To the Editor:

Ifosfamide (IFS) is a widely used antineoplastic agent, and the occurrence of ifosfamide-induced hemorrhagic cystitis (HC) continues to be a significant problem in spite of adequate uroprotection. Acrolein, the toxic metabolite of IFS, is the main molecule responsible for this side-effect and mesna (2-mercaptoethane sulfonate) is the most commonly used preventative agent. Mesna binds acrolein and prevent its direct contact with uroepithelium. Current knowledge provides information about the pathophysiologic mechanism of HC. Several transcription factors and cytokines, free radicals and non-radical reactive molecules, as well as Poly (ADP-ribose) polymerase (PARP) activation are now known to take part in its pathogenesis. Whether or not it follows chemotherapy (CP), HC is an inflammatory process. Thus, many cytokines such as tumor necrosis factor (TNF), the interleukin (IL) family, transcription factors such as nuclear factor-κB (NF-κB) and activator protein-1 (AP-1) also play a role in its pathogenesis. When these molecular factors are taken into account, pathogenesis of CP-induced bladder toxicity can be summarized in three steps: (1) acrolein rapidly enters the uroepithelial cells; (2) it then activates intracellular reactive oxygen species and nitric oxide production (directly or through NF-κB and AP-1) leading to peroxynitrite production; (3) finally, the increased peroxynitrite level damages lipids (lipid peroxidation), proteins (protein oxidation) and DNA (strand breaks) leading to activation of PARP, a DNA repair enzyme. DNA damage causes PARP overactivation, resulting in the depletion of oxidized nicotinamide–adenine dinucleotide and ATP, and consequently in necrotic cell death. For more effective prevention against HC, all pathophysiologic mechanisms must be taken into consideration.

Mild forms of HC usually resolve with supportive treatment. However, severe HC may require additional therapies, including hyperbaric oxygen treatment, amifostine, antiviral therapy such as vidarabine or cidofovir, factor VII, bladder irrigation with intravesicular instillation of ε-amincaproic acid, methyl prednisolone or formalin, cystoscopy and cauterization, and even cystectomy. In addition to acrolein, viral infections such as adenovirus and BK virus (1) have been implicated in the etiology of HC. Therefore therapies directed against viral infections may also be useful in the treatment of HC when appropriate. Further studies are needed to determine the appropriate use of antiviral therapy in virus-associated HC.

Despite the preventative use of mesna, HC is observed in 33% of the patients treated with IFS. This observation stresses the need for novel therapies as additional prevention for acrolein induced HC. Mota et al (2) investigated the role of recombinant human interleukine-11 (rhIL-11) in preventing experimental IFS-induced HC in Swiss
mice and have published their results in this issue of the journal. The observations of Mota et al\(^2\) demonstrate that rhIL-11 partially prevents IFS-induced HC, presumably due to its anti-inflammatory properties and ability to down-regulate many pro-inflammatory cytokines. Thus, rhIL-11 may be a very useful adjunct in prevention of HC induced by IFS. These data support that rhIL-11 be used in clinical trials to investigate its role in prevention of HC in humans.

REFERENCES


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