO, is the redox molecular chaperone Hsp33. HslO
protects organisms from oxidative stress that leads to protein unfolding. The loss of HslO
function also renders cells susceptible to hydrogen peroxide [12]. HslO activation is
triggered by the oxidative unfolding of the HslO redox sensor domain [18]. Further, HslO
is considered a member of a recently discovered class of chaperones that requires partial
unfolding for full chaperone activity [19]. The molecular basis for regulating HslO function
begins with the release of prebound Zn from the well-conserved Zn-binding motif Cys-Xn-
Cys-X-Cys upon oxidation and activation, causing remarkable structural changes [20,21].
Once activated, HslO protects proteins from coagulating into toxic proteins, thus preventing
cellular deaths. Using the unfolding region, HslO binds the unfolding client proteins [20].
The mechanism by which the hslO docking surface captures the coagulated body of client
proteins has been elucidated [22].

The lowering of intracellular ROS levels promotes the viability of recA polA cells. Additionally, the complete absence of RecA function is tolerated in polA cells, indicating that recA polA cell mortality or growth arrest could be related to intracellular ROS levels rather than a deficiency in DSB repair. Furthermore, redox chaperone hslO is required for recA polA cells to survive [2]. However, the relationship between DNA damage, ROS production, and cell proliferation remains unclear. In this study, we explore the growth of recAts polA cells in the presence of a reducing reagent, vitamin C. We further elucidate a potential interplay between hslO in this suppressive process of recAts polA lethality and the coordination of cellular processes. We also aimed to confirm whether ROS are an important factor in explaining the sequence of events from DNA damage to growth arrest (or lethality) observed in recAts polA cells. We found that hslO+ cells were more proficient in the suppression of lethality after treatment with vitamin C than ΔhslO counterparts. This serves as a useful key in understanding what occurred in those cells. Our findings regarding ROS, a potential inhibitor for cell growth, will generate new aspects of redox signaling to explore in E. coli.

2. Results

2.1. recAts polA Cells Form Colonies in Restrictive Temperatures in the Presence of Vitamin C

In a previous report, we demonstrated that recA polA cells were capable of colony
formation in catalase-supplemented media either at a restrictive temperature (42°C) [2] or
in a rich medium [3]. Since catalase is a hydrogen peroxide-degrading enzyme, this sug-
gested that the inability of recAts polA cells to grow at restrictive temperatures resulted from
hydrogen peroxide accumulation. Vitamin C is an antioxidant that scavenges superoxide,
hydrogen peroxide, and singlet oxygen in vitro [23]. This prompts the question of whether vitamin C affects cellular responses to hydrogen peroxide metabolism (Figure 1). In the spot
assay, E. coli cell particles were spotted on plates and then incubated at either permissible
(30°C) or restrictive temperatures (42°C). As seen in Figure 1, AB1157 derivative cells,
including TK3077 (recAts polA), TK3276 (recAts polA ΔhslO), TK3473 (recAts polA ΔhslO
pEXvec), and TK3474 (recAts polA ΔhslO pEXhslO), exhibited similar growth at 30°C. In
contrast, cells failed to grow at 42°C without vitamin C, except for TK3077 cells spotted
with $2 \times 10^6$ particles, which were confirmed to be temperature-sensitive. Interestingly, the recAts polA strain (TK3077) could grow on plates at 42 °C when plates were supplemented with either 100 μM or 1 mM, but 10 μM vitamin C was insufficient to encourage growth. A 100- to 1000-fold increase in colony-forming ability with 100 μM vitamin C was observed when compared with cells without vitamin C at 42 °C. In contrast to the TK3077 cells, the recAts polA ΔhslO strain (TK3276) failed to grow with the addition of 100 μM vitamin C at 42 °C. In addition, TK3276 cells were transformed with either vector (TK3473) or with the hslO-expressing plasmid pEXhslO (TK3474). As a result, TK3474 cells could form colonies at 42 °C when $2 \times 10^6$ bacterial particles were spotted in the presence of 100 μM vitamin C, but not for $2 \times 10^5$ particles. TK3473 cells at 42 °C failed to form colonies, even at $2 \times 10^6$ bacterial particles. Therefore, it is assumed that the colony formation ability of TK3474 cells was improved more than 10-fold compared to that of TK3473 cells. Thus, vitamin C, a ROS scavenger, restored colony formation at restrictive temperatures in recAts polA cells.

Figure 1. Effects of vitamin C addition to plate medium on colony formation of recAts polA cells. Cells were spotted with the number of particles indicated on the left-hand side and incubated at 30 °C and 42 °C for 2 days. From left to right: recAts polA (TK3077), recAts polA ΔhslO (TK3276), recAts polA ΔhslO pEXvec (TK3473), and recAts polA ΔhslO pEXhslO (TK3474). The number of particles between each spot ranged from $2 \times 10^2$ to $2 \times 10^6$ cells and varied 10-fold at each dilution. A combination of the photos is shown from left to right: without vitamin C at 30 °C; no vitamin C, 10 μM, 100 μM, and 1 mM vitamin C at 42 °C.

To directly assess the effect of vitamin C on cell growth, we used liquid media, in which cell growth could be easily measured. The bacterial culture was divided into four equal volumes in an early logarithmic phase (O.D.600 = 0.1) at 30 °C. Figure 2 shows the growth of cells under four different conditions: (a) absorbance or (b) the number of E. coli cells. As shown in Figure 2a, the recAts polA cells (TK3077) at 42 °C showed slightly improved growth at 16 h with the addition of 100 μM vitamin C (blue), compared with cells without vitamin C (red). The number of bacterial cells at 16 h is also shown in Figure 2b. However, no remarkable differences were observed regardless of vitamin C presence in TK3077 (Figure 2b). Nonetheless, a significant difference in cell particles was observed between TK3077 (in the presence of hslO) and TK3276 (in the absence of hslO) at 42 °C with the addition of 100 μM vitamin C (Welch’s t-test, $p < 0.05$). This indicates that hslO is necessary for vitamin C to stimulate a slight increment in cell growth and division. Colony formations were remarkably restored in TK3077 cells with vitamin C at 42 °C (Figure 1), in spite of only a slight increase in cell particles. Thus, it was necessary to examine whether these cell particles were membrane-damaged. Therefore, we assayed dead cells with compromised membrane integrity using propidium iodide (PI) staining. Membrane-damaged dead cells were not detected at 30 °C (Figure S1a,b) but were detected at 42 °C, as indicated by blue arrowheads (Figure S1c,d). However, contrary to our expectation, the membrane-damaged cell population (blue arrowheads) did not show a remarkable decrease following vitamin C addition. Meanwhile, a minor cell population, staining poorly for both PI and SYTO9 (Figure S1, red arrowheads), had shrunk. This suggests...
that vitamin C addition did not remarkably reduce the number of membrane-damaged cells. It is unclear whether this small change suffices to explain the observed increase in colony formation in Figure 1. Furthermore, this does not take into consideration any possible decrease in the number of anucleate cells. Therefore, we considered whether qualitative rather than quantitative changes in chromosomal DNA might be involved in the cell viability decrement at restrictive temperature in recAts polA cells.

Figure 2. Vitamin C effect on recAts polA cell growth in permissive and restrictive conditions. (a) Optical growth inhibition at a restrictive temperature in shift-up experiments. The growth curves for the shift-up experiment at the logarithmic phase are shown. The changes in turbidity (O.D. 600 arbitrary unit) over time can be seen, starting when the culture medium is divided. TK3077 and TK3276 cells are indicated using circles and squares, respectively. The progression of cell growth is indicated by lines colored gray (without vitamin C) and black (with 100 µM vitamin C) for treatments at 30 °C. Lines are colored red (without vitamin C) and blue (with 100 µM vitamin C) for treatments at 42 °C. Each point is shown as the average and standard error of the mean (SEM) (n ≥ 3). (b) Cell proliferation inhibition at a restrictive temperature in shift-up experiments. Cells cultured for 16 h were measured (particles/mL) using flow cytometry. Treatment combinations are shown from left to right: seed culture (s), cell culture without vitamin C (−C), or cell culture with 100 mM vitamin C (+C) at 30 °C and 42 °C for TK3077 and TK3276 cells. Each bar represents the average and standard error of the mean (SEM) (n ≥ 3). *p < 0.05, according to Welch’s t-test (n = 6).

2.2. Vitamin C Is Involved in Maintaining the Colony-Forming Ability of recAts polA Cells at Restrictive Temperatures

As shown in Figures 1 and 2, recAts polA cells supplemented with vitamin C were able to form colonies at 42 °C and showed slightly improved growth over those without vitamin C in a liquid medium. These results might suggest that the effect of vitamin C did not support cell growth directly, but rather indirectly promoted colony formation. Therefore, we attempted to compare the effect of vitamin C on viability, a major qualitative change that directly relates to cell growth, in both TK3077 and TK3276. We examined the effects of preincubation treatments with vitamin C at either permissive (30 °C) or restrictive temperatures (42 °C) before cells were spotted on plates. Following the treatment, the survival of cells was determined on plates at either 30 °C (Figure 3a,b) or 42 °C (Figure 3c,d). With regard to the plate culture at 30 °C, the recAts polA cells with hislo+ (TK3077) did not show any response to the addition of vitamin C with 30 °C permissive pretreatment (lanes 1 and 2). TK3077 cells decreased in colony formation in restrictive pretreatment, agreeing with the temperature-sensitive phenotype of TK3077 cells (lanes 3). Meanwhile, cells with 100 µM vitamin C supplementation and 42 °C restrictive pretreatment maintained the ability to form colonies, >10-fold better than those without vitamin C supplementation (lanes 3 and 4). Interestingly, colony-forming abilities with vitamin C for 42 °C restrictive pretreatment remained 10-fold better in comparison to those of 30 °C permissive pretreatment (lanes 2 and 4). These results indicated that TK3077 cells with 42 °C restrictive pretreatment maintained their cell integrity until colony formation, owing to the addition of vitamin
C. We further examined treatment effects on TK3276 cells (the hslO deletion derivative). These cells showed no substantial difference in colony-forming abilities between those with and without vitamin C supplementation at both 30 °C and 42 °C treatment (lanes 5–8). Interestingly, TK3276 cells with 30 °C treatment showed better colony formation than TK3077 cells (lanes 1 and 5). Similar colony-forming abilities were observed between the hslO+ and hslO deletion mutant cells after 42 °C treatment without vitamin C (lanes 3 and 7). Remarkably, TK3276 cells formed colonies 100 times better than TK3276 cells with 42 °C treatment and vitamin C (lanes 4 and 8). Regarding the plate culture at 42 °C, TK3077 cells with vitamin C formed colonies mainly at 2 × 10^6 (lanes 11 and 12). These results suggest retention of colony-forming capabilities with vitamin C in 42 °C treatment, which would not otherwise have occurred from acquiring a temperature-resistant mutation. Therefore, vitamin C was somehow maintaining cellular integrity, which likely resulted in the observed colony formation. Furthermore, hslO contributed to whether cells could grow or not in the presence of vitamin C after restrictive pretreatments.

![Figure 3](image-url)

**Figure 3.** The effects of pretreatment with vitamin C on colony formation of recAts polA cell culture. Cells were spotted with the indicated number of particles on M9GCAA plates. Spots are shown from left to right: liquid cell culture without and with vitamin C at 30 °C and 42 °C. The number of particles between each spot (indicated with arrowheads) ranged from 2 × 10^1 to 2 × 10^6 cells and varied 10-fold at each dilution. A combination of the panels is shown from left to right, pretreatment (broth) at either 30 °C or 42 °C. Plate cultures (plates) were incubated at either 30 °C or 42 °C. Posi indicates 2 × 10^3 cells of AQ10459 for spots on plates at 42 °C.

2.3. Vitamin C Reduced Accumulation of Intracellular ROS Levels at Restrictive Temperatures in hslO+ recAts polA Cells

We investigated the effect of vitamin C on the accumulation of intracellular ROS levels in recAts polA cells at a restrictive temperature (Figure 4). TK3077 (recAts polA) and TK3276 (recAts polA ΔhslO) cells were cultured at 30 °C and divided into four equal parts in the early logarithmic growth phase (O.D.600 = 0.1). Histograms showed the accumulation of intracellular ROS levels at 16 h after cultivation at either 30 °C or 42 °C with or without 100 μM vitamin C supplementation. The accumulated intracellular ROS levels in TK3077 cells were almost the same under the permissive condition (30 °C) with or without vitamin C supplementation (Figure 4a). In contrast, at the restrictive temperature (42 °C), TK3077 cells with vitamin C (blue) had lower intracellular ROS levels compared with those without
2.3. Vitamin C Reduced Accumulation of Intracellular ROS Levels at Restrictive Temperatures

We investigated the effect of vitamin C on the accumulation of intracellular ROS levels. In TK3077 cells, intracellular ROS levels decreased with the addition of vitamin C. Contrary to TK3077 cells, intracellular ROS levels of TK3276 increased even with the addition of vitamin C at 42 °C. This was in contrast to TK3077 cells, in which intracellular ROS levels decreased with the addition of vitamin C.

Next, we analyzed the dose-dependent response in mean ROS levels towards vitamin C. We observed poor dose response; however, the effect of vitamin C was observed at 100 µM in TK3077 (Figure S2). This poor dose response and unstable results at both 300 and 1000 µM might be due to a hormesis effect [24], in which a response is not always according to the dosage.

We further statistically evaluated the effect of 100 µM vitamin C on the intracellular ROS levels at 30 °C and 42 °C (Figure 5). As shown in Figure 5a, mean intracellular ROS levels were not influenced by vitamin C in TK3077 and TK3276 cells at 30 °C. However, those of TK3077 cells at 42 °C were significantly reduced with the addition of 100 µM vitamin C (Welch’s t-test; p < 0.05, n = 7). Conversely, mean intracellular ROS levels increased upon the addition of vitamin C in the TK3276 cells. Thus, the vitamin C effect on intracellular ROS levels differs between TK3077 and TK3276 cells.

Further, inoculation experiments were also performed to confirm the results above at either 30 °C or 42 °C, in the absence or presence of vitamin C. This method was especially convenient for measuring intracellular ROS levels (Figure 5b). The addition of vitamin C significantly decreased mean ROS levels in TK3077 cells (Figure 5b) (p < 0.05, n = 7). Contrary to the results in Figure 5a, ROS levels in TK3276 cells responded to vitamin C in this inoculation experiment. This might have resulted from the difference in the growth phase of the two experimental systems: the system in Figure 5a used the logarithmic growth phase, while the system in Figure 5b used the stationary phase. Growth in Figure 5a,b
corresponded to that shown in Figure 2a,b and Figure S3, respectively. In the inoculation experiment, unlike experiments divided at the logarithmic phase, neither TK3077 nor TK3276 cells showed any growth at 42 °C (Figure S3). These growth differences may have resulted from the average intracellular ROS levels in the two experimental systems. This hypothesis is studied below, at least with regard to cell growth and ROS levels.

Figure 5. Vitamin C effect on mean reactive oxygen species (ROS) levels of recAts polA cells and recAts polA ΔhslO cells in permissive and restrictive conditions. (a) Mean ROS level suppression at a restrictive temperature in the shift-up experiments. In the early logarithmic growth phase (O.D.600 = 0.1), the TK3077 (recAts polA) and TK3276 (recAts polA ΔhslO) cultures were divided into four portions, and each was incubated either with or without vitamin C at 30 °C and 42 °C. The mean ROS levels in the TK3077 and TK3276 cells were determined using CellROX Deep Red staining. The treatment combinations for each bar are shown from left to right: cell culture without vitamin C (−C) or with 100 µM vitamin C (+C) at both 30 °C and 42 °C for TK3077 and TK3276 cells. Each bar represents the average and standard error of the mean (SEM) (n ≥ 7). * p < 0.05, according to Welch’s t-test (n = 7). (b) Mean ROS level suppression at a restrictive temperature in inoculation experiments. TK3077 (recAts polA) and TK3276 (recAts polA ΔhslO) cells were inoculated with 1/100 of the overnight seed culture and incubated either with or without 100 µM vitamin C at 30 °C or 42 °C for 16 h. The cell ROS levels...
were determined, and the ROS levels of the seed and cultured cells at 30 °C or 42 °C for 16 h were compared. Treatment combinations for each bar are shown from left to right: seed culture, cell culture without vitamin C (−C), or cell culture with 100 µM vitamin C (+C) at both 30 °C and 42 °C for TK3077 and TK3276 cells. Each bar represents the average and standard error of the mean (SEM) (n ≥ 7). * p < 0.05, according to Welch’s t-test (n = 7).

2.4. Vitamin C Affected Intracellular ROS Levels in recAts polA Cells by Synchronizing ROS Levels with Growth

To interrogate the relationship between ROS levels and growth, we used results from Figure 5a to construct a scatterplot in two dimensions with growth on the vertical axis and mean ROS levels on the horizontal axis (Figure 6). When vitamin C was added at 42 °C (blue), the mean intracellular ROS levels were reduced in TK3077 cells (Figure 6a) compared with those in TK3276 cells (Figure 6b). Measurement points for TK3276 cells showed neither arbitrary units in growth at around 200–600 RFU (relative fluorescence unit, RFU) nor mean ROS levels at around 175–200 RFU. This suggests that TK3276 cells failed to alleviate ROS levels with vitamin C due to the absence of hslO, resulting in a lack of proficient growth recovery. Thus, vitamin C countered the growth inhibition of TK3077 cells at 42 °C, but not that of TK3276 cells.
Figure 6. Scatterplots of how vitamin C affects mean ROS levels and cell growth in recAts polA cells and recAts polA ΔhslO cells in permissive and restrictive conditions. (a) Effect of vitamin C on a scatterplot with mean ROS levels and cell growth in recAts polA cells (TK3077) in permissive and restrictive conditions. In the early logarithmic growth phase (O.D.\textsubscript{600} = 0.1), the TK3077 (recAts polA) culture was observed. The measurements were plotted for ROS levels and optical growth on the x- and y-axis, respectively. Round symbols indicate measurements on cells without vitamin C (−C) or with 100 µM vitamin C (+100 µM C). (b) Vitamin C effect on a scatterplot for mean ROS levels and cell growth in recAts polA ΔhslO (TK3276) cells in permissive and restrictive conditions. In the early logarithmic growth phase (O.D.\textsubscript{600} = 0.1), the TK3276 (recAts polA ΔhslO) culture was examined. The results are represented just as in (a) using square symbols. (c) Vitamin C effect on a scatterplot with mean ROS levels and cell concentrations in permissive and restrictive conditions. This replot shows ROS levels and cell concentrations on the x- and y-axis, respectively. Measurement points are indicated as in (a,b).

We wondered whether recAts polA cells synchronized in growth and ROS levels after 16 h of incubation (Figure 6a,b). We further confirmed this possibility with cell counts in culture and ROS levels. As shown in Figure 6c, both TK3077 and TK3276 cells cultured at 30 °C were aligned at ROS levels at around 175 RFU. Meanwhile, TK3276 cells cultured at 42 °C were aligned at 2.5 × 10\textsuperscript{8} cells/mL, corresponding to nearly 2.5 times as much as an initial inoculated cell concentration. Contrary to these results, TK3077 cells incubated at 42 °C without vitamin C had aligned in both ROS levels and the cell concentration, resulting in an L-figured distribution. Furthermore, TK3077 cells incubated at 42 °C with vitamin C were mainly aligned in ROS levels at 175 RFU. In summary, vitamin C, in the presence of hslO, would partly improve both growth and mean ROS levels at 42 °C. Further, these results suggest that there is a growth threshold in recAts polA cells with regard to mean intracellular ROS levels after 16 h. Thus, these results indicated that vitamin C proficiently restored cell growth in the presence of hslO among recAts polA cells. Interestingly, it is possible that cell division processes might have caused recAts polA cells’ growth failure at restrictive temperature as they divide only once. Cell division could be inhibited through multiple processes, including DNA replication and segregation, since it is the final stage of quality control. Thus, DNA levels were of interest in recAts polA cells.

2.5. Vitamin C Influences a Convergence of Cell Population with Chromosomal DNA Level Rather Than the Temperature-Sensitive Recombination of the recAts Mutation

As previously mentioned, recAts polA cells might postpone (or inhibit) cellular processes, including cell division, due to DNA damage. It was likely that DNA metabolism including a completion of DNA replication was a prerequisite for chromosome segregation and so on. The initiation of DNA replication is tightly associated with these cell masses and growth rates [25]. Thus, slowly growing cells and cells in the stationary phase possess
chromosomal DNA levels with one or two chromosomes. Accordingly, we focused on monitoring the chromosome status of recAts polA cells with their DNA levels through this cell cycle progression. The addition of vitamin C was involved in the maintenance or promotion of colony formation in recAts polA cells at restrictive temperatures. Recalling Figure S1, most recAts polA cells did not show notable membrane damage at restrictive temperatures; simultaneously, vitamin C had no effect on damaged populations. Cell death or growth arrest is unlikely to occur suddenly without any precursory phenomenon. Therefore, as a prerequisite for colony formation, we compared the effect of vitamin C on chromosome status through quantitative chromosome staining with PicoGreen; simultaneous deficiencies in recA and polA are thought to influence chromosome integrity. To compare the chromosome status, ploidy analysis using flow cytometry was performed. Cells incubated at 30 °C showed three bands corresponding to DNA quantity, as in one, two, and three ploidies (Figure 7). Cells incubated at 42 °C did not show these bands. Nonetheless, TK3077 cells supplemented with vitamin C at 42 °C still showed major converging bands with approximately two ploidies. The bands appearing from TK3077 cells with vitamin C supplementation at 42 °C were similar to those appearing at 30 °C. These results were also analyzed with histograms (Figure S4). The histograms show sharp peaks corresponding to converged chromosomes at 30 °C; however, those at 42 °C are difficult to distinguish. As described above, even in randomly growing cells, the majority of cells converged with several DNA levels owing to both the completion of DNA replication and the relatively short replicating time, thus presenting converged histograms or bands. Therefore, DNA levels in these experiments were converged in one or two ploidies. Interestingly, histograms in TK3276 cells show that DNA quantity had increased at 42 °C compared with that at 30 °C (Figure S4, lower panels). TK3077 cells with vitamin C supplementation at 42 °C showed several peaks corresponding to converged DNA content (Figure S4, upper right). In agreement with the results presented in Figure S1, the anucleate cell population was slightly decreased upon the addition of vitamin C in TK3077 cells but not in TK3276 cells (thick green arrowhead in Figure 7 and Figure S4). It is unclear whether this decrease in anucleate cells affects growth failure. We also quantified DNA content as a function of time in the presence or absence of vitamin C (Figure S5). No remarkable differences were observed between TK3077 and TK3276 cells. Thus, vitamin C mainly alters chromosome states such as chromosome ploidy rather than the amount of DNA. Naturally, the stationary phase cells have one or two chromosomes due to chromosomal replication and segregation. Therefore, chromosome ploidy could be one of the phenotypical markers for the successful progression of cellular processes towards the stationary phase. TK3077 and TK3276 cells showed irregular ploidies even in permissive temperature (30 °C), and those abnormalities were enhanced at restrictive temperature (42 °C). This enhancement of abnormalities was partially alleviated in TK3077 cells with vitamin C, even at 42 °C. These results suggest that TK3077 cells could complete their chromosomal replication in the presence of vitamin C at 42 °C, thus supporting colony formation. Without vitamin C, TK3077 and TK3276 cells failed to complete or delayed the processes toward the stationary phase. Therefore, qualitative rather than quantitative changes in chromosomes might be involved in the increase of viability with the addition of vitamin C, through synchronization between growth and ROS levels.
chronosomal replication in the presence of vitamin C at 42 °C, thus supporting colony formation. Without vitamin C, TK3077 and TK3276 cells failed to complete or delayed the processes toward the stationary phase. Therefore, qualitative rather than quantitative changes in chromosomes might be involved in the increase of viability with the addition of vitamin C, through synchronization between growth and ROS levels.

Figure 7. Effect of vitamin C on the ploidy of recA polA cells in permissive and restrictive conditions. In the early logarithmic growth phase (O.D.₆₀₀ = 0.1), the TK3077 (recA polA) and TK3276 (recA polA ΔhisO) cultures were divided into four portions, and each portion was incubated at 30 °C and 42 °C either with or without vitamin C for 16 h. The DNA levels in the cells were determined using PicoGreen staining. Strains and cultivating temperatures are indicated on the left of the panels. Whether vitamin C was added (with or without 100 µM vitamin C) is indicated at the top of the panels. Arrowheads indicate ploidies as 1 (black), 2 (red), and 3 (blue). Thick green arrowheads indicate a position of anucleate cells. In total, 20,000 particles were analyzed with flow cytometry using fluorescence, with channel (FL-1H) on the x-axis and signal/side scatter signal (SSC) SSC-H on the y-axis.
It could be argued that adding vitamin C might influence the temperature sensitivity of the recombination reaction in recAts cells. Since E. coli cells with the polA25 mutation had low P1 transduction efficiency, we used polA+ cells with recAts mutation to evaluate the effect of vitamin C on temperature sensitivity. recA+ parental cells (AQ10459) showed a transduction frequency of nearly $10^{-5}$ at both 30 °C and 42 °C, with or without 100 μM vitamin C (Figure 8). AQ10546 recAts cells showed a transduction frequency of about $5 \times 10^{-6}$ with or without vitamin C at 30 °C. Likewise, at 42 °C, the transduction frequency of AQ10546 cells (at $1 \times 10^{-7}$) did not differ between treatments with and without vitamin C. These results suggested that adding vitamin C did not influence the transduction efficiency of either recA+ or recAts cells. Therefore, we presume that suppression of temperature sensitivity did not result from a restoration of genetic recombination due to vitamin C. This agreed with our previous result that ΔrecA polA cells were viable. Thus, these results supported the hypothesis that the progression of cellular processes toward the completion of DNA replication was affected in recAts polA cells.

![Figure 8](image.png)

**Figure 8.** Effects of vitamin C on transduction frequencies in both permissive and restrictive temperatures. First, $10^9$ cells of either AQ10459 or AQ10546 cells were transduced with malB::Tn9. The cells with Cm resistance (Cm<sup>r</sup>) were subsequently selected. Further, Cm<sup>r</sup> colonies were confirmed via the phenotype of maltose assimilation on MacConkey agar plates supplemented with 1% maltose. The transduction efficiencies were calculated as the ratio of Mal<sup>+</sup> cells to the number of particles. The strains are AQ10459 (recA<sup>+</sup>) and AQ10546 (recAts). Each bar represents the average and standard error of the mean (SEM) (n ≥ 3); without (−C) or with vitamin C (+C) at 30 °C and 42 °C. **p < 0.01, according to Welch’s t-test (n = 5).

3. Discussion

Our previous study on recAts polA lethality reported that intracellular ROS accumulation was associated with lethality at the restrictive temperature [2]. However, why hslO was required to suppress recAts polA lethality was unclear. Therefore, we intended to elucidate the role that hslO played in the suppression of lethality. In this study, we determined that the accumulation of radical oxygen was partly alleviated by the addition of vitamin C and the presence of hslO. recAts polA lethality (or growth inhibition) was
ameliorated by the lowering of ROS levels in cells by \textit{hslO}. These findings are in agreement with our previous report that \textit{recAts polA lexA51} cells required \textit{hslO} for suppression of temperature sensitivity. Our present investigation has suggested that \textit{hslO}, in the presence of vitamin C, plays an important role in cellular function integrity, including the completion of chromosome replication and colony formation via lowering ROS levels. In other words, ROS production upon chromosome damage would negatively regulate normal processes of the cell (e.g., through redox signaling and prolonged damage). This leads to a gradual loss of cell functional integrity and, eventually, cell death. In this case, the reduction in ROS alone might be insufficient to accomplish the progression of cell growth. These possibilities are further discussed later.

Even though hydrogen peroxide and superoxide anions do not directly oxidize DNA, these molecules contribute directly or indirectly to the production of highly reactive hydroxyl radicals that damage bacterial chromosomes [26]. Thus, reducing the concentration of oxygen radicals early can prevent or delay damage. Due to the electron-donating ability of vitamin C, it could act as a free radical scavenger and reduce iron in its highly oxidized state to Fe$^{2+}$ [23]. The effects of vitamin C are still controversial, although many trials are underway. This might be due to hormesis-like effects [24]: Kontek et al. reported that vitamin C ameliorated DNA damage only in the presence of H$_2$O$_2$ [27]. This indicates that the addition of reducing reagents could regulate the excess oxidative intracellular state, which is consistent with our results. Thus, these results suggest that \textit{E. coli} can mediate the damage caused by ROS with the addition of vitamin C.

Furthermore, vitamin C could reduce H$_2$O$_2$, ROS, and disulfide bonds through cellular metabolism [28]. \textit{E. coli} possesses a weak dehydroascorbate reductase activity [29], but it is unknown whether the ascorbate–glutathione cycle is also functional in \textit{E. coli} cells. However, the bacterial peroxiredoxin AhpC might contribute to the cycle via the deglutathionylation cycle [13], and AhpC and vitamin C are associated at the expression level [14], suggesting a response to oxidative stress generated during the aerobic metabolism of vitamin C. Vitamin C could prevent the detrimental effects of H$_2$S, which reduces glutathione levels in \textit{E. coli} [30], and AhpC was resistant to inactivation by peroxidation [31] through suppressing aggregation of client proteins in heat shock conditions. Interestingly, AhpC was reported to function as a redox molecular chaperone, similar to HslO, to regulate the oligomer state of the target proteins. These findings suggest a possible crosstalk between vitamin C and redox molecular chaperones.

Molecular redox chaperones such as \textit{hslO} play a crucial role under oxidizing conditions. \textit{hslO} is unique for promptly detecting oxidation stress, a possible cause of the unfolding of proteins, and thus becomes activated. Furthermore, once activated by oxidation, \textit{hslO} protects proteins from coagulating and forming toxic proteins, which then protects bacterial cells from cellular death. These results agree with our observations, leading to a new direction in research of both DNA damage response and \textit{hslO}; however, direct linkages have not been fully understood yet. Using the unfolding region, the \textit{hslO} binds with the unfolding client proteins [20]. The candidates for the client proteins that possess a DNA-repairing ability have not yet been reported, although how HsLO captures client proteins on its docking surface has been elucidated [22]. It is likely that some proteins that are not clients of other redox chaperones could be associated with the Srp pathway. Protein aggregates might be involved in cellular process progression and general intracellular ROS regulation.

Notably, in our study, the abnormalities of ploidies were partially restored in TK3077 cells with vitamin C at 42 °C. Chromosome ploidies are ordinarily 2$^n$, resulting from proper termination of chromosome replication, segregation, and septation in \textit{recA$^+$} cells at the stationary phase. However, \textit{recA$^-$} cells do not show a 2$^n$ pattern in the stationary phase because of RecA function deficiencies, causing a failure of proper chromosome segregation in catenated chromosomal DNA. Even at a permissive temperature, \textit{recA200 (recAts)} mutation caused insufficient RecA function, which was exacerbated at restrictive temperatures. Lanzov et al. reported that the \textit{recAts} mutation caused hyper-recombination
at a restrictive temperature [32]. Kogoma reviewed interplays between recombination and replication [33]. Therefore, at the restrictive temperature, it is possible that *recA*-*polA* cells caused hyper-replication, and replicative stress might be elevated in *recA*-*polA* cells. Of note, *ΔrecA polA* cells failed to grow in a rich medium where replicative stress would be elevated [3].

We have reported in a previous paper that significant chromosome breakdown could not be observed [2]. However, we were not able to identify what causes growth failure of the *recA*-*polA* cells. Therefore, we instead focused on the process leading to ROS accumulation and also characterized the damaged chromosomes. As a result, *recA*-*polA* cells could grow in the presence of both the *hsLO* gene and vitamin C under restrictive conditions. Conversely, we found that *recA*-*polA* cells failed in the convergence of cell population with their DNA content at the restrictive temperature; however, this phenomenon was restored in the presence of *hsLO* and vitamin C. It is not yet known whether the growth failure and the convergence of cell population with DNA content are related to each other. It is interesting that slowly growing cells possess one or two ploidies, indicating that completion of replication was always observed for those slowly growing cells before cell proliferation via cell division. Meanwhile, the oxidation of DNA polymerase likely leads to their inactivation. Therefore, *hsLO* and vitamin C can function together to maintain cellular oxidative conditions at allowable levels, supporting cellular metabolism and the completion of DNA replication required for the convergence of the cell population with DNA contents, thus enabling cell division and cell proliferation.

Regarding the use of a synthetic lethality experimental system for chromosome damage, we observed growth arrest of *recA*-*polA* cells that was ameliorated with reducing reagents such as vitamin C. Additionally, cells were growing with synchronization in growth and ROS levels even in restrictive temperatures. This is consistent with our previous observations on the effects of catalase [2,3]. The source of ROS is unclear; however, we now know how the *recA*-*polA* cells mediate ROS stress. Our study substantially depended on analysis with chemical probes and fluorescence dyes by flow cytometry. This approach is reliable for population analysis. Yet, it is perhaps unreliable for particular molecules and fine quantitative analysis. Notably, chromosome ploidy was restored by vitamin C at the restrictive temperature. Intracellular ROS levels could influence chromosome replication and/or segregation, indicating that these cellular processes might be the targets of ROS stress. Conclusively, these findings have opened doors to new forms of interplay between the regulation in the progression of replication and DNA damage via redox signaling. These findings will contribute to understanding how DNA damage signals are transduced into chromosome replication during cell proliferation. Regulations, including modifications of particular cellular reactions or mechanisms, might suffice for now as one thread in the larger tapestry of oxidative stress response pathways. However, another thread comes from the interplays between intracellular metabolites and individual cellular processes as well. We may be on the verge of a new era in which all the threads will come together for us to see the greater picture.

4. Materials and Methods

4.1. *E. coli* Strains and Media

The *E. coli* strains used in this study are represented in Table S1 and are also described as follows: TK3077 was the same as AQ10549 but restocked. AQ10459, AQ10546, AQ10865, TK3077, TK3276, TK3473, and TK3474 were AB1157 derivatives described previously. TK3019 was constructed using phage *P1vir*-mediated transduction [34,35]. Constructed with infection of *P1vir* phage from AQ10865, the *Cm*+ colony was selected. Subsequently, a temperature-resistant strain for growth was also confirmed for *lexA*+. The resulting cells were AQ634 *malB::Tn9 lexA*+. Cells were grown at 30 °C in M9 salts–glucose minimal (M9G) media supplemented with casamino acids (CAAs) (0.2%; Difco Laboratories, Detroit, MI, USA); thymine (1 mg/mL); thiamine (1 µg/mL); appropriate amino acids (50 µg/mL):
arg, thr, leu, trp, his, pro (M9GCAA medium); and antibiotics: ampicillin (20 µg/mL), kanamycin (55 µg/mL), spectinomycin (40 µg/mL), and streptomycin (100 µg/mL).

4.2. Cultivation and Sampling Methods

M9GCAA liquid media (2 mL and 15 mL) were placed in test tubes and 100 mL Erlenmeyer flasks, respectively, and cultured aerobically at either 30 °C or 42 °C. Cells were inoculated with 1/100 volume of cells grown overnight on M9GCAA broth.

In shift-up experiments, cells were cultured in M9GCAA medium until O.D.₆₀₀ = 0.1. Cells were then divided into two to four equal portions for the addition of reagents. After the treatments, the cell culture absorbance values were determined at O.D.₆₀₀, and DNA content and ROS analyses were performed every 2 h for a total of 16 h. For the time-course experiment, typical sample volumes were 600 µL for O.D.₆₀₀, 200 µL for DNA content, and 4 µL for ROS analyses.

In inoculation experiments, cells without or with a predetermined menadione concentration were incubated at 120 spm for 16 h at either 30 °C or 42 °C.

4.3. Plasmid Construction

An in-frame hslO expression plasmid (pEXhslO) was constructed by introducing NcoI–EcoRV–NdeI site NdeI-blunted pMW119 (Nippongene, Tokyo, Japan) derivative (pTK1424) at 3 bp downstream of the Shine–Dalgano sequence of lacZ by PCR. Inverse primers were 5′-GATATCCATATGACCATGATTACGCCAAG and 5′-TGGATATCCATGGCTGTTTCCTGTGTGAAATTG. The inverse PCR product was digested with EcoRV and then self-ligated. A ligated DNA was transformed into XL-1 Blue (Agilent, California), and a blue colony was selected. NdeI-sensitive plasmids were further confirmed by sequencing and designated as pTK1434. Thus, CT sequences just before the first ATG of lacZ of pMW119 were replaced by CCATGGATATCCAT, which resulted in the addition of NcoI–EcoRV–NdeI sites to lacZ in pTK1434.

hslO/srpC gene fragments were amplified through 15 cycles with 94 °C/1 min, 40 °C/1 min, and 72 °C/2 min. Primers were 5′-TTAAGCTTAGCCATGGCTCATATGATTAGGCCGCAACATG and 5′-TTGGATCCTGTACATTAATGAACTTGCGGATC. Amplified DNA was digested with HindIII and BamHI and cloned into the HindIII and BamHI sites of pSTV28. Sequences were confirmed, and the Spc cassette was subsequently introduced into the SmaI site. This transient plasmid and pTK1434 were digested with both NcoI and KpnI and ligated. A white colony was selected, and the resultant plasmid was designated as pEXhslO. Spc cassette was also introduced into the SmaI site of pTK1434 as pEXvec.

4.4. Determination of Survival Fraction and Cell Recovery

For relative viability (RV) determination, after incubation in M9GCAA medium overnight at 30 °C, the cells were diluted in M9 medium without a nutrient source (M9B) and then plated on M9GCAA plates supplemented with appropriate antibiotics and incubated for 16 h at either 30 °C or 42 °C. Cell concentrations of 2 × 10¹, 2 × 10², 2 × 10³, 2 × 10⁴, 2 × 10⁵, and 2 × 10⁶ were spotted on plates, and the viability was determined by photographs after 16 h of cultivation at either 30 °C or 42 °C.

4.5. Flow Cytometry Analysis

Flow cytometry was conducted as previously described [2]. For ROS analysis using flow cytometry, staining was performed according to our previous studies [2,3]. Cell cultures (4 µL) were mixed with 12.5 µM CellRox Deep Red (16 µL) at indicated times, diluted with M9 medium without organic nutrients (M9B), and stained for 30 min at 25 °C. Stained cells (20 µL) were then diluted in M9B (200 µL). Except for cells from an agar plate, we stained cells with CellRox Deep Red alone. We then used a Becton Dickinson Accuri C6 (Becton, Dickinson and Company, Ann Arbor, MI, USA) with a 640 nm laser. First, we analyzed the cell culture for the gate derived from cell particles. We used identical side scatter signal/forward scatter signal (FSC) gates, designated as P3, and collected
50,000 events. In our experiments, the rate of events was less than 2500 events/s. To analyze the acquired data, we used the C6 software (version 1.0.264.21). Each sample was plotted as a histogram vs. the red channel (FL4-A with 675 ± 15 nm filter) or ROS content (fluorescence, channel FL4-A) as either autofluorescence by the green channel (FL-1A) or as a function of the cell size (as FSC above). DNA content analysis was carried out according to Ferullo et al. [36]. For 2-dimensional ploidy analysis, side scatter (SSC)-H intensities were measured and assigned to the y-axis of the dot plot. The number of particles in cultures was determined with the BD Cell Viability Kit (Becton, Dickinson and Company, 335925), following the manufacturer’s procedure.

Live–dead staining was carried out using LIVE/DEAD BacLight Bacterial Viability and Counting Kit with the manufacturer’s recommended procedure.

4.6. Statistical Analysis

The calculation of means was performed using Microsoft Excel 2019, and the standard error of the mean (SEM) was calculated with the STDEV.P function. Welch’s t-test was also performed using the Excel program, and statistical significance was set at \( p < 0.05 \).

5. Conclusions

Temperature sensitivity of recAts polA cells was suppressed with the addition of vitamin C, which is consistent with our previous observation that the addition of catalase to plate culture restored these cells. The effect of vitamin C was related to the maintenance of cell viability rather than the restoration of cell growth. This phenomenon coincides with the amelioration of ROS levels by vitamin C, resulting in the overall restoration of cellular processes and progression toward the stationary phase. These results present a possible link between ROS levels and cell viability, revealing a partial mechanism behind recAts polA lethality.

Supplementary Materials: The supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/ijms232112786/s1. References [2,3,37] are cited in the Supplementary Materials.

Author Contributions: A.K. designed the study. A.K. and N.I. performed the genetic study and flow cytometry. A.K., S.S. and T.S. performed the DNA sequence study. H.K. obtained the photographs. All authors have read and agreed to the published version of the manuscript.

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Conflicts of Interest: The authors declare no conflict of interest.

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