How can cancer cells use hydrogen peroxide to strive?

Hydrogen peroxide (H$_2$O$_2$) is a well-known antiseptic used in our homes since the 1920s. But did you know that our cells produce this molecule and that at low concentrations H$_2$O$_2$ instructs our cells to do different things, acting as a signaling molecule? In fact, H$_2$O$_2$ may oxidize several cellular components including proteins, altering their function. Healthy cells use H$_2$O$_2$ to regulate several of their behaviors, including proliferation, migration, or even cell-to-cell communication. It is, therefore, not surprising that cancer cells produce higher amounts of this reactive oxygen species and take advantage of this signaling feature of H$_2$O$_2$ to increase their proliferation and migration.

Fig. 1. RPSA oxidation by H$_2$O$_2$ results in the formation of an intramolecular disulphide (a chemical bond that occurs between two cysteine residues). This leads to increased clustering at the cell membrane level and recruitment of particular integrins to RPSA clusters (e.g. integrin ?1 subunit but not integrin ?5 subunit), where they are activated. This recruitment selectivity modulates cell adhesion properties such as alterations in binding specificities to different extracellular matrices. This might be particularly important for different events in tumor progression that depend on cell adhesion processes, such as tumor cell intravasation and extravasation leading to metastasis, conferring a selective advantage to RPSA-overexpressing cells in an oxidative environment.
Although the knowledge around this subject has been increasing in the past few years, it is still largely unknown which molecules are oxidized by H$_2$O$_2$ and how this oxidation affects their function. In our recent work, we have found that a protein called Ribosomal Protein SA (RPSA) is oxidized by H$_2$O$_2$ and that this oxidation enhances cancer properties. In fact, RPSA oxidation improves cell adhesion efficiency to the extracellular matrix and promotes cell extravasation (the passing of cells from the blood stream to surrounding tissues, a critical step in the metastatic process). The effect on cell adhesion is most likely mediated by RPSA clusters that accumulate at the cell membrane level and contain specific adhesion molecules called integrins. Interestingly, it appears that different adhesion molecules are present on those clusters depending on the RPSA redox state, suggesting that RPSA oxidation may influence the type of extracellular matrix to which cells adhere.

Our results unravel a new mechanism by which H$_2$O$_2$ modulates the cell adhesion properties and identify RPSA as the H$_2$O$_2$ sensor in this process. Elevated levels of H$_2$O$_2$ produced both in primary tumors and secondary metastatic sites, together with RPSA up-regulation, might confer a selective advantage to tumor cells in the metastatic process by contributing to alterations in cell adhesion properties such as binding specificity. The identification of redox-regulated proteins, such as RPSA and the determination of their relevance in tumor development will be essential to design more specific and efficient redox-based therapies for cancer.

_Filipe Vilas-Boas and Carla Real_
_Centro de Química e Bioquímica, Faculdade de Ciências, Universidade de Lisboa_
_Lisboa, Portugal_

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