Mechanistic Pathways of ATP Sensitive Potassium Channels Referring to Cardio-Protective Effects and Cellular Functions

Authors
Vishal Kumar Vishwakarma¹, Prabhat Kumar Upadhyay², Hridaya Shanker Chaurasiya¹, Ritesh Kumar Srivasatav³, Tarique Mahmood Ansari⁴, Vivek Srivastava¹

Affiliations
1 Department of Pharmacology, R.R.S College of Pharmacy, Amethi, Uttar Pradesh, India
2 Institute of Pharmaceutical Research, GLA University, Mathura, Uttar Pradesh, India
3 Faculty of Pharmacy, Kamla Nehru Institute of Management and Technology, Sultanpur, India
4 Department of Pharmacy, Integral University, Lucknow, Uttar Pradesh, India

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ABSTRACT
A study of potassium channels correlates the fundamentals of mechanistic pathways and various physiological functions. The knowledge of these pathways provides the background, how to determine unit cell functions and to affect cardio protection. ATP sensitive potassium channels adjust excitability of membrane and functions as per metabolic status of cell. A lot of energy consumption primarily occurred in skeletal muscles which also express high number of potassium channels. The increase in calcium release and high heat production is occurred due to loss of potassium channels. Such type of mediations determines metabolic changes and energy required in the dissipation. IPC reduces infarct size in anesthetized mice. In ischemic-reperfusion, pressure in left ventricle was watched while contractile power recovery did not happen. It was seen that elements of intact potassium channel are fundamental for Ischemic preconditioning (IPC). If more prominent is enactment of potassium channels and their cardiologic effects create high heart rate. All the more as of late, it has been suggested that mitochondrial ATP sensitive potassium channels are critical as closing stage effectors which trigger IPC as opposed to sarcolemmal potassium channels. Nevertheless, the importance of the potassium channels reconsidered in cardio-protection in present findings. These discoveries recommend that potassium channels in the adjusting ischemic-reperfusion damage in mice. The heart rate of the mouse occurred during ischemia; enhance watchful extrapolation applied to larger warm blooded animals.

Introduction
ATP-sensitive potassium (K$_{ATP}$) channels have the bend nature to alter membrane properties and control the cell metabolic status [1]. K$_{ATP}$ channels are generally spread in various tissues including cerebrum, pancreas, heart and skeletal muscle.

Different types of these channels add to particular properties of cells and tissues. Skeletal muscle K$_{ATP}$ channels are on a very basic level bound through physical relationship in 4 potassium channel [2]. A potassium channel like Kir 6.1, a weak significant modifier, with 4 working sulfonylurea receptor subunits, SUR2A, and incredibly less articulation of SUR1 and subunits of Kir6.1 [3]. The channel mediated metabolic status occurs via path of the K$^+$ pore ATP sensitivity by subunit of SUR, which play a role in activation of channel by Mg-ADP; potassium channel openers and affected by sulfonylurea drugs. Skeletal muscles and heart are highly significant sites for physical activity where K$_{ATP}$ are found in high frequency [4].
In the heart, direct opening of K\textsubscript{ATP} channel happens in light of various burdens related with either decreased vitality accessibility, for example, hypoxia or ischemia [5], or expanded vitality utilization including expanding heart rate inside the typical, physiological range. The resultant cell potassium efflux advances activity potential shortening, hence restricting the drive for calcium deluge and calcium instigated calcium discharge. This spares cell vitality that would some way or another be used for calcium homeostasis and chocking [6].

An abbreviated activity potential range (AP) furthermore achieves a more broadened diastolic interval, essential for vitality hold renewal. Along these lines, a fundamental bit of ventricular K\textsubscript{ATP} channels is alteration of the AP to streamline cardiovascular limit over a far reaching extent of workloads while avoid fatigue of cell metabolic assets that provoke damage or brokenness [7].

**Involvement of K\textsubscript{ATP} Channel in Ischemic Preconditioning**

IPC is a process by which compact irregular conditions of ischemia impenetrably protect the myocardium against different insults of ischemia, leading to decline of infarct size. At first it was recommended that initiation of sarcK\textsubscript{ATP} channels expect a basic part in IPC. In fact, a number of examinations have shown that IPC was replicated by potassium channel openers [8] and lessening through sulfonylurea K\textsubscript{ATP} channel blocker glibenclamide [9].

Grover et al. (1995) [10] proposed that it was not connected between APD shortening & cromakalamin associated cardioprotection in dogs. The class III threatening to arrhythmic medications Dofetilde and Terikalant which was not targets, each ineffectively to in dogs. The class III threatening to arrhythmic medications Dofetilde and Terikalant should not be done under crazy ischemic conditions.

Certainly both sarcK\textsubscript{ATP} and mitoK\textsubscript{ATP} channels are responsible for the cardio-protection, forgetting about way that several differentiations in their qualities among animal species and exploratory conditions. It has been questionable that sarcK\textsubscript{ATP} or mitoK\textsubscript{ATP} match some regular highlights in IPC [19]. The present examination has demonstrated that sarcK\textsubscript{ATP} channel work is crucial for IPC, which is a high-heart-rate creature assortment in any way in mouse. It was proposed that digoxin, limits sarcK\textsubscript{ATP} channel and conveys intracellular Ca\textsuperscript{2+} over-load by Na\textsuperscript{+} K\textsuperscript{+} ATPase block, repealed IPC in vivo hearts of rabbit [20]. In all terms, it has been represented to that both of mitoK\textsubscript{ATP} and sarcK\textsubscript{ATP} channels in canine heart, expect an essential limit of infarct estimated after IPC mechanism [21] and in erratic pain relieving [22]. The ischemic daamage is higher at standard in KO hearts than in WT hearts may destabilize IPC conventions in the 2 different ways. The Kir6.2-lacking mice push the criticalness of sarcK\textsubscript{ATP} that involves in cardio-security.

**Mitochondrial ATP-sensitive K\textsuperscript{+} Channel and Rodent Cardiomyocytes**

The presence of Kir6.1 protein in coronary supply routes of rodents and cardiomyocytes was predictable with the far reaching assignment of Kir6.1 mRNA of rat heart [23] and in the smooth muscle of veins [24]. The subtypes of Kir6.1 in mitochondria which were considered to results in detached part of mitochondria and myocytic ventricles observed by an examination and localized with MitoFluor red, a mitochondrial pointer [25]. The repression of Kir6.2 in ER as well as cell layer in the cardiomyocytes of rat was related to results got in electron microscopy having neurons and glial cells of cerebrum of rat [26]. The Kir6.2 proteins on ER address the site before the course of action in reasonable octamers associated with sulphonylurea receptors on cell layer [27]. A quality trade procedure did not represent in which mitochondria possess Kir6.1 or Kir6.2 proteins [28]. Additionally, an immunoblot examination without Kir6.1 or Kir6.2 proteins in mitochondria [29].

At that point, it has been kept up both Kir6.1 and Kir6.2 channels were not the parts of mitochondria. The proteins, Kir6.1 and Kir6.2 were available in mitochondria seen by immunoblot examination of mitochondrial parts following electron microscopy [30] and in separated ventricular myocytes in rats utilizing immuno-cytometry. Kuniyasu et al. (2003) [29] utilized approaches to change their antibodies and show specificities by immunoblot examination.
The Role of ATP-sensitive Potassium Channels in Cellular Defense and CVS protection

Two self-decision explore offices can put forth a defense possessing first portrayed the ATP-fragile potassium channels [31]. Some report viewed the nearness of an outside current of K⁺ ions in heart muscle cells when treated with hypoxia or metabolic dangerous substances [32]. It was turned around by ATP imbued into the cell. For all intents and purposes indistinguishable recognitions were made by another social event [33]. Such coordinates were next delineated in pancreatic beta cells, skeletal muscle [34], smooth muscle [35] and neurons. It has clarified that the channel showed electrophysiological properties and pharmacological actions [36]. In back to front fixes in ~140 mM balanced K⁺ obsessions, conductance of single-channel is represented ohm⁻¹ among a conductance of 70–80 pS. The lower regards now and again observed in composition has lower and upside down K⁺ obsessions. The subdue channel development are produced less by other adenine nucleotides without magnesium because they are ground-breaking. On the other hand, Mg²⁺, ATP and ADP are considered to produce stimulatory action. Undoubtedly, it was established that channels were known to have a rich pharmacology [37].

Surprisingly, sulphonylureas were discovered when it was well-known that antimicrobial sulphonamides were also have ability to cause hypoglycaemia and observed in animal models. It was clearly ended up that insulin release induced through beta cells of pancreas due to constraint of KATP channels. As observed in case first age administrator (e.g., tolbutamide, chlorpropamide) and second-age administrators (e.g., glibenclamide, gliclazide, glipizide) which are used in the treatment of diabetes mellitus which opens KATP channels. Furthermore, these administrators particular for KATP channels show extent of substance structures like diazoxide is a benzothiadiazine, pimidinedil a cyanoguanidine and nicorandil a pyridyl nitrate [38].

Characteristic Role of the Mitochondrial ATP-sensitive K⁺ Channel in Cardiac Functioning

Mitochondria are recognized in both bioenergetics and protection in heart against ischemia-reperfusion injury. A broad hypothesis is that the functioning of cardioprotection uses the physiological components of mitoKATP channel. This work shows that the opening of mitoKATP channels has 2 special outcomes for heart which depend upon major bioenergetic process when opening of channel occurs. At the point, when DW is high, as in case of opening of mitoKATP channels in resting conditions, extended mitochondrial ROS creation and ROS constitutes kinases inside a positive banner expansion circle provoking quality interpretation and cell improvement.

This causes the opening of mitoKATP channels lead to cardioprotection against ischemia–reperfusion damage. When, the reduction of DW occurred in ischemia which cause opening of mitoKATP channel leads to homeostasis. Other K⁺ flow through mitoKATP channel lowers fundamental force which is basic for keeping a significant transport of electrons and a low conductance of film VDAC. These results assure improvement the midst of work state as well as reperfusion after ischemia [39].

Mechanism of MitoKATP Channel Opening through Signaling Pathways of IPC and CPC

Cardioprotective effects through IPC and CPC is distressed by mitoKATP inhibitors for example, 5-HD and Glibenclamide. This shows not only that mito KATP is stressed in these techniques for protection, except for those mito KATP channel is opened by endogenous mechanisms [40].

Principally, the cell hyperpolarization occurred in vascular smooth muscle (VSM) is responsible for the opening of KATP channels [41]. A quick phosphorylation of channel subunits Kir6.1/SUR2B (like T633, S1387 and S1465 on SUR2B; S385 on Kir6.1) was observed directly by dynamic sub-nuclear examinations [42]. Additional control may occur through dephosphorylation of these developments for the Ca²⁺-subordinate phosphatase calcineurin [43]. Vasoconstrictors like angiotensin II and endothelin-1, activation of PKC which adjust KATP channel activity through such pathways [44]. The Ca²⁺-programmed through PKCε mediation organizes phosphorylation of Kir6.1 channel which is responsible for gathering of serine developments in distal C-end as main part. There may be outcomes for channel camouflage and reversing possibly by methods for caveolae [45]. The vasoconstrictors interact with PKA results extra inhibitory assurance to KATP channels [46].

One technique that has been little researched is the piece of PIP2 utilization despite PKC initiation. In some cases, it was found that the control is abolished by PKC inhibitors. Kir6.1 channel seems to have a courteously high activity for PIP2 and channel action may be kept with huge utilization. It was proposed that PKC-assisted VSM KATP activity is subject to produce small movement of vasoconstrictors. The vascular KATP channel and cloned Kir6.1/SUR2B equivalents are focused on hormonal control through direct phosphorylation of subunits. The PKC change has been included as central role in cell protection as well as preconditioning in cardiac cells [47]. In early examinations, PKC was thought to start sarcomellemal cardiovascular KATP and there is a bifacile lead with order took after by an inhibitory action contrasting with channel camouflage [48].

In any case, with unrivaled pipette Ca²⁺ in whole cell annals in flawless cells and we saw biphasic control. Inhibitory effect was a direct result of channel mask and happened due to phosphorylation of S372 in Kir6.2 protein [49]. The course of sarcomellemal KATP channels through PKA has been inspected. PKA with Kir6.2/SUR1, phosphorylation prompts increased direct development through stores in SUR1 and Kir6.2 that are homologous to those in SUR2B and Kir6.1. One charming component is the subcellular control of KATP Channels which radiate an impression of being gathered at the fore-end of T-tubule, putting forward commencement could affect excitation pressure coupling [50].

Mitochondria are a quantitatively significant wellspring of ROS, which supply to tissue harm amid ischemia, but on the other hand are middle people of IPC flagging [51]. Gathering affirmation recommends that redox flagging pathways assume an essential part in IPC [60, 61], and ready to advance mKATP actuation [52]. The es-
sential ROS created by means of mitochondria is superoxide (O$_2^•−$) [53], even as hydrogen peroxide (H$_2$O$_2$) or lipid peroxides can be shaped optionally. Both O$_2^•−$ and H$_2$O$_2$ are thought to actuate mKATP[54], while clashing reports exist with respect to O$_2$. The impact of extra peroxides on mKATP isn’t known. In addition, it is evident that a few yet not a wide range of cancer prevention agents can lessen IPC and mKATP action, justifying more examination mKATP Channel. Nitric oxide (NO•) is additionally worried in IPC and inspires a major range of cardioprotective effects [55]. NO• have been recognized in the separated mitochondrial courses of action [56] and can alternatively provide various responsive nitrogen species [57] which can provide either harmful or supportive fading parts [58].

The mKATP may be a likely center for such RNS, and whereas while high doses (10 mM) of a S-nitroso-thiol (SNT) have been displayed to activate directly in the faultless mitochondria and affirmation for more physiologically pertinent effects of NO• has observed for the foremost portion depended on the circuitous measures of channel movement [59] or examination of the channel ousted from its mitochondrial condition [60].

Biophysical Characteristics of K$_{ATP}$ Channels in Managing Vascular Diseases

Potassium channels vascular infections changed vascular potassium channel work beneath neurotic conditions may well be either a reason or an affect of contamination. Vasoconstriction and course of action that of a vein limit to broaden are after effects of damaged K$_1$ channels restrain in veins and may be direct result of an alteration in number, basic conductance, and open validity of the channel(s).

All together at this level with respect to K$_1$ channel articulation can starting at now be given just by ponders atomic and also settle prop approaches and thusly, are limited to examinations of veins or cardio-myocytes are abolished from the physiological condition [61]. In this manner, where probability permits (i.e., the accessibility of specific pharmacological instruments), it will stay basic to check such revelations with data about vascular K$_1$ coordinate limit procured in the joined physiological condition in vivo (▶ Fig. 1).

Particularly few circulated examinations have been broken down impact of infection on biophysical features of K$_1$ channels or sub units. As shown above, the greater piece of our understanding into affect of illnesses on vascular K$_1$ channel articulation remains underhanded and subject to clarification of the test information attained by utilizing K$_1$ channel modulators that are accepted to be modestly particular pharmacological specialists [62].

Contribution of K$_{ATP}$ Channels in Cerebro-Vascular Hypertension

The persistent hypertension is the most frequent thought about cardiovascular infectivity state similarly as its effects on vascular K$_1$ channel exertion. At present, there is unpredictable working of each one of the 4 vital K$_1$ channels makes in midst of Effects out of Hypertension on electrical membrane (Em). The resting Em of vascular smooth muscle cells is represented to be more depolarized in veins from hypertensive versus normotensive animals. Extended
vascular depolarization of vessels is related with overhauled myogenic tone in hails from hypertensive animals [63]. Few examinations propose that the restrain of vascular K<sub>ATP</sub> channels is prevented within the middle of hypertension. Existing K<sub>ATP</sub> Channel activators are less sensible dilators invivo in both enormous [64] and small cerebral vessels of persistently hypertensive rats [65]. This alteration shows up to most likely consolidate a weakened layer hyperpolarization response to these administrators, as essentially indistinguish-able disclosures from settle catch contemplates demonstrate that a glibenclamide-unequaled K<sub>i</sub> current start- ed by levcromakalim is reduced in mesenteric entry in smooth muscle cells of tenaciously hypertensive animals [66]. Cromakalim provoked extricating up of isolated mesenteric course is unclear prevented in case of hypertension initiated by unending NO synthase prevention [67].

The vascular K<sub>ATP</sub> channels are accepted to be inactive under most normal basal conditions and amplified vascular tone within the middle of perpetual hypertension is improbable to be related to disabled K<sub>ATP</sub> Channel effort. For improved K<sub>ATP</sub> channel effort, in continual hypertension, an extended vasodilator effect of cromakalim and significant choking of channel by glibenclamide in carotid ways from stroke-slanted SHRs against normotensive WKY rats has been considered [68].

**K<sub>ATP</sub> Channels in Diabetes and Resulting to Hypertension in Vascular Bed**

Most of information right now open for vascular K<sub>i</sub> occupies exertion in diabetes involving K<sub>ATP</sub> channels. While for never-ending hypertension, different reports of debilitated vascular relaxant properties produced to open K<sub>ATP</sub> channels which deal diabetes.

These examinations have for the mosly exploited the streptozotocin induced rat model of diabetes and have explored vessels at 2.5–4 months coming approximately to treatment with streptozotocin. In this method, the plasma glucose level are increased by 3 to 4-times; debilitated extricating up of the segregrated aorta [69] and mesenteric vascular bed [70] and decreased dilatation of far reaching [71] and little [72] cerebral passages in vivo. These developments are accepted to be the possible result leads decrease in number of K<sub>ATP</sub> channels and moreover decreased atractibility of said channels design openers. Redirection of streptozotocin induced cytotoxic effects of show up a fantastical purpose behind these movements in light of the way that, as various indications of vascular brokenness, flighty vasodilator actions through K<sub>ATP</sub> channel opening are neutralized by prescriptions that keep away from hyperglycemia.

Streptozotocin aggravated diabetes can essentially alter utilitarian response of K<sub>ATP</sub> facilitates in different tissues, with pancreatic beta-cells [73] and ventricular myocytes, [74] showing that hyperglycemia started debilitation of K<sub>ATP</sub> channels is not compelled to vasculature. Since, diabetes is characterized with raised levels of cholesterol, LDL and triglycerides in plasma. It is conceivable that some vascular difficulties of diabetes are not directly linked to hyperglycemia which is a resultant of balanced plasma lipid profile [75].

Period of exploratory hyperglycemia has each one of the saves of being indispensable parameters observed impacts of diabetes on vascular K<sub>ATP</sub> channel work in light of the way that that by disparity, reactions to K<sub>ATP</sub> channel institution are represented to be dominating in the early diabetic state (→Fig. 2). For illustration, cromakalim actuated dilatation of colossal coronary supply courses in poose are extended multi week behind treatment with alloxan, and responses of the small coronary paths are unaffected [76]. Additionally, glibenclamide produce stamped narrowing of vessels. The extended articularion and basal order of K<sub>ATP</sub>-channels occur together in the renal stream in front of schedule in midst of diabetes [77].

This condition can include to the upgrades in glomerular filtration rate and renal plasma stream, which develop in in beginning times of diabetes in both clinical and test settings [78]. An expanded K<sub>ATP</sub> Channel activity right directly may along these lines reflect a elevated metabolic state (i. e., low ATP levels) of smooth muscle cells not long after the starting of hyperglycemia. Extended K<sub>ATP</sub> arrange activity in veins through metabolic weight, for occasion, within the middle of ischemia, may important for keeping up tissue perfusion.

In this way, tissues may well be additional powerless to the ischmic heart after expanded circumstances of diabetes inferable from debilitated limit of K<sub>ATP</sub> Channels. On the other hand, differentiated impacts on K<sub>ATP</sub> channel work are represented to occur 2 months after after affirmation of hyperglycemia, most likely to a limited degree reflecting a ceaseless drop of vascular systems the middle of improvement of disease.

Since, Bouchard et al. reported that vasorelaxant response to lemakalim is weakend in coronary vessels, not in the vascular bed following 2 months of hyperglycemia. Pinacidil-started hyper-polarization of the mesenteric vein was represented to be protected consequent to 8 to 12 weeks. Zimmerman et al. [79] revealed that dilator activities leads to K<sub>ATP</sub> opening were obstructed in cerebral courses from 4 to 8-week-diabetic rats coming about from lessened basal complimentary of endothelium-induced NO. The K<sub>ATP</sub> channels restrain was re-established in diabetic vessels associated with consistent NO support [80].

Of course, such an instrument of endothelial NO subordinate K<sub>ATP</sub> Channel activation couldn’t occur in different stages, as no certification of an area was found for NO dilator activities in rodent cerebral vessels through K<sub>ATP</sub> channel opening [81].

**Hypercholesterolemia, Atherosclerosis, Cardiomyocyte Swelling and Heart Failure**

It is particularly seen that vascular blocking occurred in atherosclerosis and hypercholesterolemia. Such contamination is associated with handicap of endothelial limit. The diminished vascular enlargement of endothelium-affected NO is presumably going to participate a vital part in change of atherosclerosis [82]. As a result, supply courses may exhibit an extended vascular tone and may react insufficiently to endothelium-subordinate vasodilators. Under such condition of changed vascular reactivity, K<sub>i</sub> channel development or limit may similarly be unordinary [83].

The hypertrophy of heart is enacted by an expanded in addition to cardiovascular workload. While, transverse aortic fixing was apply in mice with heart specific over expression of SUR1 which limitless disturbs cardiovascular sarcolemmal K<sub>ATP</sub> channel utility, extended hypertrophy of left ventricle was viewed [84]. Inquisitively, there is
in every way a correspondence between cardiovascular \( K_{ATP} \) channel enunciation and the action of the PPAR-\( \gamma \) co-activator, PGC-1\( \alpha \). Reduced channel reason prompts less activity at PGC-1\( \alpha \) promoter to some degree through FOXO-1 concealment. Restored ventricular cardiomyocytes are subjected to coronary hindrance in rats show up control of KIR6.1, in the demand of the infarct zone [85]. Congestive heart dissatisfaction or restricted corruption in human hearts prompts an upgraded AP term and affectability to potassium channel openers in the 2 atria and ventricles [86].

Angiotensin II and TNF-\( \alpha \) enunciation is insistently related to that of KIR6.1 in failing rat myocardium or created cardiomyocytes and antagonistically related with KIR6.2 [87]. Moreover, cardiomyocytes were treated with diazoxide, exhibiting extended enunciation of KIR6.1/SUR2B in these cells as a noteworthy part of hypertrophy development. Over the best changes in cell volume in the heart item would have capacity to alter the essential cell credibility, cell limits and cell destruction. These movements can raise because of an intracellular metabolite sign that extension cell osmolality, empower water to enter the cell, augment the cell volume and alter molecule channel work. \( K_{ATP} \) channel has been seemed to occur in cellular changes with atrial \( K_{ATP} \) coordinates opening in view of cell swelling provoking AP shortening [88]. The non-attendance of KIR6.2 in cardiovascular cardiomyocytes led to cell swelling while in WT mice cell swelling exaggerated which is bothered by extension of diazoxide [89].

Cardioprotective Effect of Ischemic Preconditioning in Ovariectomized Rat Heart

The affirmation of caveolin, a membrane protein and a unsafe controller of nitric oxide (NO), saises after menopause Examination was anticipated to choose the impact of daidzein (DDZ), a phytoestrogen is included to appear cardioprotective effect of ischemic preconditioning (IPC) in ovariectomized rat heart.

IPC helped cardioprotection was broadly fixed in ovariectomized rats when stood out from run of the process rats, which recovered, was reestablished by the treatment of DDZ, a caveolin inhibitor (0.2 mg/kg subcutaneously) for for numerous weeks. In any case, this observed cardioprotection was amazingly diminished by perfusion of L-nitroarginine methyl ester, an endothelial nitric oxide synthase (eNOS) inhibitor (100 mM/L) and glibenclamide, an adenosine triphosphate-fragile potassium channel blocker (10 mM/L) alone or in mix, noted that expansion in myocardial infarct assess, landing of LDH and CK-MB, and other than diminish within the section of NO.

Opening of mitochondrial \( K_{ATP} \) channels shields the myocardium from I/R-affected damage [90]. Different mediators like adenosine, bradykinin, angiotensin, prostaglandins and NO, which are increased in IPC, pass on cardioprotection by the opening of mitochondrial \( K_{ATP} \) channel [91, 92]. Further, perfusion of glibenclamide, a \( K_{ATP} \) involve blocker in DDZ-pretreated ovariectomized rat heart in a general sense, dropped the cardioprotective effect of IPC without influencing the IPC-intervened entry of NO.

Fig. 2 Schematic depiction of the affect of SUs on pancreatic \( \beta \)-cells and cardiovascular myocytes. Sulfonylurea tie to sulfonylurea receptor proteins (SURs), subunits of the KATP channels.
It is discretionary that the watched cardioprotective impact of IPC in rodent and DDZ-pretreated rodent may be a coordinate result of opening of mitochondrial \(K_{\text{ATP}}\) channel. Also, perfusion of L-NAME with Gilbenclamide in DDZ-pretreated ovariectomized rodent heart was not capable to create any extra effect in examination with specific drugs. These revelations reflect that NO conveyed in light of IPC in DDZ-pretreated ovariectomized rodent heart make cardioprotection by the opening of \(K_{\text{ATP}}\) channels [93].

**Conclusion**

In summary, we have demonstrated the role of the potassium channel in mediating delayed ischemic preconditioning. It is confirmed that blockers were acting at potassium channel. Further studies will be necessary to unravel the specific signal transduction mechanism by which delayed preconditioning leads to the opening of these channels and differentiating the specific role played by each of the sarcolemmal and mitochondrial potassium channels in mediating delayed cardio protective effect in vivo.

**Clinical Implications**

It has been indicated from various disorders that mitochondrial ATP sensitive potassium channels are critical which trigger IPC as opposed to sarcolemmal potassium channels. The increment in potassium channel mediations decides metabolic changes and energy required within the dissipation.

These channels can be determined with a role in pharmacological and biophysical properties. Nevertheless, the importance of the potassium channels reconsidered in cardio-protection in present findings. The usefulness of such studies indicates a fundamental role of potassium channels in a variety of CVDs disorders including arrhythmias, hypertension and heart failures.

These findings suggest that potassium channels within the adjusting ischemic-reperfusion injury in mice. The heart rate of the mouse occurred during ischemia and probably enhances vigilant extrapolation applied to larger warm blooded animals.

**Limitations of Study**

A study of mechanistic pathways mediating potassium channels provides the knowledge of cell functions and cardio protection and limits to cardiovascacular functions and pathways. ATP sensitive potassium channels adjust excitability of membrane and control metabolic functions. Limitations of this study include missing of experimental protocols with justification and involvement of potassium channels. A lot of advancements in this field those are not proved till date which are not included. A lack of wide versatility of potassium channels in the study for physiological functions.

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**Conflict of Interest**

Authors did not have any conflict of interest in writing and publishing the manuscript.

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