The effect of hyperthermia on the DNA damage response induced by $\gamma$-rays, as determined through \textit{in situ} cell tracking

Qibin Fu$^1$, Jing Wang$^2$ and Tuchen Huang$^{1,*}$

$^1$Sino-French Institute of Nuclear Engineering and Technology, Sun Yat-sen University, Tang Jia Wan, Zhuhai 519082, P. R. China
$^2$State Key Laboratory of Nuclear Physics and Technology, School of Physics, Peking University, Beijing 100871, P. R. China
$^*$Corresponding author. Sino-French Institute of Nuclear Engineering and Technology, Sun Yat-sen University, Tang Jia Wan, Zhuhai 519082, P.R.China.
Tel: +8607563668967; Email: huangtuchen@mail.sysu.edu.cn
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\section*{ABSTRACT}
Hyperthermia (HT) acts as a cancer treatment by direct cell killing, radiosensitization, and promotion of tumor reoxygenation. The sensor proteins of the DNA damage response (DDR) are the direct targets of HT. However, the spatiotemporal properties of sensor proteins under HT are still unclear. Therefore, investigating the impact of HT on sensor proteins is of great importance. In the present study, the human fibrosarcoma cell line HT1080 stably transfected with 53BP1-GFP [the DDR protein 53BP1 fused to green fluorescent protein (GFP)] was used to investigate the real-time cellular response to DNA double-strand breaks (DSBs) induced by $\gamma$-rays. Using live-cell imaging combined with HT treatment, the spatiotemporal properties of the 53BP1 protein were directly monitored and quantitatively studied. We found that HT could delay and decrease the formation of 53BP1 ionizing radiation–induced foci (IRIF). Moreover, through the \textit{in situ} tracking of individual IRIF, it was found that HT resulted in more unrepaired IRIF over the period of observation compared with IR alone. Additionally, the unrepaired IRIF had a larger area, higher intensity, and slower repair rate. Indeed, almost every cell treated with HT had unrepaired IRIF, and the majority of these IRIF increased in area individually, while the rest increased in area by the merging of adjacent IRIF. In summary, our study demonstrated that HT could perturb the primary event in the DDR induced by IR, and this may have important implications for cancer treatment and heat radiosensitization.

\textbf{Keywords:} $\gamma$-irradiation; hyperthermia; live-cell imaging; DNA damage response; single IRIF analysis

\section*{INTRODUCTION}
Hyperthermia (HT) therapy, through increasing the temperature of tumor-loaded tissue to 40–43°C, is often applied as an adjuvant to radiotherapy \cite{1, 2}. HT acts as a cancer treatment by direct cell killing, radiosensitization, and promotion of tumor reoxygenation \cite{2}. The impact of HT on the DNA damage response (DDR) and repair is important for cancer treatment.

Accumulating evidence has shown that various DNA repair systems are targets of HT. For instance, HT could inactivate DNA-polymerase $\beta$, which is the key enzyme in base excision repair \cite{3}. For the repair of ionizing radiation (IR)-induced DNA double-strand breaks (DSBs), there are two main pathways, namely, non-homologous end joining (NHEJ) and homologous recombination (HR) \cite{4}. HT has a great impact on NHEJ through the aggregation of Ku protein, which leads to the inactivation of the DNA-binding activity \cite{5}. The effects of HT on HR are even more pronounced and diverse. Heat stress (42–45°C) could induce the translocation of MRN complex (Mre11/Rad50/Nbs1) from the nucleus to the cytoplasm \cite{6}. Moreover, the recruitment of recombinase Rad51 and protein BRCA2 are blocked at 40–42°C \cite{7, 8}.

In addition, several sensor proteins are the direct targets of HT. Based on immunofluorescent analysis, HT before or after IR could delay the formation of 53BP1 complex \cite{9}. However, traditional measurements can not reveal the dynamics of DDR within a single cell. Thus, the spatiotemporal properties of sensor proteins under HT are being discussed.
In the present study, we used human fibrosarcoma cell line HT1080 stably transfected with 53BP1-GFP [the DDR protein 53BP1 fused to green fluorescent protein (GFP)] as a DSB surrogate marker. Additionally, the confocal microscope used was equipped with a temperature chamber and CO₂ module for HT treatment. Thus, the spatiotemporal properties of sensor proteins treated with HT could be directly monitored and quantitatively studied. Using time-lapse imaging, we found that HT could delay and decrease the formation of 53BP1 IR-induced foci (IRIF). Moreover, through the in situ tracking of individual IRIF, it was observed that HT resulted in more unrepaired IRIF over the period of observation compared with treatment with IR alone. Additionally, the unrepaired IRIF had a larger area and a higher intensity as well as a slower repair rate. It is noteworthy that almost every cell treated with HT had unrepaired IRIF, and the majority of these IRIF increased in area individually, while the rest increased in area by the merging of adjacent IRIF. Taken together, the results demonstrated that HT could perturb the primary event in the DDR induced by IR, and this may have important implications for cancer treatment and heat radiosensitization.

**MATERIALS AND METHODS**

**Cell culture**
The human fibrosarcoma HT1080 cell line stably transfected with 53BP1-GFP was kindly provided by Dr David J. Chen (University of Texas Southwestern) and has previously been described [10]. The cells were cultured in DMEM high-glucose medium with 10% fetal bovine serum, 100 U/ml penicillin and 100 mg/ml streptomycin, and were incubated at 37°C in an atmosphere of 95% air and 5% CO₂.

**Cell treatments**
For IR treatment, cells were irradiated by γ-rays from a 2.6 × 10⁵ Curie ⁶⁰Co source at Peking University, with total dose of 1 Gy (dose rate 0.94 Gy/min). For IR+HT treatments, the same IR treatment procedure was performed followed by incubation in a 41°C chamber that was equipped with a confocal microscope (Carl Zeiss LSM700) as previously described [11]. The temperature was set to 41°C, which had minimal effect on cell viability.

**RESULTS**
HT delayed and decreased the formation of the 53BP1 IRIF induced by γ-irradiation
To investigate how HT affected the cellular response to DSBs induced by γ-rays, we used HT1080 cells stably transfected with 53BP1-GFP. Because HT alone did not induce 53BP1 foci, as demonstrated in a previous report [12], we could study the effect of HT on the formation of the 53BP1 IRIF induced by γ-rays. Irradiated HT1080 cells were treated at 41°C and imaged simultaneously. Generally, within 5–15 min after exposure to IR, 53BP1 localized at discrete foci [13]. However, 20 min after γ-irradiation,
53BP1 IRIF could only be observed in a few cells under HT treatment. Each selected IRIF was tracked in situ from 20 min to several hours after treatments (Fig. 1). In this way, the dynamics of 53BP1 IRIF could be studied.

As shown in Fig. 2A, ~98.6% IRIF in the IR group appeared at 20 min post irradiation, while only 58.6% IRIF existed at that time in the IR+HT group. By 55 min after exposure to the γ-rays, all IRIF had emerged in the IR+HT group (Fig. 2A), indicating the delay in 53BP1 IRIF recruitment. In addition, the number of IRIF per cell treated with or without HT were compared. In the IR group, the number of IRIF reached a peak at 20 min (Fig. 2B). However, the peak time for the IR+HT group was ~40 min (Fig. 2B), also indicating the delay in formation of 53BP1 IRIF. Moreover, the average number of IRIF in the IR+HT group was significantly fewer than that in the IR group (10 ± 3/nucleus vs 24 ± 6/nucleus, P < 0.01, Fig. 2B). Taken together, the results demonstrated that HT could delay and decrease the formation of 53BP1 IRIF.

Characterization of 53BP1 IRIF in heated–irradiated cells by individual IRIF analysis

In order to further analyze individual IRIF, the maximum values of area and intensity for each IRIF were calculated. As shown in Fig. 3, no significant difference was detected between the IR group and the IR+HT group with respect to average area (14 ± 10 for the IR group and 13 ± 15 for IR+HT group, P = 0.75) or intensity (146 ± 57 for the IR group and 156 ± 50 for the IR+HT group, P = 0.06). Interestingly, in the IR+HT group, 26% of the IRIF did not disappear at the last observed time point in the present study, which was more than twice that in the IR group. Thus, the IRIF could be divided into two categories, ‘disappeared’ (i.e. repaired) IRIF and unrepaired IRIF. As shown in Fig. 3B, the recruitment of unrepaired IRIF was clearly delayed compared with that of disappeared IRIF. Additionally, the disappeared IRIF lost half their peak intensity within 80 min, while there was no obvious decrease in the intensity of the unrepaired IRIF over the period of observation (Fig. 3B). Furthermore, compared with the disappeared IRIF, a significantly larger area (28 ± 23 vs 8 ± 6, P < 0.01) and higher intensity (200 ± 15 vs 141 ± 49, P < 0.01) were observed in the unrepaired IRIF (Fig. 3C). Collectively, the results suggested that larger and brighter IRIF may be associated with the slower repair process.

It is of interest to note that almost every cell treated with HT had unrepaired IRIF. The spatiotemporal properties of each unrepaired IRIF were further analyzed in situ. Interestingly, we found that ~83% of these unrepaired IRIF gradually increased in size with time after exposure (Fig. 4A), which was reflective of the persistence of complex DSB lesion, while for the remaining IRIF, the apparent growth in IRIF area was due to the clustering of adjacent IRIF (Fig. 4B). As shown in Fig. 4B, the proximal IRIF clustered in the x, y and z directions.

DISCUSSION

Many studies have shown that HT has a great impact on the response of IR-induced DNA damage. Notably, the sensor proteins of the DDR are the direct targets of HT. Andrei Laszlo et al. reported that heat induced the delay of 53BP1 formation, and that this delay was modulated by Hsp 70 protein [9]. However, traditionally, DDR has been investigated using biochemical approaches that are based on measurements of populations of cells. Such measurements cannot reveal the detailed dynamics within a single cell. Also, the effect of HT on the spatiotemporal properties of sensor proteins is difficult to observe. In the present study, we used live-cell imaging to investigate the effect of HT on the spatiotemporal properties of 53BP1 IRIF induced by γ-rays. We found that HT inhibited the response for IR-induced DSBs in HT1080 cells, delaying 53BP1 IRIF formation and reducing the number of IRIF (Fig. 2). Moreover, HT-induced delay in 53BP1 complex formation could perturb the repair of DSBs [14], which would result in a higher probability of the persistence of unrejoined DSBs and/or the generation of misrejoined DSBs [15]. Furthermore, mild hyperthermia
Fig. 3. Analysis of single IRIF. (A) The correlation of maximum IRIF area and maximum IRIF intensity. Orange circles and green triangles represent single IRIF in the IR+HT group and the IR group, respectively. (B) The average intensity of unrepaired IRIF (red squares) and ‘disappeared’ (i.e. repaired) IRIF (blue dots) in the IR+HT group at indicated times. (C) The correlation of maximum IRIF intensity and maximum IRIF area in the IR+HT group. Red squares and blue circles represent single unrepaired IRIF and ‘disappeared’ (i.e. repaired) IRIF, respectively.

Fig. 4. Different ways to increase the area of unrepaired IRIF in the IR+HT group. (A) Representative time-lapse images show the growth in individual IRIF area in situ (white arrows). Bar = 5 μm. (B) Representative images of the merging of adjacent IRIF in the x-y plane (upper panel) and x-z plane (lower panel). White arrows show the clustering process of the proximal IRIF. Bar = 5 μm.
(41°C) efficiently induces degradation of BRCA2 and inhibits HR [8]. The inhibition of HR might stimulate the NHEJ or backup pathways of NHEJ [16–18] that was responsible for the error-prone repair [19]. Whether IR+HT causes more misrepaired DSBs in HT1080 cells is still unclear; therefore, further studies need to be performed by evaluating the induction of chromosome rearrangements.

Interestingly, through in situ tracking of individual IRIF from the IR+HT group, we found that HT resulted in more unrepaired IRIF over the period of observation compared with treatment with IR alone. Additionally, the unrepaired IRIF had a larger area and a higher intensity as well as a slower repair rate (Fig. 3). Previous studies have shown that larger IRIF may represent the machinery for slower repair of complex DSBs [20]. Moreover, larger IRIF are more persistent and are closely correlated with cell lethality [21].

Moreover, low-intensity foci are believed to represent simple DSBs, which are repaired quickly [22]. If the IRIF were always persistent in the nucleus, they can be viewed as ‘residual IRIF’. Several previous studies have reported a possible correlation between residual IRIF and cellular radiosensitivity [21, 23]. Furthermore, based on in situ tracking of individual IRIF, we found that the apparent change in area of unrepaired IRIF was mainly due to the area of individual IRIF increasing, rather than the clustering of proximal IRIF (Fig. 4). This could result in relocating of repair proteins to persisting lesions [24], or the expansion of chromatin in the vicinity of DSBs [25].

In summary, we found that HT could delay and decrease the formation of 53BP1 IRIF. Additionally, almost every heated cell contained a type of IRIF displaying larger area, higher intensity and slower repair rate compared with the unheated cells. Moreover, the majority of these IRIF had an apparent increase in area. Our results suggested that HT could perturb the primary event of the DDR induced by IR, and this may be responsible for the heated cells’ inability to repair DSBs.

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CONFLICT OF INTEREST

No potential conflicts of interest were disclosed.

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