Does Cellular Hydrogen Peroxide Diffuse or Act Locally?

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Abstract

Understanding of redox signaling requires data on the spatiotemporal distribution of hydrogen peroxide (H₂O₂) within the cell. The fluorescent reporter HyPer is a powerful instrument for H₂O₂ imaging. However, rapid diffusion of HyPer throughout the nucleocytoplasmic compartment does not allow visualization of H₂O₂ gradients on the micrometer scale. Here we dramatically improved the spatial resolution of H₂O₂ imaging by applying subcytoplasmic targeting of HyPer. The membrane-attached reporters identified “microdomains” of elevated H₂O₂ levels within the cytoplasm of the cells exposed to growth factors. We demonstrate that diffusion of H₂O₂ across the cytoplasm was strongly limited, providing evidence that H₂O₂ acts locally inside cells. Antioxid. Redox Signal. 14, 1–7.

Introduction

Cells continuously produce reactive oxygen species (ROS). Among ROS, superoxide anion radicals, hydroxyl radicals, hydrogen peroxide (H₂O₂), and nitric oxide are the most investigated species in biology (7). ROS are responsible for oxidative damage of DNA, proteins, and lipids and are associated with disease states such as inflammation, cancer, and neurodegenerative diseases, among others. Physiological processes, such as aging, differentiation, cell motility, and others, are known to be significantly associated with intracellular ROS production.

A decade ago, H₂O₂ was positively identified as a messenger molecule acting through selective modification of thiol groups in proteins regulating signal transduction from growth factor and cytokine receptors (16, 20). H₂O₂ is not a radical; therefore, it is less reactive and, hence, much more stable than other ROS (21). H₂O₂ is small enough to travel over long distances within or even between cells. Although the degree of permeability of biomembranes for H₂O₂ is still under debate (4), properties of H₂O₂ such as its transient appearance make this molecule a good candidate for being a “messenger.” However, lack of data on spatiotemporal mapping of H₂O₂ at the level of cellular subcompartments significantly challenges the understanding of its role and regulation in signaling and disease.

Intracellular H₂O₂ production can be efficiently visualized with fluorescent probes, based on either small molecules (14) or genetically encoded constructs (3, 10). The first provides the advantages of simplicity and high signal-to-background ratio, whereas the second class of sensors give often higher specificity and avoid cumbersome chemistry. Reversibility of the genetically encoded probes enables real-time imaging of the H₂O₂ dynamics. Currently, two genetically encoded H₂O₂ sensors are available. One is HyPer (3) and the other is Orp1-redox-sensitive green fluorescent protein 2 (roGFP2) (10). HyPer is based on circularly permuted yellow fluorescent protein (YFP) inserted into the regulatory domain of the Escherichia coli OxyR protein. It reacts directly with sub-micromolar H₂O₂ by forming a disulphide bridge, which leads to conformational changes in the protein and, hence, to a YFP spectral change. In the Orp1-roGFP2 fusion, the peroxidase Orp1 oxidizes a cysteine in roGFP2 using H₂O₂ as a substrate. The resulting disulphide changes the roGFP2 β-barrel structure, leading to spectral changes. However, spatial resolution of the H₂O₂ signal imaged with both genetically encoded probes is low. The rapid diffusion of the sensor within the nucleocytoplasmic compartment delocalizes the signal even in the case of local H₂O₂ production.

Peptide growth factors stimulate cellular response via activation of receptor tyrosine kinases (RTKs) localized on the plasma membrane (PM) of cells (19), followed by phosphorylation of various downstream targets. Ligand-bound receptors become rapidly internalized into endosomes and are subsequently either degraded or recycled to the PM (17). Protein tyrosine phosphatase (PTP-1B) attached to the cytosolic face of the endoplasmic reticulum (ER) membrane via a C-terminal “tail anchor” sequence (1) is responsible for
dephosphorylation of many RTKs, including epidermal growth factor receptor (EGFR) and platelet-derived growth factor receptor (PDGFR) (9, 11), especially after internalization (22), and helps switching off the kinase cascade. To allow the propagation of RTK activity, it is necessary to inhibit PTPs including PTP-1B. Cells extensively utilize H₂O₂ to control PTP activity via reversible oxidation of their active site thiolates (6, 13, 18). Although PTP-1B oxidation by H₂O₂ was shown both in vitro and in vivo (6, 12), the mechanism of oxidation of this and other PTPs remains enigmatic. It is not clear how PTPs, which have a rate constant of reaction with H₂O₂ of compartmentalized redox signaling. According to this model, thiolate-containing targets are oxidized by H₂O₂ produced by neighboring NADPH oxidase (Nox) family enzymes residing in the same subcompartment. In this case, the local H₂O₂ concentration could be high enough to temporarily oxidize any nearby peroxiredoxins, allowing further oxidation of PTPs (21). This model, however, requires experimental verification by monitoring real-time dynamics of H₂O₂ production in various cellular compartments upon RTKs activation. An especially controversial issue is which cellular compartment is responsible for Nox activation and H₂O₂ production. The PM, the endosomal compartment, or even the ER membrane are candidates, because all of these compartments have been shown to contain Nox (2, 5).

Therefore, we decided to improve the spatial resolution of H₂O₂ imaging by developing an imaging platform based on immobilized HyPer. We immobilized HyPer probes within certain subcompartments in the cytoplasm and used this strategy to identify localization of the H₂O₂ signal during activation of RTKs. We demonstrated that H₂O₂ produced in RTK signaling does not diffuse freely across the cytoplasm, but instead localizes to microdomains near the sites of its generation.

**Results, Discussion, and Future Directions**

We proposed that H₂O₂ generation should colocalize with either activated receptor or PTP-1B, or both. To evaluate the first possibility, we targeted HyPer to the PM and endosomes by fusing it with the C-termini of EGFR and PDGFR, and the EGFR (1-1210)-HyPer and PDGFR (1-1106)-HyPer fusions were expressed in HeLa and NIH-3T3 cells, respectively. Upon serum starvation, most of the fluorescence was localized at the PM in both cell types. In many NIH-3T3 cells, PDGFR-HyPer localized both at the PM and in endosome-like endomembranes (Supplementary Fig. 1; see www.liebertonline.com/ars).

Stimulation of HeLa cells with EGF resulted in internalization of EGFR-HyPer into endocytic vesicles (Fig. 1A; Supplementary Video 1; see www.liebertonline.com/ars). Importantly, a fraction of the fluorescence retained at its initial membrane localization reflects the noninternalized receptors. We observed that a substantial elevation of H₂O₂ levels in the endosomal fraction started within 7–10 min after stimulation. H₂O₂ concentrations reached a plateau within the next 10–15 min, followed by the second wave of H₂O₂ production (Fig. 1C). However, the fraction of the probe residing at the PM demonstrated no increase in F500/F420 ratio. In fact, the probe associated with the PM became more reduced, suggesting either termination of H₂O₂-producing activity on the PM or translocation of the H₂O₂-generating system from PM to the endosomes. This observation suggested that the endosomes containing internalized receptor served as the compartment responsible for H₂O₂ generation. Addition of the Nox inhibitor diphenyleneiodonium (DPI) at concentrations as low as 5 μM rapidly and completely inhibited H₂O₂ production, indicating Nox as a most likely source of ROS in endosomes (Supplementary Fig. 2; see www.liebertonline.com/ars). The absence of PM-associated HyPer oxidation demonstrated that, first, there was no Nox activation and ROS production on the PM and, second, the diffusion volume of H₂O₂ is limited to a few micrometers. Indeed, in case H₂O₂ would diffuse freely across the cytoplasm, PM-associated HyPer would be oxidized by H₂O₂ produced by endosomes. Importantly, HyPer ratio values often differ between neighboring endosomes (Fig. 1B, D), arguing for the strict limitation of H₂O₂ diffusion. Therefore, we suspect that cellular antioxidant systems degrade H₂O₂ before it is able to reach distant compartments.

H₂O₂ is generated upon activation of many different RTKs. However, it is not known if the spatiotemporal patterns of H₂O₂ production differ between cell types expressing different receptors. To address this question, we stimulated NIH-3T3 fibroblasts expressing PDGFR-HyPer with PDGF. In contrast to EGFR-HyPer–expressing HeLa cells, we did not observe intensive internalization of PDGFR-HyPer into endosomes, but rather intensive filopodia formation and changes of the cell shape in response to the stimulus. The spatiotemporal pattern of H₂O₂ generation also differed from that in HeLa cells. In PDGFR-HyPer–expressing cells, the majority of H₂O₂ production was associated with the PM (Fig. 2A; Supplementary Video 2; see www.liebertonline.com/ars), including filopodia (Fig. 2B, C, E; Supplementary Video 3; see www.liebertonline.com/ars). H₂O₂ started to rise within 1 min after PDGF addition (Fig. 2D, E) and peaked after 10 min, followed by partial reduction of the probe within the next 10 min and the subsequent start of the second H₂O₂ wave (Fig. 2D). Fluorescence of HyPer associated with PDGFR-containing endomembranes remained almost unchanged during the first 20 min after stimulation and then started to rise (Fig. 2D). This showed that PM contained the pool of active Nox in NIH-3T3 cells upon RDGFR stimulation. The absence of diffusion of H₂O₂ from the PM to endomembranes, at least below some [H₂O₂] threshold at the PM, resembled that in HeLa cells, indicating rapid degradation of H₂O₂ by intracellular oxidant defense systems. The elevation of H₂O₂ in 3T3-NIH fibroblasts appeared to be DPI sensitive, supporting that Nox may function as a source of ROS (data not shown).

Being a target of H₂O₂, phosphatase PTP-1B becomes oxidized during RTK activation (12, 13). Membranes hosting activated EGFR and PDGFR have been previously shown to contact with microdomains of PTP-1B–containing ER membrane where receptor dephosphorylation occurs (11). At the same time, membranes containing activated receptors also possessed strong H₂O₂ production, likely because of Nox activity (Figs. 1 and 2), for PTP-1B inhibition. Another possible source of H₂O₂ for PTP-1B oxidation was the ER-residing Nox pool. We proposed that [H₂O₂] dynamics at the ER membrane could provide some information to support either ER or non-ER Nox to be a source of the oxidant. For this
purpose, we produced HyPer-tail anchor (TA). HyPer targeted to the cytoplasmic surface of the ER membrane, by fusing the indicator with the TA of PTP-1B (24 C-terminal amino acid residues) (Supplementary Fig. 1). When expressed in HeLa and NIH-3T3 cells, HyPer-TA localized to the ER (Fig. 3A and Supplementary Fig. 1), demonstrating the same localization pattern as the EYFP-tagged PTP-1B (1) and recently reported HyPer-ERCyto construct (8). Upon stimulation of EGFR in HeLa cells, HyPer-TA detected increases of H$_2$O$_2$ near the ER membrane (Fig. 3A; Supplementary Video 4; see www.liebertonline.com/ars). However, the temporal pattern of the fluorescent response differed from that of EGFR-HyPer.

The first wave of H$_2$O$_2$ started with less than a minute delay after the stimulus and lasted for 10–15 min before a second wave of H$_2$O$_2$ of much higher amplitude followed (Fig. 3C). This second wave of HyPer-TA oxidation reflected either direct ER-produced H$_2$O$_2$ or, alternatively, the endosomal pool of the oxidant reaching the ER membrane. The first H$_2$O$_2$ peak, despite being of low amplitude, started before the endosome-associated H$_2$O$_2$ pool became detectable. This indicated that the ER-associated H$_2$O$_2$ production becomes activated faster than endosomal H$_2$O$_2$ appears.

Addition of PDGF to the NIH-3T3 cells expressing HyPer-TA caused a significant change in the probe ratio, reflecting intensive H$_2$O$_2$ production near the ER of the fibroblasts (Fig. 3B; Supplementary Video 5; see www.liebertonline.com/ars). The shape and amplitude of H$_2$O$_2$ change closely resembled those observed on the PM using PDGFR-HyPer (compare Fig. 3D with Fig. 2D). Together with the observation that the HyPer ratio at the PDGFR-containing endomembranes remained unchanged at the same time (Fig. 2D), HyPer-TA oxidation suggests the existence of a specific ER-residing Nox pool.

Immobilized HyPer variants, unlike the nontargeted probe, allow a significant enhancement of the spatial resolution in H$_2$O$_2$ imaging. Using this approach, we demonstrated for the first time the microdomains of H$_2$O$_2$ production in RTK signaling. Upon activation of EGFR in HeLa cells, a substantial amount of H$_2$O$_2$ was generated on endosomes. Therefore, endosomes carrying internalized EGFR possess both activities required for maintaining downstream target phosphorylation after tyrosine kinase activation: RTK phosphorylates substrates, whereas Nox releases ROS exactly at the same place, inhibiting PTPs and other targets. Such endosomes represent powerful functional units of intracellular EGFR and redox signaling. Rapid rises of H$_2$O$_2$ at the ER might be necessary for the immediate inhibition of PTP-1B on these membranes. However, the mechanism required for the initial activation of ER-resident Nox remains to be identified.

The absence of PM-associated HyPer oxidation in HeLa cells suggests that H$_2$O$_2$ production and many of the H$_2$O$_2$ targets are localized within the cell. In contrast, activation of Nox at the PM in fibroblasts argues for possible extracellular targets of H$_2$O$_2$ produced upon PDGFR activation (15). Indeed, as the diffusion of H$_2$O$_2$ within the cytoplasm is limited to a few microns, H$_2$O$_2$ for paracrine signaling should be produced by PM-residing Nox.

Oxidation of HyPer-TA in both EGFR and PDGFR signaling provides a reliable model to study the PTP-1B inhibition by H$_2$O$_2$. Comparison of timing and amplitude of HyPer-TA oxidation with that of EGFR-HyPer and PDGFR-HyPer fusions revealed a high probability of ER-resident Nox activation in both cell types. For PDGFR signaling in NIH-3T3 fibroblasts, Nox in the ER could be the major source of PTP-1B inhibition because of the apparent absence of receptor internalization and the limitation of intracellular H$_2$O$_2$ diffusion.

Taken together, imaging by immobilized HyPer variants in complementation with recent research on tissue-scale H$_2$O$_2$ patterning (15) shows how biological systems control highly reactive and diffusible molecules and use them for signaling.

Materials and Methods

**Materials**

H$_2$O$_2$, D$_2$O, EGF, and PDGF-BB were purchased from Sigma. Dulbecco-phosphate-buffered saline, Dulbecco’s modified Eagle’s medium, Opti–minimal essential medium (MEM), MEM, fetal calf serum, and FuGene6 transfection reagent were from Invitrogen. Glass-bottomed dishes were from MatTec. NIH-3T3 cells were from ATCC. HeLa-Kyoto cell line was provided by EMBL. Encyclo PCR kit and HyPer expression vectors were from Evrogen. Restriction endonucleases were from SibEnzyme.

**Methods**

Preparation of fusion constructs. To make EGFR-HyPer, the YFP-coding region of pEGRF-YFP vector was replaced with the coding region of Hyper. Hyper was amplified from HyPer-cyto (Evrogen) with the primers 5’-ATCTCGATCG GTGGAGGTTGGAGCGGAGATGGCGAACCGACCGACCGC-3’ and 5’-CCGACGCGGCCGCTTAAACCGCCTGTTTTAAAACTT-3’; the PCR product was digested with PvuI and NotI and cloned into EGFR-YFP in place of YFP.

PDGFR-HyPer was produced by replacing the GFP-coding region of PDGFR-GFP by HyPer-coding DNA amplified from HyPer-cyto vector using the primers 5’-ATCTGGTACCGAGTCTCGATCG GTGGAGGTTGGAGCGGAGATGGCGAACCGACCGACCGC-3’ and 5’-CCGACGCGGCCGCTTAAACCGCCTGTTTTAAAACTT-3’; the PCR product was digested with PvuI and NotI and cloned into EGFR-YFP vector.

PDGFR-HyPer was produced by replacing the coding region of PDGFR-GFP by HyPer-coding DNA amplified from HyPer-cyto vector using the primers 5’-ATCTGGTACCGAGTCTCGATCG GTGGAGGTTGGAGCGGAGATGGCGAACCGACCGACCGC-3’ and 5’-CCGACGCGGCCGCTTAAACCGCCTGTTTTAAAACTT-3’; the PCR product was digested with PvuI and NotI and cloned into EGFR-YFP vector.

All constructs were confirmed by sequence analysis.

Cell culture and transfection. HeLa-Kyoto and NIH-3T3 cells were cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal calf serum at 37°C in an atmosphere containing 95% air and 5% CO$_2$. Cells were split every four hours later, cells were transfected with a mixture of vector DNA and FuGene6 transfection reagent according to the manufacturer’s recommendations.

Stimulation of the cells and imaging. Twenty-four hours after transfection, the culture medium was changed to MEM without phenol red and serum. After 2 h (HeLa-Kyoto) or 6 h (NIH-3T3) of incubation, the glass-bottomed dishes were transferred onto the microscopic stage of either a confocal (laser scanning confocal inverted microscope Carl Zeiss LSM 510-META or PerkinElmer Ultraview VoX confocal inverted microscope) or a widefield (Leica 6000) microscope equipped with an HCX PL APO lbd.BL 63×1.4NA oil objective and an environmental chamber (EMBL). HyPer fluorescence was
FIG. 1. Microdomains of elevated H$_2$O$_2$ generation upon activation of EGFR. (A) Confocal images of EGFR-HyPer–expressing HeLa-Kyoto cells at indicated time points (in minutes) after stimulation of the cells with 50 ng/ml EGF. Upper row of images represent subcellular distribution of HyPer ratio. Lower row of images show subcellular distribution of EGFR-HyPer. Cell boundaries and nucleus of one of the cells are highlighted. Scale bar: 15 μm. (B) Confocal image of a selected region inside the cell containing internalized receptor. Image represents HyPer ratio values. Note that neighboring endosomes demonstrate different HyPer ratio. Signal intensity profiles along the lines are shown on (D). Scale bar: 5 μm. (C) Time course of H$_2$O$_2$ production associated with endosomes and PM in the cells shown on (A). (D) HyPer ratio values along the upper (red) and lower (black) lines shown on (B). The figure is representative of 37 cells from three experiments. H$_2$O$_2$, hydrogen peroxide; PM, plasma membrane; EGFR, epidermal growth factor receptor. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article at www.liebertonline.com/ars).
FIG. 3. Dynamics of H$_2$O$_2$ near the cytoplasmic surface of the endoplasmic reticulum membrane upon activation of the cells with growth factors. (A) Widefield fluorescence images of HyPer-TA–expressing HeLa-Kyoto cells at indicated time points (in minutes) after stimulation of the cells with 50 ng/ml EGF. Upper row of images represent subcellular distribution of HyPer ratio. Middle and lower rows of images show subcellular distribution of HyPer-TA and changes in fluorescence intensity in each imaging channel. Scale bar: 15 µm. (B) Confocal images of HyPer-TA–expressing cells at indicated time points (in minutes) after stimulation of the cells with 10 ng/ml PDGF. Upper row of images represent subcellular distribution of HyPer ratio. Middle and lower rows of images show subcellular distribution of HyPer-TA and changes in fluorescence intensity in each imaging channel. Scale bar: 10 µm. (C) Time course of H$_2$O$_2$ production by the cells shown on (A). (D) Time course of H$_2$O$_2$ production by the cell shown on (B). The figure is representative of 11 HeLa-Kyoto cells from three experiments and 43 NIH-3T3 cells from three experiments. TA, tail anchor. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article at www.liebertonline.com/ars).

FIG. 2. Microdomains of elevated H$_2$O$_2$ generation upon activation of PDGFR. (A) Confocal images of PDGFR-HyPer–expressing NIH-3T3 cell at indicated time points (in minutes) after stimulation of the cells with 10 ng/ml PDGF. Upper row of images represent subcellular distribution of HyPer ratio. Lower rows of images show subcellular distribution of EGFR-HyPer and fluorescence intensities in two imaging channels excited with 405 and 488 nm lasers (see Materials and Methods section for details). Scale bar: 15 µm. (B) H$_2$O$_2$ generation in filopodia of PDGFR-HyPer–expressing NIH-3T3 cell after stimulation with PDGF. Scale bar: 15 µm. (C) Illustration of the events visible on (B). Upon addition of PDGF, cell starts changing its shape. This results in simultaneous movement of the nucleus away from the confocal optical section and appearance of the “dorsal” PM in the optical slice. At this moment, intensive filopodia formation on the PM is clearly visible as well as production of H$_2$O$_2$ by the receptor localized on PM. (D) Time course of H$_2$O$_2$ production by receptor-containing endomembranes and PM in the cell shown on (A). (E) Time course of H$_2$O$_2$ production by filopodia of stimulated NIH-3T3 cell shown on (B). The figure, except (B), (C), and (E), is representative of 27 cells from three experiments. PDGFR, platelet-derived growth factor receptor. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of this article at www.liebertonline.com/ars).
excited sequentially with 405 and 488 nm lasers of confocal fluorescent microscope or 427/10 and 504/12 band-pass excitation filters of the widefield fluorescent microscope. Emission of the probe was collected every 30 s using a 527 W/55 band-pass emission filter of the confocal system or a 525/50 band-pass emission filter of the widefield fluorescent microscope. After 3–5 images were acquired, 50 ng/ml EGF or 10 ng/ml PDGF was added.

**Time series processing.** Time series were analyzed using ImageJ free software downloaded from EMBL ALMF website (www.embl.de/almf/). For confocal images, 420- and 500-nm stacks (correspond to two HyPer excitation peaks) were Gaussian filtered and the background was subtracted. Images were converted to 32 bits and a 420-nm stack was thresholded to remove pixel values from background (Not-a-Number function). A 500-nm stack was divided into a 420-nm stack frame-by-frame. The resulting stack was depicted in pseudocolours using a “Ratio” lookup table. Time course of HyPer fluorescence was calculated for regions of interest inside the imaged cell.

To calculate average HyPer ratio values in the endosomes, we used the fact that the ratio in endosomes is higher than the ratio associated with the PM. Therefore, we thresholded images eliminating ratio values below 0.7. Upon thresholding, mostly endosomes remained visible in the images. After that, the average ratio values for each frame in the stack were calculated.

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### Abbreviations Used

- DPI = diphenyleniodonium
- EGFR = epidermal growth factor receptor
- ER = endoplasmic reticulum
- H₂O₂ = hydrogen peroxide
- MEM = minimal essential medium
- Nox = NADPH oxidase
- PDGFR = platelet-derived growth factor receptor
- PM = plasma membrane
- PTP = protein tyrosine phosphatase
- roGFP2 = redox-sensitive green fluorescent protein 2
- ROS = reactive oxygen species
- RTK = receptor tyrosine kinase
- TA = tail anchor
- YFP = yellow fluorescent protein