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Evaluation of three simple direct or indirect carbonyl detection methods for characterization of oxidative modifications of proteins

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Abstract
Among disruptions induced by oxidative stress, modifications of proteins, particularly irreversible carbonylation, are associated with the development of several diseases, including cardiovascular diseases, neurodegenerative diseases, and cancer. Carbonylation of proteins can occur directly or indirectly through the addition of lipid oxidation products. In this study, three classical and easy-to-perform techniques to detect direct or indirect carbonylation of proteins were compared. A model protein apomyoglobin and a complex mixture of rat liver cytosolic proteins were exposed to cumene hydroperoxide oxidation or adduction to the lipid peroxidation product 4-hydroxynonenal in order to test direct or indirect carbonylation, respectively. The technique using a specific anti-4-hydroxynonenal-histidine adduct antibody was effective to detect in vitro modification of model apomyoglobin and cytosolic proteins by 4-hydroxynonenal but not by direct carbonylation which was achieved by techniques using biotin-coupled hydrazide or dinitrophenylhydrazine derivatization of carbonyls. Sequential use of these methods enabled the detection of both direct and indirect carbonyl modification in proteins, although constitutively biotinylated proteins were detected by biotin-hydrazide. Although rather classical and efficient, methods for carbonyl detection on proteins in oxidative stress studies may be biased by some artifactual detections and complicated by proteins multimerizations. The use of more and more specific available antibodies is recommended to complete detection of lipid peroxidation product adducts on proteins.

Keywords: Protein carbonyls, 4-hydroxynonenal, lipid peroxidation, biotin-hydrazide, dinitrophenylhydrazine

Introduction
Oxidative stress occurs when the levels of intracellular reactive oxygen species (ROS) exceed the detoxification capacity of the cell. It induces disruption of redox regulation trough disturbances in cell signaling and protein-protein interactions and damages macromolecule. Oxidative protein modifications play central role in major human diseases, such as atherosclerosis, diabetes, cardiovascular diseases, cancer, neurodegenerative disorders, and chronic liver and lung diseases (Nystrom 2005; Valko et al. 2007; Waris and Ahsan 2006; Negre-Salvayre et al. 2010). Oxidative stress induces significant biological damage in cells through the modification of DNA, proteins, lipids, and carbohydrates. In proteins, the modification can be induced directly by ROS or indirectly by reaction of intermediates due to oxidation of other cellular components, such as lipids or carbohydrates, yielding advanced lipoxidation or glycation end products.

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Modification resulting in the addition of reactive functional carbonyl groups is called "protein carbonylation."

Radicals react directly with amino acids, such as arginine, threonine, proline, and lysine (Dalle-Donne et al. 2003), causing the formation of semi-aldehyde amino acids. Direct carbonylation of proteins can also arise through oxidative cleavage of proteins (α-amidation pathway or oxidation of glutamine side chains) (Aldini et al. 2007). Indirect carbonylation of proteins results from Michael addition of lipid peroxidation products involving α, β-unsaturated aldehydes such as 4-hydroxynonenal (HNE), 4-oxonononal (ONE), 2-propenal (acrolein) and the nucleophilic side chains of Cys, His, or Lys residues (Stadtmann and Berlett 1997; Aldini et al. 2006) or by reaction of reactive carbonyl derivatives (ketamines, ketoaldehydes, and deoxyosones) with lysine residues (Dalle-Donne et al. 2003).

Carbonylation is an irreversible, nonenzymatic modification of proteins. Protein carbonylation results in structural and conformational changes that lead to the malfunction of many enzymes, damage to membrane transporters, and impaired interactions with receptors. This modification is also related to the formation of protein cross-linkages; the resulting aggregates may accumulate in cells leading to degenerative diseases (Dalle-Donne 2006b). Carbonylation of specific amino acid side chains is one of the most abundant consequences of oxidative stress (Requena et al. 2003).

One of the most abundant and toxic α, β-unsaturated aldehyde product of omega-6 polyunsaturated fatty acids is HNE. It is formed at basal concentrations under physiological conditions, and its cellular concentration ranges from 0.2 to 3 μM. Because HNE is an amphiphilic compound with strong lipophilic properties, it accumulates in biomembranes and reacts with phospholipids and membrane proteins (Poli et al. 2008), such as receptors and transporters. In addition, HNE has a wide range of biological activities, including inhibition of protein and DNA synthesis, disruption of Ca²⁺ homeostasis, membrane damage, cell death, and activation of stress signaling pathways. Most of these effects are thought to be mediated by covalent linking of HNE to proteins (Petersen and Doorn 2004; Poli et al. 2008). Previous works have shown that the C-3 carbon of HNE, which participates in the double bond, is a soft electrophilic center that interacts with the soft nucleophilic sulfur atom of cysteine. This carbon can also react with the imidazole nitrogen of histidine or, although to a lesser extent, the amine nitrogen of lysine to form Michael adducts (Esterbauer et al. 1991; Doorn and Petersen 2003; Grimsrud et al. 2008; LoPachin et al. 2009). Structures of HNE-protein adducts have been elucidated from chemical studies and also from in vitro or in vivo studies followed by complete acid or enzymatic hydrolysis and mass spectroscopic evaluation. Although HNE is much more reactive toward cysteine than histidine, it has been postulated that the HNE adduct of histidine is more stable than cysteine or lysine conjugates toward retro-Michael addition (Nadkarni and Sayre 1995; Doorn and Petersen 2002; Petersen and Doorn 2004; Sayre et al. 2006; Grimsrud et al. 2008; Lesgards et al. 2009).

Such Michael adducts have also been observed with thiol groups of glutathione. In model systems, mass spectrometric analysis has demonstrated that > 99% of proteins modified in vitro by HNE present a free carbonyl group (Bruenner et al. 1995). One of the most studied lipid peroxidation products in biological fluids or tissues is HNE, and it is now considered as one of the best ways to evaluate ongoing oxidative stress in vivo through measurement of HNE-protein adducts (Poli et al. 2008). Advanced methods are needed to elucidate the biological role of carbonyl protein modifications, the interplay of oxidative stress, and degradation of the oxidatively modified proteins as well as the molecular mechanisms involved in the development and progression of diseases. The methods must be in agreement with the projected objective, ensure practicality and reproducibility, and maintain the quality and quantity of the protein for further identification. Precise identification of modified sites constitutes an ultimate objective. Global approaches, such as proteomics, remain advantageous because they provide comparative information on the constant changing of the cell proteome through its biochemical interactions with the genome and environment. Thereby, proteomics is useful for identification of carbonylated proteins in a complex mixture.

Among several assays, which have been developed for detection of protein carbonyls, the most widely used involves derivatization of the carbonyl group with 2,4-dinitrophenylhydrazine (DNPH) followed by detection of the stable hydrazone product formed 2,4-dinitrophenyl (DNP) (Levine et al. 1990). The DNP adduct can be detected by various techniques, including spectrophotometric assay (Levine et al. 1990, 1994), enzyme-linked immunosorbent assay (ELISA), and one-dimensional- or
two-dimensional-electrophoresis coupled to western blot.

To select the best methods for protein carbonyl isolation from a complex mixture of proteins, the authors only used methods that allow for evaluation of the protein changes in one step without a priori digestion. These methods detect HNE-protein adducts using an anti-HNE-His antibody (Ab-HNE) or protein carbonyl derivatization with biotin-hydrazide (BH), followed by streptavidin detection or 2, 4-DNPH and anti-DNP antibody detection. These methods, which can be easily used as they do not involve sophisticated techniques, allow the detection of whole proteins by electrophoresis methods. Here, easy, sensitive, and improved methods are presented and compared to detect direct and indirect protein carbonylation in the context of oxidative stress studies.

Material and methods

Animals

Male Wistar rats (230–250 g) were obtained from Harlan, Granat, France. The rats had free access to water and to a semipurified diet (AIN 76, from UAR, Epinay, France). All experiments were performed in accordance with the institutional guidelines governing animal experimentation.

Chemicals and reagents

EZ-Link® Biotin Hydrazide, MES Buffer [(2-N-morpholino) ethanesulfonic acid] and desalting spin columns were from Pierce (Rockford, IL, USA). Myoglobin from equine heart, cumene hydroperoxide (CH), and 2,4-DNPH were from Sigma Aldrich (St Quentin Fallavier, France). The Lowry assay kit for protein concentration determination was from Bio-Rad (Richmond, CA, USA). COMPLETE protease inhibitor cocktail tablets were provided by Roche Molecular Biochemicals (Mannheim, Germany). Anti-dinitrophenyl (Anti-DNP) antibody was from Sigma Aldrich. The mouse specific anti-HNE-His adducts antibody was a generous gift from Dr Waeg. It has been produced and tested as described (Waeg et al. 1996). The IRDye 800CW streptavidin and IRDye 800CW Goat anti-mouse antibodies were from Li-Cor Bioscience (Lincoln, NE, USA). CF680 Goat anti-rabbit antibody was from Biotium (Hayward, CA, USA).

HNE preparation

The stable HNE precursor 4-[(tetrahydro-2H-pyran-2-yl)oxy]-2(E)-nonenal (4-OTHP-HNE) was synthesized according to a published method (Jouanin et al. 2008). The protecting group tetrahydropyranol (THP) was removed by dilution in methanol (150 µL of a 10 mM solution) and addition of PTSA (p-toluenesulfonic acid: 300 µL of a 10 mg/mL solution in methanol). After stirring at room temperature for 3 h, the methanol was evaporated and water was added. The HNE was extracted by methylene chloride, washed with brine, and dried with magnesium sulfate. The solution was evaporated and then 110 µL of water was added. The concentration was determined by UV absorption (223 nm, ε13750 L.mol⁻¹.cm⁻¹) of a diluted solution (1/1000 dilution).

In vitro oxidation and addition methods

Apomysoglobin (apo) was prepared from horse heart myoglobin, which was dissolved in water (~ 10 mg/mL) and precipitated in cold acetone containing 0.3%, 2 N HCl for 30 min at −20°C. After centrifugation for 1 min at 1000 × g, the precipitate was dried under a nitrogen stream and dissolved in ultra pure water. The bovine serum albumin (BSA) was directly dissolved in 50 mM sodium phosphate, pH 7.4. Liver tissue samples (~100 mg) were homogenized in a 10 mM HEPES buffer containing 137 mM NaCl, 4.6 mM KCl, 1.1 mM KH₂PO₄, and 0.1 mM EDTA, pH 7.4 and supplemented with 0.1 mM PMSF and protease inhibitor cocktail prepared according to the manufacturer’s directions. The cytosolic fraction was obtained from liver homogenates by successive centrifugations of 20 min at 14,000 × g and 1 h at 100,000 × g. The soluble fraction was taken as the cytosol. Before in vitro oxidation, the different extracts were passed through desalting columns to replace the cellular fractionation buffer. The oxidation procedure was performed on 250 µg of protein. Addition of HNE to apo, BSA and total proteins from liver cytosolic fraction was performed as described hereafter. The APO and BSA (2 mg/mL) were incubated for 2 h at 37°C or at room temperature in 50 mM sodium phosphate buffer, pH 7.4 containing 1.3 mM HNE. Cytosolic proteins (5 mg/mL) were exposed to 2 mM HNE for 2 h at room temperature. Treatment with CH was carried out according to the previously described method of Arguelles et al. with minor modifications (Arguelles et al. 2006). Briefly, APO and protein extracts (100 µg) were incubated for 20 min at 37°C with 1 mM CH in PBS.

Carbonyl adducts detection

Detection with BH reagent was adapted from the method described by Sayre et al. (2006) and Roe et al. (2007). Briefly, controlled and modified APO, BSA, and the cytosolic proteins were filtered on Sephadex G25 (Ø1 × 10 cm column) or desalting spin columns equilibrated in 100 mM sodium acetate buffer, NaCl 150 mM, pH 5.5. Next, 50 mM BH reagent in dimethylsulfoxide was added to a 1 mg/mL protein solution to yield a final concentration of 5 mM. Incubation was performed for 2 h at room temperature with vigorous mixing (400 rpm). The controls were incubated with DMSO. Reaction with DNPH was performed according to the method described by Levine et al. (1994). Briefly, 40-µg samples of control or modified proteins were incubated in 2 N HCl additioned with 20 µL of 12% SDS and 200 µL of 10 mM DNPH for 20 min at room temperature. After the reaction, the pH was neutralized with 2 M Tris in 30% glycerol. The samples were
then prepared for western blotting and two-dimensional electrophoresis.

Western blotting

Samples were boiled for 5 min in classical Laemmli buffer prior to western blotting. Proteins were separated by SDS-PAGE (12.5 %) at 200 V for about 1 h using the Mini-Protean III™ device (Bio-Rad) and then transferred to a Trans-Blot® Transfer Medium nitrocellulose membrane (Bio-Rad) using the Transblot-Blot® SD Semi-Dry Transfer Cell (Bio-Rad) at constant 250 mA (65 V) for 2 h and 15 min in an ice bath. The membranes were blocked for 1 h at room temperature in PBS containing 0.1% Tween 20 and 5% nonfat milk (blocking buffer) to avoid nonspecific labeling. The HNE-His protein adducts were detected on the blot after incubation with primary mouse anti-HNE-His antibody (1/100), followed by secondary goat anti-mouse antibody (1/5000) coupled to IR-Dye® goat anti-rabbit antibody (1/10,000) in blocking buffer. The blot membranes were successively incubated for 1 h at room temperature with a primary rabbit antibody (1:1000) specific for DNP-protein adducts (Sigma-Aldrich) followed by a secondary goat anti-rabbit antibody coupled to IR-Dye® 800CW (Li-Cor Biosciences) at 1:10,000 concentration in blocking buffer. Antibody dilutions were made in blocking buffer. The blots were scanned using an Odyssey Infrared Imaging System (Li-Cor Biosciences).

Two-dimensional electrophoresis

The protein concentration was estimated using the Lowry method (RC DC Protein Assay, Bio-Rad) or Amido-black staining (Schaffner and Weissmann 1973). For the first dimension, protein samples (100 µg) were loaded onto a 110-mm pH 3–10 ReadyStrip™ IEF strip (Bio-Rad, Marnes-La-Coquette, France) and covered with oil. Isoelectric focusing (IEF) was achieved on an Ettan IPGphor Isoelectric Focusing System (GE Biosciences-Amersham) by active rehydration for 18 h at 50 V in 125 µL of buffer containing 8 M urea, 2% ampholytes, 10 mM dithiothreitol (DTT), 2% Chaps, and 2% bromophenol blue, followed by stepwise application of 50 V for 12 h, 300 V for 1 h, 500 V for 1 h, 1000 V for 1 h and finally a linear gradient to 8000 V for 5 h, for a total of at least 35,000 Vh on the same instrument. The strips were stored at −80°C until the second-dimension electrophoresis. For the second dimension, the strips were thawed and equilibrated with gentle shaking in an equilibration solution (50 mM Tris-HCl, 6 M urea, 30% glycerol, 2% SDS, pH 8.8) with 1% DTT for 15 min and then in an equilibration solution with 2.5% iodoacetamide for 15 min. The strips were loaded onto 12.5% SDS-PAGE gels; in addition, 3 µL of PageRuler™ prestained molecular weight marker (Fermentas, Saint-Rémy-Lès-Chevreuse, France) was loaded against the IEF-strips before sealing with 0.5% agarose. Gels were run for 2 min at 250 V and then for 2 h at a constant voltage of 100 V. Proteins were then transferred to nitrocellulose using the same protocol used for western blotting.

Results

Experiments using model proteins, that is, APO and BSA, and cellular cytosolic extracts were designed to improve the efficacy of two different modifying agents, HNE and CH, in the formation of protein carbonyls. Three detection methods among the most routinely used techniques were tested and compared, namely histidine-HNE conjugate monoclonal antibody (Ab-HNE) immunodetection, BH derivatization followed by streptavidin detection and 2,4-DNPH conjugates.

Comparison of the three methods for the detection of APO modifications by HNE and CH

To evaluate the efficacy of the different detection methods for a pure protein, in vitro addition by HNE and oxidation by CH on a model protein, APO, by detecting HNE adducts and carbonyl formation using western blotting was first investigated. As shown in Figure 1A, the formation of HNE adducts on APO (band at 17 kDa) after exposure to 4-HNE using the Ab-HNE, BH, or DNPH methods was clearly detected. Immunoreactive extra bands migrating close to 35 and 68 kDa may represent the formation of dimers and tetramers, respectively, during the HNE-protein reaction in these experimental conditions. This generation of multimeric species is likely due to the formation of crosslinks between proteins because of the double reactivity of HNE, which can also undergo Michael additions with Cys, His, or Lys residues of one protein molecule and Schiff bases with the Lys residue of another (Roede et al. 2008). The absence of trimeric forms suggests that specific spatial conformations may be required for aggregation. As expected, oxidation of APO with CH did not induce the formation of HNE-His adducts, but CH was able to oxidize APO by a carbonylation process. Indeed, bands were detected using the BH and DNPH methods but not with Ab-HNE (Figure 1B). The observed bands are similar to those detected on HNE-treated APO, suggesting similar multimerization of the oxidized protein and confirming the involvement of carbonyl groups in this process.

Detection of cytosolic proteins modified by in vitro treatment with HNE and CH

In vitro HNE modification and protein carbonylation of rat liver total cytosolic proteins were investigated using the same Ab-HNE, BH, and DNPH methods, as described above for APO. An increase of HNE adducts as compared to basal level in proteins ranging in
size from 25 to 250 kDa was clearly detected with the Ab-HNE method combined with western blotting analyses (Figure 2A). In addition, the level of carbonylated proteins was increased by HNE treatment as observed when using the DNPH method. A pattern similar to that obtained with the BH method, but with slightly higher global intensity and specific band difference, such as the ones at 30–33 kDa or 25 kDa, observed after HNE treatment was detected (Figure 2A). Because false positive results are possible with the BH method, detection using IRDye-coupled streptavidin on samples not treated with BH to evaluate the possible constitutive expression of biotinylated proteins in the cytosolic fractions was performed. Western blots revealed positive staining for the BH-untreated HNE-treated cytosolic proteins at two major bands of 70 and 120 kDa. Such staining was not observed for unbiotinylated model proteins (i.e., BSA and APO) (Figure 3). Western blot analysis after CH oxidation revealed reactive proteins ranging from 27 to 250 kDa (Figure 2B). As expected Ab-HNE technique confirmed the inability of CH to induce the formation of HNE-His adducts on lipid-deprived cytosolic extracts (Figure 1B). In parallel, the BH and DNPH methods revealed that the carbonyl groups were quantitatively enhanced after CH treatment in comparison with the controls.

**Carbonylated protein detection in two-dimensional gel electrophoresis by Ab-HNE, BH, and DNPH methods**

Two-dimensional gel electrophoresis (2DGE) method, which has been frequently used to visualize and identify carbonylated proteins in complex mixtures, was
performed. The ability of the Ab-HNE, BH, and DNPH methods to identify directly and indirectly carbonylated proteins in the same cytosolic fraction (Figures 4 and 5) was compared. Separation of carbonylated proteins using 2DGE revealed an increased occurrence of each reactive protein and showed “string of pearls” shapes, which usually occurs for differently charged isoforms of similar protein backbones. This observation supports an enhanced modification of protein charges resulting from heterogeneous multisite oxidation. Using 2DGE, the results obtained by western blotting analyses with a main group of modified protein ranging from 25 kDa to 55 kDa were confirmed. Additional information on protein charges brought by the IEF step reveals that proteins migrating around 100 kDa in western blot analysis are charged negatively (Figures 4 and 5). As expected, CH-oxidized proteins were found unchanged, when tested with the Ab-HNE method (not shown in Figure 5).

**Discussion**

Oxidative stress is a disturbance in the balance of the redox state of the cell, inducing damage in membranes, DNA and proteins through disturbance of intracellular signaling, enzymes efficiency, and protein–protein interactions. Protein oxidation induced by oxidative stress is likely involved in the evolution of many diseases (Kohen and Nyska 2002). However, the identification of
oxidized proteins is incomplete. Therefore, a targeted proteomic analysis of oxidized proteins is necessary to elucidate their relationships with pathological status and the protein-level mechanisms that induce disease. The potential utility of oxidized proteins as biomarkers of disease has initiated much interest in the development of methodologies to characterize protein modifications on a proteome-wide scale (Dalle-Donne et al. 2003, 2006a). This study reports the application of three different methods for labeling carboxylated proteins directly or indirectly for identification by proteomic analysis. Carboxylated proteins are classic indicators of oxidative stress damage. The incubation of sample proteins with DNPH can be used to measure protein carboxylation by colorimetry, immunohistochemistry, mass spectrometry, and western blotting. The DNPH method can detect not only carbonyl groups but also specific modifications, such as HNE, acrolein, ONE, and malondialdehyde (Sayre et al. 2006). Using standard streptavadin-coupled detection techniques, BH has been shown to be able to detect protein carbonyl groups, including the aldehyde group of HNE when it is added to protein by Michael addition. On the other hand, anti-His-HNE antibodies allow specific visualization of HNE-histidine adducts either in tissues by western blot and immunohistochemistry or with ELISA techniques (Negre-Salvayre et al. 2008; Spickett et al. 2010).

First a conventional immunostaining method using an antibody that recognizes HNE-His adducts, which have been proposed as a marker for the progression of several diseases (Loguerio and Federico 2003; Reed et al. 2008) was evaluated. The results validate the efficiency of the HNE-His antibody method in a single protein assay or in a complex protein mixture by western blotting analysis or by two-dimensional electrophoresis with antibody detection. This classical method provides an easy way to identify the modification of proteins by Michael addition of HNE to His. Specific antibodies for HNE adducts have been used most for detecting HNE-protein adducts in cells and tissues. Such immunohistochemical detection of the HNE-His adducts in various human and animal tissues revealed both physiological presence of the HNE-modified proteins as well as their pathological distribution in particular in malignant (cancer), inflammatory, and degenerative processes (Spickett et al. 2010). Interestingly, comparison of the quantitative (the HNE-His ELISA) and qualitative immunocytochemical analyses in vitro with the qualitative, immunohistochemical analyses in vivo revealed the presence of the HNE-modified proteins in vivo resembling those obtained in vitro for the cell exposed to the 1–10 µM HNE concentration range (Borovic et al. 2006; Zarkovic et al. 2009). This concentration range is much above the values of HNE previously considered to be physiological (0.1–1 µM) indicating physiological importance of lipid peroxidation, while specific protein modification associated with pathology of lipid peroxidation generating higher levels of HNE should be carried out as was done in this study. Namely, when APO or the cytosolic protein extract was treated by CH, there was no detection by the HNE-His Ab method, likely because there was no fatty acid in the purified protein or the cytosolic extract to yield HNE upon oxidation. Mass spectrometric analyses of proteins modified under conditions of oxidative stress revealed that His residues are the major sites for HNE adduction in heme proteins. Liquid chromatography–electrospray ionization–mass spectrometry (LC–ESI–MS)/MS analysis confirmed that HNE adduction occurs exclusively on His residues in bovine APO (Alderton et al. 2003). Although HNE-His is not the most common HNE-protein adduct, HNE-His should reflect the increased presence of HNE during acute oxidative stress or during the late stages of chronic diseases when addition to the cysteine thiolate pool is saturated (LoPachin et al. 2009). Some methods have been developed to evaluate HNE protein adducts on different amino acid residues (Veronneau et al. 2002; Lesgards et al. 2009). However, these methods involve sophisticated methodology and the necessary instrumentation is not widely available.

Two-dimensional electrophoresis has been used frequently for the identification of modified proteins in complex mixtures. To demonstrate the feasibility of using the HNE-His antibody, two-dimensional electrophoresis on a cytosolic protein extract treated in vitro by HNE was used. The HNE-modified protein spots could be clearly visualized after immunoblotting. The number of spots that was obtained indicated that a large number of proteins can be modified by HNE in vitro, which confirms previous data by other authors (Roede et al. 2008). The BH method includes a detection step that uses labeled streptavadin and is based on the reaction of carboxyl groups with hydrazide to form a hydrazone. Although this is a new method, the approach based on hydrazide chemistry has been used by some authors for the enrichment of carbonyl-containing peptides obtained after modification by the lipid peroxidation product HNE (Vila et al. 2008; Codreanu et al. 2009). Here BH was employed to label the HNE adducts and other carboxylated proteins. The BH method has been reported by others to be advantageous over the HNE-His antibody method for detection of not only HNE-His adducts but also other Michael adducts of HNE and direct carboxylated proteins (Vila et al. 2008). Several authors have detected HNE-proteins with this BH method (Roe et al. 2007; Codreanu et al. 2009). However, positive staining in liver cytosolic fractions without reaction with the biotin hydrazine reagent was observed. These bands likely reveal the presence of naturally occurring biotinylated proteins. The apparent molecular weights of these proteins correspond to those of pyruvate carboxylase and propionyl-CoA carboxylase, which were previously characterized in western blots of rat liver samples as naturally biotinylated proteins (Salto et al. 1999). Thus, these artifacts are inherent in streptavadin-biotin detection systems and should be taken into
account in the evaluation of protein carbonylation with the BH method.

The DNPH method was able to identify proteins modified in vitro by HNE and CH. The DNPH clearly detected the HNE-His adduct formed in APO and also in the cytosolic proteins treated with HNE, demonstrating high sensitivity for not only HNE adducts but also other oxidatively modified proteins, which could correspond to carbonylated proteins. The DNPH is one of the older techniques used for detecting protein carbonylation. The use of DNPH has been reported for proteins modified in vitro and in vivo, enabling identification of carbonyl proteins in various traumatic injuries and disease states that involve cellular oxidative stress, including neuronal degeneration and Alzheimer’s disease (Vaishnav et al. 2007; Butterfield et al. 2009). It is herein shown that this method is able to target both direct and indirect protein carbonylation.

The three tested methods allow a first step of evaluation without a priori digestion and give spot profiles by two-dimensional electrophoresis. Image software can be used to match and analyze the protein spots visualized among differential two-dimensional gels, followed by statistic analyses to determine which of the modified proteins may have biological importance. In this way, only the proteins considered significantly different could be subjected to in-gel trypsin digestion and subsequent mass spectrometric identification analyses. In the assays used, the clearest two-dimensional image was observed when using the HNE-His Ab method rather than the BH and DNPH methods. However, staining by HNE-His Ab is recommended to specifically detect protein modifications by oxidative damage.

Conclusion

There was a need to compare methods for detection of protein carbonylation in the context of oxidative stress studies, especially when occurring through distinct direct and indirect processes. As expected, the anti-HNE antibody specifically revealed HNE adducts and not CH-oxidized proteins. A clear separation of HNE-modified proteins using 2DGE and HNE-His Ab was observed. In contrast, the DNPH method detected both CH-oxidized proteins and HNE adducts and could probably detect adducts coming from other bifunctional aldehydes originating from lipid peroxidation, such as 4-hydroxyhexenal and 4-oxononalen. The authors found the BH method to be less specific and to generate artifactual staining of unoxidized proteins with streptavidin coupled dyes. The HNE-His Ab and DNPH methods sound complementary: DNPH method has the advantage to give a global vision of oxidative stress consequences on cellular proteins, while HNE-His Ab method is ideal to get precise information concerning HNE adducts formation, particularly on low-abundant proteins. Although rather classical, methods for carbonyl detection on proteins may be biased by artifactual detections and complicated by proteins multimerizations. The use of more and more specific available antibodies in oxidative stress studies involving detection of lipid peroxidation product adducts on proteins are recommended.

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