Over past several years, the cold plasma-stimulated medium (PSM) has shown its remarkable anti-cancer capacity in par with the direct cold plasma irradiation on cancer cells or tumor tissues. Independent of the cold plasma device, PSM has noticeable advantage of being a flexible platform in cancer treatment. Currently, the largest disadvantage of PSM is its degradation during the storage over a wide temperature range. So far, to stabilize PSM, it must be remained frozen at $-80^\circ \text{C}$. In this study, we first reveal that the degradation of PSM is mainly due to the reaction between the reactive species and specific amino acids; mainly cysteine and methionine in medium. Based on this finding, both H$_2$O$_2$ in PSM and the anti-cancer capacity of PSM can be significantly stabilized during the storage at 8°C and $-25^\circ \text{C}$ for at least 3 days by using phosphate-buffered saline (PBS) and cysteine/methionine-free Dulbecco’s Modified Eagle Medium (DMEM). In addition, we demonstrate that adding a tyrosine derivative, 3-Nitro-L-tyrosine, into DMEM can mitigate the degradation of PSM at 8°C during 3 days of storage. This study provides a solid foundation for the future anti-cancer application of PSM.

Over the past decade, cold atmospheric plasma has shown a selective anti-cancer capacity both in vitro$^{1-8}$ and in vivo$^{9-13}$. Various type of cold plasma devices were used to directly irradiate cancer cells cultured in the multi-well plates$^{4,5}$, petri-dishes$^{6,7}$, or tumor tissues$^{8,12}$. Recently, plasma-stimulated medium (PSM) proved to exhibit a significant anti-cancer capacity as strong as the direct cold plasma treatment on glioblastoma cells$^{16-18}$, lung carcinoma cells$^{19}$, and bladder cancer cells$^{20}$. Very recently, it is reported that microsecond-pulsed plasma-activated media is able to selectively inhibit the growth of lung cancer (H460) cells rather than normal lung cancer (L132) cells$^{21}$. Another very recent study about the selective apoptosis in the PSM treated glioblastoma cells$^{22}$, further confirming that PSM is a selective anti-cancer tool. The injection of PSM into mice also significantly inhibits the growth of tumor$^{23}$. Thus, PSM may have wide application in cancer treatment including specific situations where cold atmospheric plasma cannot reach deep seated tumors or when the cold plasma device is not portable.

The cold plasma-originated reactive species are thought to be the main factor in cancer cell death and growth inhibition$^{24}$. In fact, the feasibility of using PSM to kill cancer cells is a direct evidence to support this conclusion. When cold plasma interacts with the medium both reactive oxygen species (ROS) such as hydroxyl free radicals (OH)$^{25}$ and hydrogen peroxide (H$_2$O$_2$)$^{19,26}$ and reactive nitrogen species (RNS) such as nitric oxide (NO)$^{27}$ and nitrite (NO$_2^{-}$)$^{17,28}$ are dissolved in the aqueous solution. Among them, H$_2$O$_2$ has been found to mainly contribute to the death of cancer cells after the direct cold plasma irradiation on cancer cells$^{19,31}$ or the indirect cold plasma irradiation on the culture medium$^{18,19}$.

To date, the degradation of PSM during the storage is its largest disadvantage for future clinical application. For pharmaceutical reasons, PSM should be stably stored for relative long time. However, as it is known PSM gradually loses its anti-cancer capacity during the storage at the room temperature$^{17,19}$ or down to the temperature a few degrees above the freezing point of water$^{19}$. Accordingly, the H$_2$O$_2$ concentration in PSM also gradually

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decreases during the storage under these conditions\textsuperscript{19}. Because the degradation of the anti-cancer capacity of PSM has been reported by different groups using different plasma devices and treatment doses\textsuperscript{17,19}, the instability of PSM during the storage can be regarded as a basic feature of PSM.

So far, the sole strategy to inhibit the degradation of PSM during the storage is freeze at an adequately low temperature. Tetsuo Adachi et al. reported that a freeze at $-80 \degree C$ was able to stabilize the anti-cancer capacity of cold plasma-stimulated Dulbecco's Modified Eagle Medium (DMEM) for a week\textsuperscript{19}. However, we found that freezing PSM at around $-20 \degree C$ would significantly accelerate the degradation of PSM\textsuperscript{17}. In fact, Tetsuo Adachi et al. also reported that the noticeable degradation of PSM still existed during the storage at $-30 \degree C$\textsuperscript{19}. These observations indicate that the degradation of PSM during the storage may be due to a slow temperature-dependent reaction which can be inhibited in an adequately cold atmosphere. Nonetheless, according to the description of most manufacturers of cell culture media, the ideal storage temperature range for medium is over 2 \degree C to 8 \degree C, rather than the freezing condition. Thus, considering the clinical application prospective, PSM at least should be stable at such a temperature range. So far, no method has been reported about improving the stability of PSM over 2 \degree C to 8 \degree C.

In this study, by comparing the $H_2O_2$ concentration in the cold plasma-stimulated phosphate-buffered saline (PBS) and the cold plasma-stimulated DMEM during the storage at 8 \degree C, 22 \degree C and $-25 \degree C$, we found that the degradation of PSM was mainly due to the reaction between the plasma-originated reactive species and components in DMEM. A further detailed investigation showed that cysteine and methioine were the main components in DMEM contributing to this process. Based on these principles, we further proved that the plasma-stimulated cysteine/methionine-free DMEM was much more stable than the plasma-stimulated standard DMEM during the storage at 8 \degree C, 22 \degree C and $-25 \degree C$. In addition, adding a tyrosine derivative, 3-Nitro-L-tyrosine, could also improve the stability of cold plasma-stimulated DMEM at 8 \degree C but not at $-25 \degree C$.

**Results**

The cold plasma device (Fig. 1a) was a typical helium cold atmospheric plasma jet generator which has been used in a series of studies\textsuperscript{17,18,32-34}. Cold plasma jet was generated between the central electrode and the grounded ring electrode. The carrying gas was helium with a flow of 4.7 L/min and was controlled by a flow meter. The output voltage was 3.16 kV. The general research strategy is illustrated in Fig. 1b. 1 mL of PSM was made by the vertical irradiation on the medium in a well on a 12-well plate. Then, PSM was transferred into a 1.5 mL of centrifuge tube by pipette. The centrifuge tubes were stored in refrigerators with different internal temperatures (8 \degree C and $-25 \degree C$) for 3 days. Similarly, the centrifuge tubes were stored at 22 \degree C when we investigated the degradation of PSM at the room temperature (not shown in Fig. 1b). Ultimately, PSM was either transferred to culture the seeded cancer cells in a 96-well plate or transferred into a black-wall 96-well plate to perform the measurement for $H_2O_2$. The detailed protocols for the transfer are illustrated in Methods and Fig. S1a,b. Because PBS is not suitable for cell culture, the protocols used to transfer the plasma-stimulated PBS is slightly different from the protocols used to transfer the plasma-stimulated DMEM (Fig. S1).

To understand the mechanism of PSM degradation, we first compared the $H_2O_2$ generation in the cold plasma-stimulated DMEM and PBS after 26 hours of storage and found that the $H_2O_2$ in plasma-stimulated PBS was quite stable during the storage while most $H_2O_2$ in the plasma-stimulated DMEM had been lost after 26 hours at 8 \degree C and 22 \degree C (Fig. 2a). The higher storage temperature for PSM, the stronger degradation occurs in PSM (Fig. 2a). Obviously, the instability of PSM is mainly due to the reaction between component of DMEM and reactive species generated by the cold plasma. Recently, we have demonstrated that the $H_2O_2$ in PSM tended to react with cysteine among all 20 amino acids after the plasma irradiation\textsuperscript{18}. Thus, the plasma-stimulated PBS has higher $H_2O_2$ concentration than the plasma-stimulated DMEM (Fig. 2a). We further found that the $H_2O_2$ concentration in the plasma-stimulated PBS was not only stable over 7 days of storage at 8 \degree C (Fig. 2b) but also stable over 3 days at $-25 \degree C$ (Fig. 2c). The stable anti-cancer capacity of the plasma-stimulated PBS over 3 days
at 8 °C and −25 °C was further confirmed by using the corresponding treated PBS to affect the growth of breast cancer (MDA-MB-231) cells (Fig. 2d). In addition, we demonstrated that the plasma-stimulated PBS could also effectively inhibit the growth of glioblastoma (U87MG) cells and pancreatic cancer (PA-TU-8988T) cells with a

Figure 2. The plasma-stimulated PBS is a stable and effective anti-cancer tool. (a) The change of H$_2$O$_2$ concentration in the cold plasma-stimulated DMEM and PBS during the storage at 8 °C and 22 °C for 26 hours. (b) The change of H$_2$O$_2$ concentration in the cold plasma-stimulated PBS during the storage at 8 °C for 7 days. (c) The change of H$_2$O$_2$ concentration in the cold plasma-stimulated PBS during the storage at 8 °C and −25 °C for 3 days. (d) The change of anti-cancer capacity of the cold plasma-stimulated PBS during the storage at 8 °C and −25 °C for 3 days. (e) The effect of the cold plasma-stimulated PBS on the cell viability of U87MG, PA-TU-8988T, and MDA-MB-231 cells cultured in the cold plasma-stimulated PBS. (f) The change of H$_2$O$_2$ concentration in 37.3 μM H$_2$O$_2$-containing PBS during the storage at 8 °C for 3 days. The volume of solution for each well was 1 mL for all experiments. The treatment time for (a–d) was 1 min and 2 min, respectively. Results are presented as the mean ± s.d. of three independently repeated experiments performed in triplicate (a–c,f) or in sextuplicate (d,e). For the cell viability, the data have been normalized to corresponding control group. Student’s t-test was performed and the significance is indicated as ***p < 0.005.
dose-dependent manner (Fig. 2e), making it a stable anti-cancer PSM over a wide temperature range. Ultimately, we tested whether the H$_2$O$_2$-containing PBS would be as stable as the cold plasma-stimulated PBS, and found that the H$_2$O$_2$-containing PBS just experienced a slight degradation over the storage at 8 °C for 3 days (Fig. 2f), which is similar to the slight degradation observed in the plasma-stimulated PBS (Fig. 2c). The H$_2$O$_2$ in PBS will experience a natural degradation during the storage at 8 °C. Thus, the gradual H$_2$O$_2$ consumption in the cold plasma-stimulated DMEM is due to the combined effect of the natural degradation of H$_2$O$_2$ in aqueous solution and the reaction between H$_2$O$_2$ and the component in DMEM. The latter mainly controls the degradation at 8 °C.

Furthermore, the effect of specific components in DMEM on the PSM degradation was comprehensively investigated. A comparison between DMEM and PBS reveals that 15 amino acids, glucose, calcium ion, magnesium ion, and phenol red are the main composition difference between DMEM and PBS (Fig. S2). Thus, 15 specific amino acids solutions, 1 glucose solution, 2 saline solutions (CaCl$_2$, MgCl$_2$), and 1 phenol red solution were respectively made by adding or dissolving specific component in PBS according to their concentrations in the standard DMEM (Fig. S3). After the plasma treatment and subsequent 26 hours of storage at 22 °C, the H$_2$O$_2$ concentrations in these solutions were measured and compared with corresponding H$_2$O$_2$ concentration in the solution just after the cold plasma treatment (Fig. 3a). The H$_2$O$_2$ in the plasma-stimulated PBS is quite

Figure 3. Cysteine and methionine mainly contribute to the degradation of the plasma-stimulated DMEM at 22 °C and 8 °C. (a) The change of H$_2$O$_2$ concentration in the cold plasma-stimulated PBS containing specific component in DMEM during the storage at 22 °C for 26 hours. (b) The change of H$_2$O$_2$ concentration in the cold plasma-stimulated cysteine/methionine-free DMEM, cysteine-free DMEM, methionine-free DMEM, and standard DMEM during the storage at 8 °C for 3 days. (c) The change of anti-cancer capacity of the cold plasma-stimulated cysteine/methionine-free DMEM, cysteine-free DMEM, methionine-free DMEM, and standard DMEM during the storage at 8 °C for 3 days. For all experiments, the volume of solution and the treatment time in each well was 1 mL and 1 min, respectively. Results are presented as the mean ± s.d. of three independently repeated experiments performed in triplicate (a,b) or in sextuplicate (c). For the cell viability, the data have been normalized to corresponding control group. Student's t-test was performed and the significance is indicated as *p < 0.05, **p < 0.01, ***p < 0.005.
stable over the storage. In contrast, most of the studied components cause different levels of $H_2O_2$ degradation in the plasma-stimulated PBS-based solution, except tryptophan, tyrosine, and lysine. Cysteine and methionine cause the most significant degradation of $H_2O_2$ during the storage at room temperature. Phenylalanine also cause the noticeably degradation of $H_2O_2$. Based on these trends, it is reasonable to predict that a modified DMEM without cysteine, methionine, phenylalanine or other components will be a stable PSM. So far, however, the modified DMEM without phenylalanine is not available on the market. Thus, we just focused on studying the role of cysteine and methionine on the PSM degradation. We first compared the plasma-stimulated cysteine/methionine/glutamine-free DMEM with the plasma-stimulated arginine/lysine/glutamine-free DMEM and found that only the former was stable during the storage at 22 °C for 26 hours (Fig. S4). The absence of arginine and lysine did not inhibit the $H_2O_2$ degradation in PSM (Fig. S4). These results are consistent with the trends revealed in Fig. 3a.

To make cysteine/methionine-free DMEM, 4 mM glutamine was added in the purchased cysteine/methionine/glutamine-free DMEM. Based on these cysteine/methionine-free DMEM, we further compared the modified DMEM just without cysteine and the modified DMEM just without methionine according to the description in Methods. Then, the specific effect of cysteine and methionine on the instability of PSM was investigated. After three days of storage at 8 °C, only the cold plasma-stimulated cysteine/methionine-free DMEM shows strong capacity inhibiting the $H_2O_2$ degradation (Fig. 3b). For cysteine and methionine, either of them will result in noticeable $H_2O_2$ degradation in PSM during the storage (Fig. 3b). Furthermore, the anti-cancer capacity of these PSM on breast cancer (MDA-MB-231) cells was investigated. In contrast to the experiments performed in the immediately treated medium, 3 days of storage at 8 °C causes noticeable degradation of the anti-cancer capacity of PSM even the modified DMEM does not contain cysteine and methionine (Fig. 3c). However, compared with other three cases, the cysteine/methionine-free DMEM still shows the strongest capacity to retain the effective species in PSM. The cysteine/methionine-free DMEM does not completely inhibit the $H_2O_2$ degradation, which may be due to two reasons. One is that many components other than cysteine and methionine in DME also contribute to the instability of PSM during the storage (Fig. 3a). The modified DMEM without all these reactive components may be as stable as PBS after the plasma irradiation. Another possibility is that some reactive species other than $H_2O_2$ may also contribute to the anti-cancer capacity of PSM.

Despite the optimized storage temperature for the culture medium should be between 2 °C to 8 °C, the cryopreservation of PSM may still be necessary at some special circumstances. For most of refrigerators served in pharmacies, the temperature in the freezing chamber is around ~20 °C. To date, however, PSM has been found even more degradable when stored at temperature around ~20°C than that at 22 °C or 4 °C. General observation is that freezing aggravates the degradation of PSM. As we demonstrated above, both the $H_2O_2$ concentration (Fig. 2c) and the anti-cancer capacity (Fig. 2d) of the cold plasma-stimulated PBS are stably retained during the storage at ~25 °C. Thus, the degradation of the plasma-stimulated DMEM should also be due to the reaction between component in DMEM and the plasma-originated reactive species such as $H_2O_2$. To verify this proposition, we first investigated the stability of $H_2O_2$-containing DMEM and $H_2O_2$-containing PBS under the freezing conditions for 3 days. It is found that, in contrast to the immediately prepared $H_2O_2$-containing DMEM and $H_2O_2$-containing PBS, $H_2O_2$ has been completely consumed (Fig. 4a) and has been well retained (Fig. 4b) in the frozen DMEM and the frozen PBS at ~25 °C, respectively. Thus, the plasma-originated or other sources-originated $H_2O_2$ naturally tend to strongly react with specific component in DMEM under the freezing conditions. In addition, similar to the trend shown in Fig. 2f, $H_2O_2$ also shows a natural tend to be slightly consumed in PBS during the freezing storage. Thus, the degradation of $H_2O_2$ in PSM under the freezing condition is also due to the reaction between $H_2O_2$ and the components in media and a natural degradation in aqueous solution. The former is the main factor.

Similar to the research strategy described above, we further investigated which component in DME contribute to the degradation of PSM during the freezing storage. This was done by comparing the degradation of $H_2O_2$ in 15 amino acids solutions, 1 glucose solution, 2 saline solutions (CaCl$_2$, MgCl$_2$), 1 phenol red solution, and PBS. These solutions were made by previously mentioned methods according to the concentration shown in Fig. S3. As presented in Fig. 4c, cysteine and methionine are still the two most reactive components in DME consuming the plasma-originated $H_2O_2$ during the freezing storage. Different from the trend shown in Fig. 3a, all studied components tend to aggravate the degradation of $H_2O_2$ in the cold plasma-stimulated PBS at ~25 °C, including tyrosine, tryptophan, and lysine, which in fact inhibit the degradation of $H_2O_2$ in the cold plasma-stimulated PBS at 8 °C (Fig. 3a). Under the freezing conditions, the $H_2O_2$ degradation in the methionine solution is stronger than that in the cysteine solution. Furthermore, we investigated the anti-degradation effect of the cysteine/methionine-free DME under the freezing conditions using the above mentioned methods. The appearance of either of cysteine and methionine in DME will result in the complete consumption of $H_2O_2$ after 3 days of storage at ~25 °C (Fig. 4d). The cysteine-free DME is slightly more resistant to the $H_2O_2$ degradation than the methionine-free DME, which is consistent with the trend revealed in Fig. 4c that methionine is more reactive with $H_2O_2$ than cysteine during the freezing storage. For the cysteine/methionine-free DME, more than half of $H_2O_2$ has been consumed during the 3 days of storage at ~25 °C, which is also consisted with trend shown in Fig. 4c that many other components in DME also contribute to the instability of PSM. Ultimately, we compared the anti-cancer capacity of these modified DME on MDA-MB-231 cells. When the plasma treatment time is just 1 min (Fig. 4e), the difference between four experimental groups is not obvious. Such phenomenon may be due to the low concentration of residual $H_2O_2$ or reactive species in frozen plasma-stimulated cysteine/methionine-free DME, which are too weak to cause noticeable anti-cancer effect (Fig. 4d). When the plasma treatment time extends to 2 min (Fig. 4e), the appearance of cysteine or methionine will significantly aggravate the degradation of the anti-cancer capacity of PSM. In contrast, the plasma-stimulated cysteine/methionine-free DME still causes a 60% of anti-cancer growth effect.
Figure 4. Cysteine and methionine mainly contribute to the degradation of the plasma-stimulated DMEM under the freezing condition. (a) The change of H$_2$O$_2$ concentration in 34.5 μM H$_2$O$_2$-containing DMEM during the storage at −25 °C for 3 days. (b) The change of H$_2$O$_2$ concentration in 37.3 μM H$_2$O$_2$-containing PBS during the storage at −25 °C for 3 days. (c) The change of H$_2$O$_2$ concentration in the cold plasma-stimulated PBS containing specific component in DMEM during the storage at −25 °C for 26 hours. (d) The change of H$_2$O$_2$ concentration in the cold plasma-stimulated cysteine/methionine-free DMEM, cysteine-free DMEM, methionine-free DMEM, and standard DMEM during the storage at −25 °C for 3 days. (e) The anti-cancer capacity of the cold plasma-stimulated cysteine/methionine-free DMEM, cysteine-free DMEM, methionine-free DMEM, and standard DMEM during the storage at −25 °C for 3 days. For all experiments, the volume of solution in each well was 1 mL. The treatment time was 1 min for (c,d). Results are presented as the mean ± s.d. of three independently repeated experiments performed in triplicate (a–d) or in sextuplicate (e). For the cell viabiity, the data have been normalized to corresponding control group. Student’s t-test was performed and the significance is indicated as ***p < 0.005.
In addition to above mentioned methods, we presented a third anti-degradation method based on the results shown in Fig. 3a. It is clear that tyrosine is able to slightly increase the H₂O₂ concentration in the plasma-stimulated PBS during the storage at 8°C. Among 15 amino acids in DMEM, only tyrosine and phenylalanine have aromatic ring structures (Fig. 5a). However, phenylalanine causes noticeable H₂O₂ degradation in the cold plasma-stimulated PBS during the storage at 8°C (Fig. 3a). The only difference between tyrosine and phenylalanine is a hydroxyl on the aromatic ring (Fig. 5a), indicating that the modification on the aromatic ring of amino acids may endow the amino acids with a capacity inhibiting the degradation of H₂O₂ during the storage at 8°C. We utilized a tyrosine derivative, 3-Nitro-L-tyrosine, which had a nitro group on the aromatic ring of tyrosine (Fig. 5a). 2 mM 3-Nitro-L-tyrosine-containing DMEM, 2 mM phenylalanine-containing DMEM, and 2 mM tyrosine-containing DMEM were prepared by dissolving purchased 3-Nitro-L-tyrosine, phenylalanine, and tyrosine powders in the standard DMEM, respectively. Subsequently, the stability of H₂O₂ in the plasma-stimulated DMEM, tyrosine-containing DMEM, phenylalanine-containing DMEM, and 3-Nitro-L-tyrosine-containing DMEM stored at 8°C were compared. 3-Nitro-L-tyrosine significantly inhibits the degradation of H₂O₂ in PSM (Fig. 5b). On the contrary, tyrosine and phenylalanine do not improve the stability of PSM (Fig. 5b). Furthermore, it is found that noticeable H₂O₂ degradation still occur in the cold plasma-stimulated 3-Nitro-L-tyrosine-containing DMEM and the cold plasma-stimulated standard DMEM after the storage at 8°C for 3 days (Fig. 5c). Despite 60% of plasma-originated H₂O₂ has been lost during the storage, it is more effective than the plasma-stimulated standard DMEM, in which 74% of plasma-originated H₂O₂ has been lost during the 3 days of storage. Nonetheless, after the storage, the cold plasma-stimulated 3-Nitro-L-tyrosine-containing DMEM still shows a noticeable anti-breast cancer cells capacity as strong as the immediately treated 3-Nitro-L-tyrosine-containing DMEM (Fig. 5d). In contrast, the anti-cancer capacity of the plasma-stimulated standard DMEM significantly decreases during the freezing storage.

To reveal the anti-degradation mechanism of 3-Nitro-L-tyrosine in the plasma-stimulated DMEM, we first investigated whether 3-Nitro-L-tyrosine was able to inhibit the H₂O₂ degradation in 0.2 mM cysteine-containing PBS, 0.2 mM methionine-containing PBS, and 0.4 mM phenylalanine-containing PBS. As shown in Fig. 5e, 2 mM 3-Nitro-L-tyrosine is able to completely inhibit the H₂O₂ degradation in all three cold plasma-stimulated amino acids-containing PBS solutions at 22°C. Thus, the cold plasma-stimulated DMEM may be stabilized by 3-Nitro-L-tyrosine through inhibiting the reaction between some components in DMEM such as cysteine, methionine, or phenylalanine with the cold plasma-originated H₂O₂. Furthermore, we studied the concentration-effect of 3-Nitro-L-tyrosine on its anti-degradation effect in the plasma-stimulated DMEM. For the plasma-stimulated 3-Nitro-L-tyrosine-containing DMEM, the residual H₂O₂ after 3 days of storage increases as the concentration of 3-Nitro-L-tyrosine increases from 1 mM to 4 mM (Fig. 5f). After 3 days of storage at 8°C, 4 mM 3-Nitro-L-tyrosine-containing DMEM loses 30% of the H₂O₂ originated from the plasma treatment. In contrast, the standard DMEM loses 81% of the plasma-originated H₂O₂ (Fig. 5f). Finally, we identified that the degradation of H₂O₂ in the 3-Nitro-L-tyrosine-containing DMEM during the freezing storage is as extensive as that occurs in the standard DMEM after the plasma treatment (Fig. 5g). In short, adding 3-Nitro-L-tyrosine in DMEM weakens the degradation of PSM stored at 8°C, though it cannot weaken the degradation of PSM stored at −25°C.

Discussion
Understanding the interaction between the cold atmospheric plasma and molecules in an aqueous solution builds the foundation of the plasma medicine. Current experimentation is mainly limited to studying chemical reactions between the cold plasma-originated reactive species and components in an aqueous solutions such as cell culture medium. Among diverse chemicals, amino acids are not only the building blocks of all proteins involving in nearly all biological processes but also the main component in most cell culture media utilized in the studies in vitro. The interaction between the cold plasma and the amino acids in culture media has been extensively investigated in past two years. A general investigation regarding the chemical modification and change of 20 amino acids after the cold plasma treatment has been performed via high-resolution mass spectrometry. It was found that hydroxylation, nitration, sulfonation, disulfide formation and sulfoxidation were the main chemical modification of 14 amino acids upon the cold plasma treatment. In addition, sulfur-containing amino acids such as cysteine and methionine and aromatic amino acids such as tryptophan, phenylalanine, and tyrosine were preferentially consumed during the plasma treatment. It was also demonstrated that proteins in FBS consume the reactive species generated by cold plasma treatment significantly. Recently, we further demonstrated that among 20 amino acids, cysteine was most reactive to plasma produced species including H₂O₂. These results clearly revealed that the plasma-originated reactive species will be partially consumed immediately after their formation in PSM. Furthermore, in this study, we proved that many components of medium particularly cysteine and methionine will slowly consume the plasma-originated reactive species such as H₂O₂ during storage. Thus, in designing the ideal PSM, cysteine and methionine should be avoided. Such slow reaction between reactive species and components in medium should be temperature-dependent, because it can be completely inhibited through an adequately low temperature freeze at −80°C.

The discovery of the anti-degradation property of 3-Nitro-L-tyrosine provides a promising strategy to improve the stability of PSM because the original component of DMEM does not need to be removed. Despite the fact that the chemical mechanism of such anti-degradation capacity is unknown yet, adding modifications to the aromatic ring of tyrosine or phenylalanine may be the general strategy to obtain new derivatives capable to preserve H₂O₂ and perhaps other plasma produced reactive species. In addition, the anti-degradation effect of 3-Nitro-L-tyrosine first reveals that the interaction between the cold plasma and media cannot be simply described as the consumption of plasma-originated reactive species such as H₂O₂ or the modification of component such as cysteine and methionine in the medium. Indeed, most of components in the medium tend to consume the plasma-originated reactive species; however, specific amino acid derivatives are able to inhibit the
Figure 5. 3-Nitro-L-tyrosine inhibits the degradation of cold plasma-stimulated DMEM. (a) Chemical formulas of phenylalanine, tyrosine, and 3-Nitro-L-tyrosine. (b) The change of \( \text{H}_2\text{O}_2 \) concentration in the cold plasma-stimulated standard DMEM, phenylalanine-containing DMEM, tyrosine-containing DMEM, and 3-Nitro-L-tyrosine-containing DMEM during the storage at 8°C for 26 hours. (c) The change of \( \text{H}_2\text{O}_2 \) concentration in the cold plasma-stimulated standard DMEM and 3-Nitro-L-tyrosine-containing DMEM during the storage at 8°C for 3 days. (d) The change of anti-cancer capacity of the cold plasma-stimulated
standard DMEM and 3-Nitro-L-tyrosine-containing DMEM during the storage at 8 °C for 3 days. (e) The change of H₂O₂ concentration in the cold plasma-stimulated single amino acids-containing PBS and double amino acids-containing PBS during the storage at 22 °C for 26 hours. Corresponding concentration of cysteine, methionine, phenylalanine, and 3-Nitro-L-tyrosine in PBS were 0.2 mM, 0.2 mM, 0.4 mM and 2 mM, respectively. (f) The change of H₂O₂ concentration in the cold plasma-stimulated 3-Nitro-L-tyrosine-containing DMEM with different concentrations during the storage at 8 °C for 3 days. (g) The change of H₂O₂ concentration in the cold plasma-stimulated standard DMEM and 3-Nitro-L-tyrosine-containing DMEM during the storage at −25 °C for 3 days. For all experiments, the volume of solution in each well was 1 mL. The treatment time was 1 min for (b,c,f) and was 2 min for (c,d,g), respectively. Results are presented as the mean ± s.d. of three independently repeated experiments performed in triplicate (b,c,e–g) or in sextuplicate (d). For the cell viability, the data have been normalized to corresponding control group. Student’s t-test was performed and the significance is indicated as *p < 0.05, **p < 0.01, ***p < 0.005.

reaction between plasma-originated reactive species and specific components in the medium such as cysteine and methionine. These results demonstrate that the chemical essence of interaction between cold plasma and aqueous solution is far from clear. We not only need to measure the immediate reaction in media due to the plasma irradiation but we also have to monitor the long term dynamic reactions in PSM after the plasma irradiation.

The essence of the anti-cancer reactive species in cold plasma or PSM is still disputable. A simple comparison between the concentration of H₂O₂ (Fig. 3b) and the anti-cancer effect (Fig. 3c) in the plasma-stimulated cysteine/methionine-free DMEM stored at 8 °C for 3 days reveals that though H₂O₂ just loses about 23% during the storage, corresponding anti-cancer capacity loses about 45% during the same storage. Perhaps other plasma-originated reactive species also contributes to the anti-cancer effect of PSM and decay faster than H₂O₂. Currently, H₂O₂ has just been regarded as the main plasma-originated reactive species contributing to the anti-cancer capacity of cold plasma²⁹,³⁰. However, evidence exist that both cellular-level¹⁸,²⁶,²⁷ and molecular-level²⁹,³⁶ response of cancer cells to the cold plasma treatment are different from that positive control using H₂O₂. The cold plasma-originated atomic oxygen has been proven to be a main toxic reactive species to epithelial kidney cells under a specific plasma treatment condition²⁷. So far, based on the accumulated data in the literature, it is premature to reach final conclusion regarding the mechanism of action of cold atmospheric plasma. A further comprehensive investigation on the role of key plasma-originated reactive species on the anti-cancer effect of cold plasma is necessary and urgent.

Conclusions
The degradation of PSM during storage is due to the reaction between the plasma-originated reactive species and components of the cell culture medium. We have found that cysteine and methionine are two most reactive components in DMEM contributing to the consumption of plasma-originated H₂O₂. The cold plasma-stimulated PBS and the cold plasma-stimulated cysteine/methionine-free DMEM can be stably stored at 8 °C and −25 °C for at least 3 days. In addition, adding 3-Nitro-L-tyrosine into DMEM could weaken the degradation of the cold plasma-stimulated DMEM at 8 °C for 3 days. These stabilizing strategies build a solid foundation for the application of PSM in future cancer treatment.

Methods
Medium and cell cultures. Standard Dulbecco’s modified Eagle’s medium (11965-118), the modified cysteine/methionine/glutamine-free DMEM (21013-024), the modified arginine/lysine/glutamine-free DMEM (A14431-01) and PBS (14040-133) were purchased from Gibco (Life Technologies). In addition, all DMEM and PBS were mixed with 1% (v/v) antibiotic (penicillin and streptomycin) (Life Technologies) before any experiments. Human glioblastoma (U87MG) cells and pancreatic cancer (PA-TU-8898T) cells were provided by Dr. Murad’s Lab at the George Washington University. Human breast cancer (MDA-MB-231) cells were provided by Dr. Zhang’s Lab at the George Washington University. All cancer cell lines were seeded with a confluence of 3 × 10⁴ cells/mL and a volume of 100 μL in each well on 96-well plate and were cultured for 6 hours in a complete media composed of Dulbecco’s modified Eagle’s medium (Life Technologies) supplemented with 10% (v/v) fetal bovine serum (ThermoFisher Scientific) and 1% (v/v) antibiotic (penicillin and streptomycin) solution (Life Technologies) under the standard cell culture conditions (a humidified, 37 °C, 5% CO₂ environment). In each experiment, 6 wells in a single column on 96-well plate would be seeded with cancer cells.

Making plasma-stimulated PBS and DMEM. In this study, the protocols to make PSM are the same among different experiments. 1 mL of PBS or DMEM or other modified DMEM such as cysteine/methionine-free DMEM in a well on 12-well plate (Falcon) was treated by cold plasma for 1 min or 2 min. The gap between the end of dielectric plasma tube and the bottom of 12-well plate was 3 cm.

Making H₂O₂-containing PBS and DMEM. The H₂O₂-containing PBS and H₂O₂-containing DMEM were prepared by adding purchased 30 wt% H₂O₂ solution (Sigma-Aldrich) into PBS and DMEM, respectively. The H₂O₂ concentration in H₂O₂-containing PBS and H₂O₂-containing DMEM were close to the H₂O₂ concentration in the 1 min of cold plasma-treated 1 mL of PBS or DMEM in 12-well plate, respectively.

Making the specific component-containing PBS and DMEM. The amino acids-containing PBS, the calcium-containing PBS, the magnesium-containing PBS, the glucose-containing PBS, the phenol red-containing PBS, and amino acids-containing DMEM were prepared by adding and dissolving purchased specific amino
 acids, calcium chloride solution, magnesium chloride solution, D-glucose and phenol red solution or amino acids derivative such as 3-Nitro-L-tyrosine into PBS and DMEM, respectively. Except PBS and DMEM, all above chemicals were purchased from Sigma-Aldrich.

Affecting the growth of cancer cells seeded in 96-well plate by the cold plasma-stimulated DMEM or PBS. First, the initial culture medium which has been cultured cells for 6 hours has been removed before this step. Then, for the plasma-stimulated DMEM (Fig. S1a), 100 μL of treated DMEM was transferred from a well on 12-well plate or other tubes stored in refrigerator into a well on 96-well plate, in which 3 × 10^3 cancer cells were seeded and has been cultured for 6 hours. In each experiment, 6 wells in a single column on 96-well plate have been seeded with cancer cells. In the control group, the DMEM used to culture cancer cells was the untreated DMEM. For the plasma-stimulated PBS (Fig. S1b), considering that cancer cells cannot grow in PBS over a long time range, 100 μL of untreated DMEM was first transferred into the well seeded with 3 × 10^3 cancer cells on 96-well plate. Then, 100 μL of treated PBS was transferred into the well which contained 100 μL of untreated DMEM and 3 × 10^3 cancer cells. Thus, after this step, the total volume of the mixed medium in each well seeded with cancer cells were 200 μL. For the control group, the PBS transferred into the well would not be treated. Ultimately, for above two cases, the cancer cells on 96-well plate were cultured under a standard cell culture condition (a humidified, 37 °C, 5% CO_2 environment) for 3 days.

Measuring and processing the cell viability. After the experiments, the cancer cells in 96-well plates were cultured in 96-well plate under the standard cell culture conditions (a humidified, 37 °C, 5% CO_2 environment) for 3 days. Then, according to the standard two-steps MTT measurement protocols, the viability of cancer cells was qualified by MTT test and was read by an H1 microplate reader (Hybrid Technology) at the absorbance of 570 nm. In each independent experiment, 6 wells in a single column on 96-well plate would be seeded with cancer cells and would be taken cell viability test. The average value of measured cell viability from 6 wells was regarded as the cell viability measured from one independent experiment. To facilitate the understanding on data, all data about cell viability shown in figures has been normalized to the control group by dividing the measured cell viability of experimental group by the measured cell viability of control group. The final data shown in all figures are the mean ± s.d. of the normalized cell viability from three independently repeated experiments.

Measuring H_2O_2 concentration. 50 μL of medium or PBS which needs to be measured was transferred to a well on a black-well 96-well clear bottom plate (Falcon) in triplicate. Next, according to the standard protocols provided by Sigma-Aldrich, the H_2O_2 concentration in the cold plasma-stimulated medium was measured. The fluorescence at 540/590 nm was read using an H1 microplate reader (Hybrid Technology).

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