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Inflammation: More Than One Explanation

I read with interest the *EHP* supplement on oxygen radicals and lung injury (vol. 102, supplement 10). I would like to take this opportunity to comment about this supplement and raise a key issue concerning the major concepts regarding the mechanisms of cellular injury in inflammatory diseases.

As an active investigator in this field of research, I cannot fully understand why there was no mention in the supplement about the basic understanding that cellular damage in inflammation is multifactorial. The nonexpert reader of this supplement might receive an erroneous impression that oxygen radicals, per se, are the exclusive toxic agonists that induce cellular injury. Many in this field share the view that cellular damage in inflammatory diseases might be caused by a "coordinated cross-talk" among oxidants, membrane-damaging agents, proteinases, arachidonic acid metabolites, phospholipases, cationic proteins, and cytokines. All these agents are likely to be present in sites of infection and inflammation. But sadly, none of the publications elaborating on this multifactorial view are quoted in modern textbooks or in symposia on inflammation and inflammatory diseases. Instead, the literature is filled with publications that insist on a single agonist, be it an oxidant, a

protease, a cytokine, etc., in experimental models. No attempt to integrate the various agonists into the full picture is made.

Several of our publications (1–7) deal with synergistic interactions among multiple proinflammatory agonists in cellular injury during inflammation. I believe that this issue is important, timely, and might contribute to an understanding of how drugs, chemicals, and xenobiotics function *in vivo*.

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Response

We appreciate the interest shown by Dr. Ginsburg in our recent conference proceedings (*EHP* 102, supplement 10). As stated in the preface of those proceedings, The Oxygen Radicals and Lung Injury Conference was the first of its kind dedicated to pulmonary science. Therefore, in this conference, the primary attempt was to focus on oxygen radicals and their involvement in toxic insults and the ensuing pathological processes in the lung. We did

not ignore the importance of multifactorial relationships of other cellular reactions and products involved in cellular damage and injury. In fact, these issues were addressed in the presentations of Ward (1), Holian et al. (2), Repine (3), Torphy et al. (4), and Demers and Kuhn (5). The complex network of micromolecular reactions have not been fully defined to understand the coordination, modulation, and integration of cellular functions. In many pulmonary diseases (e.g., cancer, emphysema, pneumoconiosis) in which oxygen radicals are implicated, the disease becomes evident only after several years. Subtle damage or changes to biomolecules and their relationships to the coordination and interactions of oxygen radical generation and degradation are important issues to be dealt with in greater detail to understand the synergistic concepts of lung diseases. We hope that future conferences will address these and other issues.

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MTBE: Not Carcinogenic

Subsequent to publication of *EHP*'s timely article on the toxicological potential of methyl-*tert*-butyl ether (MTBE; vol. 103, pp. 666–670), the long-awaited study from the Ramazzini Foundation of Oncology and Environmental Sciences appeared in print (1). This was a landmark publication because for months we in the scientific community had been advised that the data predicted dire health hazards for humans exposed