Role of Oxidative Stress and Cytokines in Renal Ischemia Reperfusion Injury

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Abstract

Ischemia reperfusion injury is the principal and the most severe cause of acute kidney injury. This is the result of a generalized or localized degeneration in the transportation of oxygen and nutrients to cells and the removal of waste products from kidney cells. Ischemia reperfusion injury is the main chance element for the development of persistent kidney disorder; this is explained in terms of renal or instantaneous system irregularity for more than 3 months. Throughout various clinical cases, kidneys are exposed to the ischemia reperfusion injury, interpreted by interruption of the kidney blood out glide by continued by successive perfusion development, such as resuscitation circulatory arrest, cardiac and vascular surgery, kidney transplantation and trauma. This article offers the literature's basic information to evaluate the most relevant details about the Oxidative stress and cytokines that act in renal ischemia reperfusion injury.

Keywords: Oxidative stress , cytokines, Ischemia reperfusion injury


Introduction

Renal Ischemia/Reperfusion Injury

Ischemia-Reperfusion Injury (IRI) is the main and commonplace reason of acute kidney injury (AKI). It is the effects of a generalized or localized deterioration of oxygen and nutrient transport to cells and the elimination of waste products from the cells of kidneys (1). The imbalance of local tissues O2 demand, resources also accumulation of wastes output of cells metabolism. Sequence of this imbalance, tubular epithelial cell endure damage, if it is critical dying through necrosis and apoptosis result in ATN, which includes organ feature deterioration in electrolyte, water homeostasis and reduced excretion of wastes output of metabolism. There are massive range of medicinal drugs and path physiological states that could relate to localized or generalized ischemia (2), ischemia-Reperfusion Injury causation of AKI that is recognized with high mortality and morbidity fee with expanded costs of therapy in both adult and pediatric population (3). IRI is the principle chance element for the development of persistent kidney disorder; this is explained as irregularity in renal features or systems instantaneous for more than 3 months (4). Kidneys are uncovered to the IRI in numerous scientific situations, represented via disruption of the kidney blood out glide and persisted by way of the successive improvement of perfusion, like circulatory arrest with resuscitation, cardiac and vascular surgical operation, kidney transplantation and trauma. IRI in kidney transplantation evokes descend of pathophysiology transformation that results in a behind schedule alloimmune precise response to the graft and graft function (DGF), finally ensuing in acute rejection after which expand to chronic allograft nephropathy (CAN) (Figure 1) (5).
**Pathophysiology of Renal Ischemia/Reperfusion Injury**

Inflammation is greater not unusual abnormality related to many mobile species and has a extreme position in pathophysiology of kidney ischemia. Inflammatory cascade brought on in ischemia that is worried in immoderate kidney damage; consequently, blocking off of inflammatory reaction is therapeutic process to guard kidney tissues (6). In the kidney dysfunction, the inflammatory mediators together with interleukin-1β (IL-1β), interleukin-6 (IL-6) and tumor necrosis factor-α (TNF-α) own a full-size function (7-8). Inflammatory mediators recruit Neutrophil infiltration and leukocyte in post ischemic renal tissues, then force to promote leukocyte-endothelial interaction that may permit swelling, damage of the endothelial mobile and physically impair blood glide (9). In addition, there is a discounted vasodilatation in reaction to nitric oxide, bradykinine and acetylcholine (10). Endothelial lesion consequences in loss of glycocalyx, alternation in endothelial mobile-cell contacts, disruption of the acting cytoskeleton, breakdown of the perivascular permeability and lack of fluid into interstitial for the duration of AKI (11). Moreover, intended that the innate immune gadget is reactivated in a very early stage of ischemia/reperfusion lesion as demonstrated through accumulation of macrophages, dendritic cells and Neutrophils observed by the liberation of big amounts of inflammatory mediators (12) and make the tubular cells more vulnerable to ROS (13)

**Oxidative stress and lipid peroxidation**

Oxidative strain response is noticed with the aid of the increase in the amount of interactive oxygen unfastened radicals. ROS are particles of high have interaction with an electron that is not related to the maximum of the outer orbital which make the molecule risky and toxic to the cells. This unfastened radical reasons membrane damage and activates the sign pathway inside cells that finally lead to cell damage and demise (14). The precept deliver of reactive oxygen species is the reduced form of O2. Oxygen reduction occurs via cytochrome oxidase CO enzyme in mitochondrial (ETC) resulting in the formation of hydroxyl radical, H2O2 and superoxide anion (15). There may be no particular intention for reactive oxygen species; however, attacking fats, amino acids and proteins leads to the production of precarious molecules appearing like radical, eventually turning into compounds with a couple of metabolic outcomes. Additionally, the down regulation of antioxidant enzymes like superoxide dismutase, glutathione peroxidase and catalase may be answerable for the pathophysiology of IRI. Preventing this pathway or preventing unfastened radical formation is a technique for tissue safety at some point of ischemia reperfusion (Figure 2) (16).
Chemokines

The chemotactic Chemokines are the cytokines which have been performed a essential performance in the infection and reparation processes afterward RIRI thru the leading of the leukocytes’ immigration and stimulation inside the infection’s places (7).

High-Mobility Group Box-1 Protein

High-mobility institution box-1 protein (HMGB1) is a harm-associated molecular pattern molecule (DAMP), an inflammatory cytokine that has been cautioned to normalize the inflammation in one-of-a-kind tissues (18). It binds to chromatin, however upon signaling it undergoes a nuclear–cytoplasmic translocation and a launch from the renal cells into the venous circulatory machine in a while RIRI. HMGB1 induces a rapid go with the flow in the systemic, and the mobile circulating Chemokines which have been protected TNF-α and other mediators (19). HMGB1 has been said to be expressed thru the renal tissues following IRI (20). An important function of the speedy release for HMGB1 after the mobile injury is suggesting that it can have an vital role on this method (21). As soon as it has been expressed to the circulatory gadget, it is able to interact with special receptors consisting of the superior glycation quit products (RAGE), TLR2 and TLR4 on the goal cells leading to the stimulation of the nuclear aspect kabba B (NFKB) and different molecules (22). It had been showed that the neutralized HMGB1 reduces the releasing of TNFα, IL6, and MCP1 at some point of IRI (19).

Interleukin-1 beta (IL-1β) is produced by means of monocytes and macrophages upon activation. It is a strong proinflammatory cytokine this is regulated by means of the ones immune cells (23). The Mechanisms of irritation were regulated by IL-1β (inflammasomopathies) and NFKB. So, because the NLR family pyrin domain containing three protein (NLRP3) has been interacted with now not simplest the apoptosis-related speck-like protein containing (ASC) but also with the caspase-1. These interactions were shaped a complicated that's referred to as the inflammasome. On cell stage, it's far a large molecule that converts Pro-IL-1β (precursor) into its activated structure (IL-1β) (24). Also, the special cytosolic danger signals translate via Pathogen-associated molecular pattern proteins (PAMPs) and NLR own family pyrin domain containing 1 protein (NLRP1) into the caspase1-established secretion of IL-1β. The moves may be produced through RIRI (25).
Furthermore, its activation has a major function in the Neuroinflammation that has caused the scientific syndrome of AKI. The Mechanisms for IL-1β activation were defined in info within the discern (1.3) (Figure 3) (26).

**Figure (3):** The regulation of IL-1β in the inflammatory diseases which adopted from

**Monocyte Chemoattractant Protein-1**

Monocyte chemoattractant protein-1 (MCP-1) is the key cytokine which has been regulated now not simplest the migration however also the infiltration of monocytes, reminiscence T lymphocytes and herbal killer (NK) cells, it is produced due to constitutive way or due to the inductive method as in OS. The receptors for MCP-1 encompass an extracellular N-terminal region and a C-terminal intracellular area (27). Some of the molecular changes, that associated with tumor, had been protected the increasing of the expression of MCP-1. It tremendously expressed within the breast tumor cells (28). The gathered statistics advise that MCP-1 expression might be accelerated in sufferers of the inflammatory bowel disorder, the allergic allergies and the rheumatoid arthritis (29). Also, the significance of MCP-1 and its receptor is not constrained to the exhibition of the cardio logical problems but may be anticipated in other inflammatory instances along with inside the neurological problems such in which is increased in astrocytic leading to neuronal dying due to ischemic accidents (30). Overall, the skill ability of MCP-1 might also specify that point and spatial factors are involved in indication whether MCP-1 may act as mediator for infection or survival (31).

**Conclusion**

In conclusion, this review showed those different aspects of oxidative stress production, and the expression of different cytokines and biomarkers that may be used in clinical settings and future perspectives in oxidative stress control.

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