In the early 1980s, we proposed a unifying model for β-cell damage (The OKAMOTO model), in which poly(ADP-ribose) synthetase/polymerase (PARP) activation plays an essential role in the consumption of NAD+, which leads to energy depletion and necrotic cell death. In 1984, we demonstrated that the administration of PARP inhibitors to 90% depancreatized rats induces islet regeneration. From the regenerating islet-derived cDNA library we isolated Reg (Regenerating Gene) and demonstrated that Reg protein induces β-cell replication via the Reg receptor and ameliorates experimental diabetes. More recently, we showed that the combined addition of IL-6 and dexamethasone induces the Reg gene expression in β-cells and that PARP inhibitors enhance the expression. In 1993, we found that cyclic ADP-ribose (cADPR), a product synthesized from NAD+, is a second messenger for intracellular Ca²⁺ mobilization for insulin secretion by glucose, and proposed a novel mechanism of insulin secretion, the CD38-cADPR signaling system. Therefore, PARP inhibitors prevent β-cell necrosis, induce β-cell replication and maintain insulin secretion.

Key Words: Poly(ADP-ribose) synthetase/polymerase (PARP), Cyclic ADP-ribose, Reg gene, The OKAMOTO model, Necrosis
In this paper, we would like to present a perspective view based on our studies concerning cell death, cell regeneration, and cell function, especially on insulin-producing pancreatic β-cells, in the processes of which poly(ADP-ribose) synthetase/polymerase (PARP) and cyclic ADP-ribose (cADPR) are functioning.

**PANCREATIC β-CELL DEATH BY PARP ACTIVATION**

In 1981, we published two papers, one of which was concerned with *in vitro* experiments [1] and the other with *in vivo* experiments [2], describing that streptozotocin and alloxan induce DNA strand breaks and PARP in pancreatic islets, and proposed a unifying model for the action of diabetogenic agents, streptozotocin and alloxan, on pancreatic β-cells [3]. Central to the model are breaks in the nuclear DNA of β-cells, resulting from either an accumulation of free radicals or from alkylation of DNA. These breaks induce DNA repair involving the activation of PARP, which uses cellular NAD⁺ as a substrate. As a result, the intracellular levels of NAD⁺ fall dramatically. The fall in cellular NAD⁺ inhibits cellular functions including insulin synthesis and secretion, and thus the β-cell ultimately dies. Thus, this appears to be a suicide response for β-cells to repair DNA. The NAD⁺ depletion and the decrease in β-cell functions, induced by alloxan and streptozotocin, were shown to be prevent-
ed by radical scavengers such as superoxide dismutase and catalase and by PARP inhibitors [4, 5].

Interest in the model for the mechanism of action of alloxan and streptozotocin has been heightened by its possible extension to the effects of viruses and inflammation, especially immune-mediated events on β-cells [3, 6-10]. Thus, since the early 1980s, we have thought that, although type 1 (insulin-dependent) diabetes can be caused by many different agents such as immunologic abnormalities, inflammatory tissue damage, and β-cytotoxic chemical substances, the final pathway for the toxic agents is the same (Figure 1). This pathway involves DNA damage, PARP activation, and NAD⁺ depletion. The fall in cellular NAD⁺ inhibits cellular activities. Therefore, type 1 (insulin-dependent) diabetes is theoretically preventable by suppressing immune reactions, scavenging free radicals, and inhibiting PARP by nicotinamide and 3-aminobenzamide. Concerning nitric oxide (see Figure 1), we produced transgenic mice expressing nitric oxide synthase constitutively in pancreatic β-cells and found that the β-cell mass was markedly reduced and that the transgenic mice developed severe diabetes [11]. In 1999, using PARP deficient mice, three independent groups in Germany, Japan and U.S.A. provided irrefutable support for the model shown in Figure 1: PARP deficient mice were remarkably resistant to streptozotocin and did not show the β-cell death [12-15]. More recently, many
other tissues and cells have been reported to die by the same mechanism as in pancreatic β-cell death [16-35].

The cell death caused by PARP activation described above is thought of in terms of necrosis [36]. In apoptotic cell death, PARP is cleaved by caspases and inactivated. Therefore, PARP inhibitors can be effective in preventing necrosis but ineffective in preventing apoptosis (Figure 2). “Whether to die from necrosis or to die from apoptosis” may depend on the severity and duration of the cell damage, differences in death signals, and the species of cells. A recent report from Bhardwaj’s laboratory suggests that dendritic cells distinguish between two types of cell death, with necrosis providing a control that is critical for the initiation of immunity [37]. Therefore, immunological abnormalities, which are frequently observed in type 1 diabetes, may be triggered by the preceding necrotic cell death, and then cause apoptotic death of β-cells.

**β-CELL REGENERATION AND REG GENE**

As described above, alloxan and streptozotocin diabetes can be prevented by PARP inhibitors. Concerning experimental diabetes, at the end of the 19th century von Mering and Minkowski in Strasbourg found that a dog became glycosuric and hyperglycemic by pancreatectomy. This observation stimulated many workers to try to isolate the active pancreatic principle as a possible treatment for diabetes. In 1984, we demonstrated that PARP inhibitors induce the regeneration of pancreatic β-cells, thereby ameliorating surgical diabetes [38]. Male Wistar rats were 90% depancreatized, and nicotinamide or 3-aminobenzamide was injected intraperitoneally every day. The administration of PARP inhibitors ameliorated the surgical diabetes, and the islets in the remaining pancreases of rats that had received PARP inhibitors for 3 months were extremely large, and almost the entire areas of the enlarged islets were stained for insulin.

We isolated the regenerating islets and constructed a cDNA library. In screening the regenerating islet-derived cDNA library, we came across a novel gene expressed in regenerating islets. The cDNA had one large open reading frame which encoded a 165-amino acid protein. The deduced protein has a signal sequence. We propose to name the novel gene Reg, that is, regenerating gene, with the implication that the gene may be involved in islet regeneration [39]. We subsequently isolated human REG gene [39, 40]. Rat Reg protein increases [3H]thymidine incorporation in rat islets, and mitosis was often observed [41]. We intraperitoneally injected rat Reg protein (1 mg/kg/day) to 90% depancreatized rats. On the 30th and 60th postoperative day, the fasting plasma glucose level of the rats receiving Reg protein was significantly lower than that of the 90% depancreatized control rats. After 2 months, almost all the islets of the 90% depancreatized control rats were destroyed. In contrast, the islets of the remaining pancreas in the Reg protein-treated rats were enlarged and the enlarged islets were densely and almost entirely stained for insulin [41]. These results indicate that Reg protein stimulates the regeneration and/or growth of pancreatic β-cells, thereby ameliorating the surgical diabetes.

Recently, we isolated a Reg protein receptor cDNA from a λZAP II rat islet cDNA expression library [42]. The cDNA encoded a 919-amino acid protein, and the amino acid sequence suggested that the protein is a type II transmembrane protein with a long extracellular domain. We also isolated a human cDNA that shows over 97% amino acid identity to the rat homologue. The rat Reg receptor-expressing CHO cells bound rat Reg protein with high affinity (Kd = 4.4 nM). The binding of 125I-labeled rat Reg protein was displaced by
increasing the concentration of unlabeled rat Reg protein. Human REG protein, which shows 70% amino acid identity to rat Reg protein, also bound to the CHO cells ($K_d = 14.0$ nM), but higher concentrations of human REG protein were required for the displacement of the rat Reg protein. We established several cell lines of RINm5F cells overexpressing the Reg receptor. The cell lines showed significant increases in BrdU incorporation in the presence of 0.3-100 nM rat Reg protein. Moreover, the cell numbers were increased in response to Reg protein. The receptor mRNA was expressed in normal pancreatic islets, regenerating islets and a pancreatic ductal cell line, ARIP cells, that proliferate in a Reg protein-dependent manner.

The receptor mRNA expression was unchanged during islet regeneration [42]. This suggests that the regeneration and proliferation of pancreatic β-cells are primarily regulated by the Reg gene expression. Accordingly, the transcriptional activation is of great importance in β-cell regeneration. More recently, we found that Reg gene is activated by interleukin-6 (IL-6), dexamethasone, and PARP inhibitors [43]. The combined addition of IL-6 and dexamethasone increased the Reg mRNA level, and further addition of nicotinamide or 3-aminobenzamide increased the mRNA even more. Progressive deletion of the 5'-flanking region of rat Reg gene revealed that the region between nucleotides -81 and -70 is essential for the Reg gene promoter activity. The sequence is “TGCCCCCTCCCAT”. Similar GC box-like sequences were also observed in mouse and human Reg genes. The site-directed mutated luciferase construct “TGCCCCCTAACAT” abolished the induction. The mutant (“TGCC-CGGCCCAT”), which changed the sequence to a GC box, and the mutant (“TGCCCCACC-CAT”), which changed the sequence of the rat Reg promoter to those of human REG genes, REG Iα [40] and REG Iβ [44], showed the induction. In gel mobility shift assays (GMSA) with the GC box-like sequence, the intensity of the band, which was detected in the nuclear extracts of RINm5F cells treated with IL-6, dexamethasone and/or nicotinamide, was correlated with the luciferase activity [43]. The addition of NAD+ to nuclear extracts attenuated the band, and nicotinamide and 3-aminobenzamide quenched the effect of NAD+. These results suggest that PARP participates in the formation of the active transcriptional DNA/protein complex and that the formation of the active complex was inhibited by the poly(ADP-ribosyl)ation of nuclear proteins. The involvement of PARP in the active transcriptional complex was evidenced by the fact that the active transcriptional complex was stained by an anti-PARP antibody after GMSA analysis. The involvement of PARP in the active complex was further evidenced by the immunodepletion of PARP [43]. Southwestern experiments showed that a 113 kDa nuclear protein, the molecular weight of which corresponds to PARP, bound the GC box-like sequence. The band was recognized by the antibody to PARP. In fact, a purified recombinant PARP bound the cis-element. When nuclear extracts were incubated in the GMSA reaction in the presence of [α-32P]NAD+ and the reaction products were analyzed, only PARP was labeled. This suggests that PARP in the transcriptional complex auto-poly(ADP-ribosyl)ates itself.

Thus, as shown in Figure 3, inflammatory mediators, IL-6, and glucocorticoids induce the formation of an active transcriptional complex for Reg gene, in which PARP is involved, and the Reg gene transcription proceeds. On the other hand, during inflammation, superoxide ($O_2^-$) and nitric oxide (NO·) are produced and cause DNA damage. In this case, PARP is activated by DNA nicks for the DNA repair. Then, PARP poly(ADP-ribosyl)ates PARP itself, the poly(ADP-ribose) chains on the PARP protein inhibit the formation of the active transcriptional complex, and the Reg gene transcription
is stopped. In the presence of PARP inhibitors such as nicotinamide, the PARP is not poly(ADP-ribosyl)ated, the transcriptional complex is stabilized, and the Reg gene transcription proceeds. Therefore, PARP inhibitors keep PARP active as a transcription factor for β-cell regeneration. This can account for the previous observation of islet regeneration in 90% depancreatized rats treated with PARP inhibitors [38] and also supports our previous proposition that the restriction of β-cell replication is relieved by PARP inhibitors [6]. When DNA is massively damaged, PARP is rapidly activated to repair the DNA, as mentioned in the first part of this paper, and the complex for Reg gene transcription is not formed at all.

Recently, Reg and Reg-related genes have been isolated and revealed to constitute a multigene family, the Reg gene family [44-63]. Based on the primary structures of the Reg proteins, the members of the family are grouped into three subclasses, type I, II, III [45, 52]. Type I (and Type II) Reg proteins, about which we have discussed above, are expressed in regenerating islets [45]. Type III Reg proteins have also been suggested to be involved in cellular proliferation in intestinal cells, hepatic cells, and neuronal cells. In fact, a Cambridge group reported that mouse Reg III is a Schwann cell mitogen accompanying the regeneration of motor neurons [64], and a French group recently reported that Reg protein functions as a neurotrophic factor for motor neurons [65]. A Kyoto group reported that regenerating gene
protein may mediate the gastric mucosal proliferation induced by hypergastrinemia in rats [66-68]. The expression of Reg protein receptor mRNA was also detected in liver, kidney, stomach, small intestine, colon, adrenal gland, pituitary gland, and brain [42], suggesting the possible involvement of the Reg protein-Reg receptor signal system in a variety of cell types other than pancreatic β-cells.

**THE CD38-cADPR SIGNAL SYSTEM FOR INSULIN SECRETION IN β-CELLS**

cADPR is synthesized from NAD⁺, and our results have shown that cADPR is a second messenger for intracellular Ca²⁺ mobilization for insulin secretion in pancreatic β-cells. Therefore, decreases in the NAD⁺ level (see Figure 1) can cause decreases in cADPR and then in insulin secretion.

Glucose induces an increase in the intracellular Ca²⁺ concentration in pancreatic β-cells of the islets of Langerhans to cause the secretion of insulin. This increase in the Ca²⁺ concentration was first explained in 1984 by the hypothesis of Ashcroft et al. of Oxford University [69], in which Ca²⁺ is provided extracellularly. That is, millimolar concentrations of ATP, produced in the process of glucose metabolism,
inhibit the potassium channel, inducing membrane depolarization and the opening of the voltage-dependent Ca\textsuperscript{2+} channels. In 1993, we proposed another model of insulin secretion by glucose via cADPR-mediated Ca\textsuperscript{2+} mobilization from an intracellular Ca\textsuperscript{2+} pool, the endoplasmic reticulum [70], as shown in Figure 4. That is, ATP inhibits the cADPR hydrolase of CD38, causing the accumulation of cADPR, which acts as a second messenger for Ca\textsuperscript{2+} mobilization from the endoplasmic reticulum for insulin secretion. The first important issue is whether the accumulation of cADPR is actually caused by glucose stimulation in pancreatic islets. We incubated normal rat (Wistar) and mouse (C57BL/6J) islets with low (2.8 mM) glucose and high (20 mM) glucose, and assayed the cADPR content in the islets by radioimmunoassay using an anti-cADPR antibody. The cADPR content of islets incubated with high glucose was increased within 5 min, whereas the cADPR content of islets incubated with low glucose was not [71]. Next, we used rat pancreatic islet microsomes as a cell-free system to study Ca\textsuperscript{2+} release and found that cADPR released Ca\textsuperscript{2+} from islet microsomes, as indicated by the observed prompt increase in fluo 3 fluorescence [70, 71]. Inositol 1,4,5-trisphosphate (IP\textsubscript{3}) did not cause the release of Ca\textsuperscript{2+}, and at this point, the islet microsomes were still responsive to cADPR. We then used rat cerebellum microsomes. IP\textsubscript{3} caused a release of Ca\textsuperscript{2+}.
from cerebellum microsomes. cADPR also caused a release of Ca\(^{2+}\). Heparin, an inhibitor of IP\(_3\) binding to its receptor, blocked the IP\(_3\)-induced Ca\(^{2+}\) release from cerebellum microsomes, but did not block the cADPR-induced Ca\(^{2+}\) release. These results indicate that islet microsomes respond to cADPR but not to IP\(_3\). In contrast, cerebellum microsomes respond to both cADPR and IP\(_3\), but cADPR induces the Ca\(^{2+}\) release via a different mechanism than that utilized by IP\(_3\). We then examined the effect of cADPR on insulin secretion using digitonin-permeabilized pancreatic islets. cADPR as well as Ca\(^{2+}\) induced insulin secretion, but IP\(_3\) did not. The combined addition of cADPR and Ca\(^{2+}\) did not induce significantly more insulin secretion than the addition of cADPR or Ca\(^{2+}\) alone. The cADPR-induced insulin secretion was inhibited by the addition of EGTA. These results suggested that the cADPR-induced insulin secretion was mediated by Ca\(^{2+}\) mobilization from islet microsomes [70]. Thus, we proposed that glucose stimuli induce cADPR formation from NAD\(^+\), cADPR then mobilizes Ca\(^{2+}\) from the endoplasmic reticulum, serving as a second messenger for insulin secretion.

The next question is why the glucose stimulus induces the formation of cADPR. CD38 is a 300-amino acid protein and was first recognized as a leukocyte antigen. We found that CD38 is expressed in a variety of tissues including pancreatic \(\beta\)-cells [72, 73]. We and others have found that CD38 has both ADP-ribosyl cyclase, synthesizing cADPR from NAD\(^+\), and cADPR hydrolase to produce ADP-ribose [72, 74, 75]. We purified human CD38 protein and found that millimolar concentrations of ATP inhibit the cADPR hydrolase activity of CD38, competing with the substrate, cADPR [76]. The competitive inhibition of the cADPR hydrolysis by ATP suggests that ATP and cADPR bind to the same site of CD38. We then labeled the purified CD38 with an ATP analogue, 5'-p-fluoroosulfonylbenzoyladenosine, and identified the binding site for ATP and/or cADPR as the lysine-129 of CD38 [76]. From these results and other available evidence, we proposed that CD38 catalyzes the formation of cADPR from NAD\(^+\) and also the hydrolysis of cADPR to ADP-ribose. As shown in Figure 5, lysine-129 of CD38 is the cADPR binding site, and ATP competes with cADPR for the binding site, resulting in the inhibition of the hydrolysis of cADPR and then, in the accumulation of cADPR [76]. Cysteine-119 and Cysteine-201 are essential for the hydrolase reaction [77], and glutamic acid-226 for the NAD\(^+\) binding [78].

cADPR has been thought to activate the ryanodine receptor to release Ca\(^{2+}\) from the intracellular stores, the endoplasmic reticulum [70, 79, 80]. We have shown that the type 2 ryanodine receptor is expressed in rat pancreatic islets [71]. Our experiments indicated that cADPR did not bind directly to the ryanodine receptor but may act on the receptor through a mediator such as FK506-binding protein 12.6, FKBP12.6, to release Ca\(^{2+}\). FK506 is one of the most widely used immunosuppressive agents. The cellular target for FK506 is thought to be FKBP12 and FKBP12.6. Rat FKBP12 is composed of 108 amino acids and is highly conserved among human, mouse, bovine, and rabbit FKBP12. Rat FKBP12.6 is also a 108-amino acid protein as are human and bovine FKBP12.6. Rat islet microsomes contained FKBP12.6, but did not contain FKBP12. It is of great interest that cADPR was found to bind to FKBP12.6 at a \(K_d\) value of 35 nM. The binding of radiolabeled cADPR was inhibited by cold FK506 as well as cADPR and neither structurally nor functionally related analogues of cADPR inhibited the binding to FKBP12.6 [81]. These results indicate that FKBP12.6 acts as a cADPR-binding protein and strongly suggest that cADPR is the actual ligand for FKBP12.6 since FK506 does not nor-
mally exist in mammalian cells. FKBP12.6 occurs in rat islet microsomes. However, when rat islet microsomes were treated with cADPR, FKBP12.6 dissociated from the microsomes and moved to the supernatant, releasing Ca\textsuperscript{2+} from the intracellular stores [81]. From these results together with other experiments, it is strongly suggested that, when cADPR binds to FKBP12.6 in the ryanodine receptor and causes the dissociation of FKBP12.6 from the ryanodine receptor to form the FKBP12.6-cADPR complex, the channel activity of the ryanodine receptor is thereby increased to release Ca\textsuperscript{2+} from the endoplasmic reticulum. As you can also see in Figure 4, when FK506 is present, cADPR cannot act on the ryanodine receptor to release Ca\textsuperscript{2+} and the glucose-induced insulin secreting machinery ceases to function. In fact, when FK506 was used as an immuno-suppres-sant in kidney transplantation, hyperglycemia was observed in 20-35 per cent of the recipients [82, 83]. The diabetogenic side effect of FK506 may be explained by the mechanism shown in Figure 4. Furthermore, in the presence of calmodulin, islet microsomes were sensitized to cADPR at much lower concentrations for Ca\textsuperscript{2+} release, and the Ca\textsuperscript{2+} release was greatly increased [84, 85]. These results and other available evidence suggest that the cADPR-mediated Ca\textsuperscript{2+} mobilization for insulin secretion is achieved by the calmodulin-activated CaM kinase II. Possibly, the activated kinase phosphorylates the ryanodine receptor to sensi-tize the Ca\textsuperscript{2+} channel for the cADPR signal.

To verify a novel mechanism of insulin secretion, the CD38—cADPR signal system, we created CD38 knockout mice [86]. The pancreatic islets of CD38 knockout mice showed almost no ADP-ribosyl cyclase activity. The glucose-induced increase in the intracellular Ca\textsuperscript{2+} concentration was severely impaired in the knockout mouse islets, and the glucose-induced insulin secretion was severely decreased. The knockout islets, however, responded normally to the extracellular Ca\textsuperscript{2+} influx stimulants tolbutamide and KCl to secrete insulin [86, 87]. This suggests that the CD38—cADPR signal system functions in the Ca\textsuperscript{2+} mobilization from intracellular Ca\textsuperscript{2+} stores. The paradigm of insulin secretion based on the CD38—cADPR signal system, so far described, relies on a wide body of evidence obtained in rat and mouse. Our recent results indicate that the CD38—cADPR signal system functions in insulin secretion in man. We identified a missense mutation in the CD38 gene [88, 89] in Japanese diabetic patients [90]. The resulting CD38 protein showed altered catalytic activities, with a decreased production of cADPR. Furthermore, circulating anti-CD38 autoantibodies have been detected in 10-14% of Japanese [91] as well as Caucasian diabetic patients [92-94]. The autoantibody altered the enzymic activity of islet CD38 and insulin secretion in vitro. These findings strongly suggest that the CD38—cADPR signal system functions in insulin secretion by glucose in man.

The CD38—cADPR signal system for insulin secretion is different from the conventional hypothesis [69] in which Ca\textsuperscript{2+} influx from extracellular sources plays a role in insulin secretion by glucose. Furthermore, the CD38—cADPR signal system is also different from the hypothesis proposed by Berridge and Irvine of Cambridge University [95], in which IP\textsubscript{3} induces Ca\textsuperscript{2+} release from the intracellular pool, the endoplasmic reticulum. In this context, the CD38—cADPR signal system was the focus of intense debate [96-100]. Controversial results were reported using diabetic β-cells such as ob/ob mouse islets and RINm5F cells, which have been traditionally used for studying insulin secretion in Europe and U.S.A. We revealed that the Ca\textsuperscript{2+} release responses of these diabetic β-cell microsomes were quite different from those of normal islet microsomes [71]. Microsomes from normal C57BL mouse islets released Ca\textsuperscript{2+} in response to cADPR but scarce-
ly in response to IP$_3$. This response to cADPR was completely attenuated by the prior addition of 8-amino-cADPR, an antagonist of cADPR [101]. In contrast to normal islet microsomes, ob/ob mouse islet microsomes released only a small amount of Ca$^{2+}$ by cADPR but released much Ca$^{2+}$ by IP$_3$. RINm5F cell microsomes responded well to IP$_3$ to release Ca$^{2+}$ but did not respond to cADPR. RINm5F cells are rat insulinoma-derived immortal cells and show almost no glucose-induced insulin secreting ability. Furthermore, the CD38 mRNA level was significantly decreased in ob/ob islets [71], and in RINm5F cells, CD38 mRNA was not expressed [73]. These results indicate that the CD38-cADPR signal system for insulin secretion is used under normal physiological conditions, and is replaced by the IP$_3$ system in diabetic β-cells such as ob/ob mouse islets and RINm5F cells. In fact, Balb/c mouse islets showed distinct increases in intracellular cADPR, Ca$^{2+}$, and insulin secretion by glucose [102], and MIN6 cells showed a dramatic Ca$^{2+}$ mobilization in response to cADPR via the ryanodine receptor despite the fact that no response to IP$_3$ was observed [80].

Although IP$_3$ has been thought to be a second messenger for Ca$^{2+}$ mobilization from intracellular stores, as described above, cADPR induces Ca$^{2+}$ release from pancreatic islet microsomes but IP$_3$ does not. In cerebellum microsomes, both cADPR and IP$_3$ induced Ca$^{2+}$ release. Therefore, cells can utilize two second messengers, IP$_3$ and cADPR, for Ca$^{2+}$ mobilization, depending on the species of cells as well as
differences in cellular conditions, physiological or pathological, performing a variety of cellular functions. Recently, various physiological phenomena from animal to plant cells become understandable in terms of this novel signal system [103-127]. In pancreatic acinar cells of CD38-knockout mice, the acetylcholine-induced Ca\(^{2+}\) oscillation was greatly reduced or completely disappeared under a physiological concentration of acetylcholine [126]. Furthermore, acetylcholine induced the cADPR formation in normal acinar cells, but not in CD38 knockout acinar cells. The IP\(_3\) formation was very small in the presence of a physiological concentration of acetylcholine and there was no difference between normal and CD38 knockout cells. Probably, acetylcholine induces the cADPR formation via the G-protein-coupled CD38 system [128]. In pancreatic β-cells, glucose is metabolized and induces the CD38-cADPR system to secrete insulin. In many other cells, hormones and neurotransmitters may regulate the CD38-cADPR signal system in a receptor-coupled manner, such as in a G-protein-coupled manner, to express various physiological responses.

CONCLUSION AND FUTURE PROSPECTS

In the first part of this paper, we described that PARP activation causes NAD\(^+\) depletion to form poly(ADP-ribose), resulting in necrotic β-cell death. More recently, accumulating evidence has confirmed that the mechanism proposed for β-cell death is involved in the process of the cell death of many tissues and cells. In the second part, we described β-cell regeneration and Reg gene, showing that PARP acts as a transcription factor for Reg gene, and that the active transcriptional complex for Reg gene is not formed when PARP is activated and auto-poly(ADP-ribosyl)ated. Recently, Reg proteins have been shown to be a growth factor for Schwann cells, neuronal cells, and gastrointestinal cells. In the last part, we described that the cADPR formation from NAD\(^+\) is essential for insulin secretion by glucose in β-cells. Recently, various physiological phenomena from animal to plant cells become understandable in terms of a novel signal system, the CD38-cADPR signal system. Therefore, the inhibition of the PARP activity by PARP inhibitors results in at least three important events in the cell: PARP inhibitors prevent the necrotic cell death, keep PARP active as a transcription factor for cell regeneration, and maintain the formation of a second messenger, cADPR, to achieve the cell function (Figure 6).

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