Resolving the topological enigma in \(Ca^{2+}\)-signaling by cyclic ADP-ribose and NAADP

Hon Cheung LEE*, Yong Juan ZHAO

State Key Laboratory of Chemical Oncogenomics, Key Laboratory of Chemical Genomics, Peking University Shenzhen Graduate School, Shenzhen, China, 518055

*Corresponding authors: leehoncheung@gmail.com

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Abstract

Cyclic ADP-ribose (cADPR) and nicotinic acid adenine dinucleotide phosphate (NAADP) are two structurally distinct messengers that mobilize the endoplasmic and endo-lysosomal \(Ca^{2+}\)-stores, respectively. Both are synthesized by CD38 molecule (CD38), which has long been thought to be a type II membrane protein whose catalytic domain, intriguingly, faces to the outside of the cell. Accordingly, for more than 20 years it has remained unresolved how CD38 can use cytosolic substrates such as NAD and NADP to produce messengers that target intracellular \(Ca^{2+}\)-stores. The discovery of type III CD38 whose catalytic domain faces the cytosol has now begun to clarify this topological conundrum. This article reviews the ideas and clues leading to the discovery of the type III CD38; highlights an innovative approach for uncovering its natural existence, and discusses the regulators of its activity, folding and degradation. We also review the compartmentalization of cADPR and NAADP biogenesis. We further discuss the possible mechanisms that promote type III CD38 expression and appraise a proposal of a \(Ca^{2+}\)-signaling mechanism based on substrate limitation and product translocation. The surprising finding of another enzyme that produces cADPR and NAADP, sterile alpha and TIR motif containing 1 (SARM1), is described. SARM1 regulates axonal degeneration and has no sequence similarity with CD38, but can catalyze the same set of multi-reactions and has the same cytosolic orientation as the type III CD38. The intriguing finding that SARM1 is activated by nicotinamide mononucleotide to produce cADPR and NAADP suggests that it may function as a regulated \(Ca^{2+}\)-signaling enzyme like CD38.
**Introduction**

Mobilization intracellular Ca\(^{2+}\)-stores is a universal signaling mechanism cells employ for responding to a wide range of external stimuli. Ligands, such as hormones, bind to surface receptors and activate production of second messengers in the cytosol, which in turn activate the Ca\(^{2+}\)-release mechanisms of the stores and result in cytosolic Ca\(^{2+}\) changes. The first such messengers identified was inositol trisphosphate (IP3), which targets the cytosolic portion of its receptor channel in the endoplasmic reticulum (ER) (1,2). Cells, however, possess multiple Ca\(^{2+}\)-stores. Soon after, two other Ca\(^{2+}\)-messengers, cyclic ADP-ribose (cADPR) (3,4) and nicotinic acid adenine dinucleotide phosphate (NAADP) (5,6) were discovered, with cADPR acting on the ryanodine channel (7,8) present also in the ER, while NAADP targets the Ca\(^{2+}\)-releasing mechanisms in the endo-lysosomes (9,10). Like IP3, both nucleotide messengers are known to regulate a wide range of physiological functions and in a variety of cell types spanning three biological kingdoms (reviewed in (11-16)), including mediating the adrenergic receptor activation in the salivary gland (17) and the hippocampal mGluR1-dependent long-term potentiation (18), shown recently.

The synthesis enzymes for both nucleotide messengers were soon identified, even though both messengers were unknown molecules that had never been described before. The first enzyme shown to be able to cyclize NAD to produce cADPR was named ADP-ribosyl cyclase (cyclase) (19,20). It is a small soluble protein of about 30 kDa found only in *Aplysia* (mollusks). Soon afterward, sequence comparison revealed that the carboxyl (C)-domain of human CD38 was homologous (19). CD38 is a single pass membrane protein consists of a short amino tail, a transmembrane segment and a C-domain (21).

Surprisingly, despite their distinct differences in structure and function, both messengers are produced by the same enzymes, CD38 and the cyclase (22,23). Both are multi-functional enzymes that not only can cyclize NAD to produce cADPR but also can catalyze a base-exchange reaction to produce of NAADP from NADP and nicotinic acid (24-27). Crystallography in combination with mutagenesis, identified the catalytic site in the C-domain where NAD binds (Fig. 1) and delineated the mechanism of both messenger production (22,23,28). CD38 has since remained the only fully characterized enzyme for producing cADPR and NAADP. Although CD157, a glycosylphosphatidylinositol-anchored homolog of CD38, also can produce cADPR from NAD, its activity is much lower than CD38 (29,30). Both its catalysis and function remain under-characterized.

Ablation of CD38 gene in mice produces multiple physiological defects (reviewed in (11-14)), including the altering response of skeletal muscle contractile force to beta-adrenergic stimulation (31) and impaired development of astrocytes and oligodendrocytes (32), confirming its biological relevance. Gene ablation also provides strong evidence that CD38 is responsible for producing cADPR and NAADP. For example, in lymphokine-activated killer (LAK) cells, interleukin-8 induced NAADP formation after cADPR production, resulting in Ca\(^{2+}\) changes and activation of cell migration in these cells. Formation of both nucleotide messengers was abrogated if the CD38 gene was deleted, indicating that CD38 is responsible their synthesis (33).

CD38 has long been thought to be a type II membrane protein expressing mostly on
cell surface and with its C-domain facing outside (Fig. 1) (21). In fact, it was first identified as an antigen on the surface of lymphocytes, which expression varies with the developmental stages (34), but strangely, with no known biological function. Since the C-domain is also the catalytic domain of CD38, this raises a topological puzzle; how a type II protein with its catalytic domain facing outside can use cytosolic substrates, NAD and NADP, to produce messengers that target the cytosolic portions of the Ca^{2+} release machineries.

Cells are organized and separated into two main compartments, cytosol and extra-cytosol, by lipid bilayer that also defines the topology of all transmembrane proteins. The lumen of organelles, such as ER and endo-lysosomes, is topologically contiguous with the outside and constitutes part of the extra-cytosolic compartment. In Ca^{2+}-signaling, surface receptors with their ligand-binding portions facing outside serve to transduce external signals, such as interleukin-8 described above, into the production of messenger molecules, such as NAADP and cADPR, in the cytosolic compartment.

The topological enigma raised by CD38 being a type II protein has defied resolution for more than twenty years. This has recently begun to be clarified by studies focusing on the membrane orientation of CD38, which have led to the proposal of the existence of type III CD38 with an opposite membrane orientation. With its catalytic domain facing the cytosol, type III CD38 is fully consistent with the regular topology of a Ca^{2+} signaling enzyme.

**Type III CD38**

*Clues for the type III CD38.* Based on sequence analyses, the possible existence of type III CD38 with its catalytic domain facing the cytosol was first proposed (11,35). If correct, it should readily resolve the topological issue. An accepted rule that governs the orientation of single pass transmembrane proteins is the “positive inside rule” (36), which states that the more positively charged side of the two sides of the transmembrane segment is the one more likely to be in the cytosol. In the case of CD38, the N-terminal side has, overall, about three net positive charges (red R and K, Fig. 1) and two on the carboxyl side of the transmembrane segment, consistent with CD38 being preferentially expressed as a type II protein. The difference in the charges is minimal though, less than one charge, suggesting a good possibility of CD38 being expressed also in the opposite orientation, as a type III protein.

Proteins expressing in two opposite membrane orientations are not common but have many documented cases nonetheless (15), including prion (37), epoxide hydrolase (38) and ductin (39). Likewise, bacterial EmrE (40) and melanocortin-2 receptor accessory protein (41), both are expressed as a homo-dimer with the two monomers in opposite orientation. In the case of cytochrome P450 2E1, it is localized to the ER but has opposite transmembrane orientation when transported to the plasma membrane (42).

**Cytosolic CD38 is enzymatically active.** Whether type III CD38 exists or not is physiologically relevant only if it is enzymatically active. The issue is far from straightforward because of the two critical characteristics of CD38 that must be considered, namely glycosylation and disulfides (Fig. 1). Glycosylation is immaterial for activity since recombinant CD38 with the glycosylation sites mutated and produced in yeast is fully active (43,44). On the other hand, disulfides are known to
be important for the enzymatic activities of CD38. It is not a certainty that the critical disulfides of the C-domain can be formed when it is facing the cytosol in the type III orientation.

The issue of disulfide was directly tested by expressing the catalytic domain in the cytosol. An inducible construct was made by splicing the catalytic domain of CD38 to EGFP, which directed the expression of the construct in the cytosol (35). Induction by tetracycline resulted in progressive elevation of cellular cADPR levels, indicating that the catalytic C-domain expressed in the cytosol is active and can cyclize cytosolic NAD to cADPR. A combination of mutational and immuno- analyses showed that the disulfides of the catalytic domain are formed and remain intact in the cytosol (35). Cytosolic proteins with disulfides are not common but an analysis of disulfide bonds in the Protein Data Bank shows that 298 cytosolic or nuclear proteins contain 509 structurally-defined disulfide bonds (45). Indeed, many cytosolic proteins such as chaperones, glycolytic enzymes, and kinases are found to contain disulfides (46).

Construction of type III CD38. The next step in verifying the idea of type III CD38 is to actually construct it. If the charges indeed govern the orientation of CD38, mutating the positive residues in its N-terminal tail should result in expression of the CD38 in the type III orientation. This is the case (47,48). Mutation of even one of the positive residues to aspartate resulted in expression of significant amount of type III CD38. With all mutated, more than ninety five percent of CD38 were expressed as type III (Fig 1). It was expressed in the ER instead of the plasma membrane where the type II CD38 was localized. The membrane orientation of type III CD38 was verified by independent assays (47). As expected, type III CD38 was shown to be non-glycosylated, as the C-domain was facing the cytosol instead of the lumen of the ER where glycosylation normally occurs (47).

Most importantly, the constructed type III CD38 was enzymatically active with specific activity similar to the wildtype (47,48), indicating the expressed type III CD38 was folded correctly even with its catalytic domain facing the cytosol. In live cells, cellular cADPR contents progressively increased concomitant with the induced expression of the constructed type III CD38 (47).

Endogenous Type III CD38. The construction of the active type III CD38 represents a definite proof of principle for the proposal of the type III CD38. Whether it does exist naturally needs to be established. That the endogenous type III CD38 has so far defied detection is because most of the CD38 is expressed preferentially as type II and that it is identical to type III CD38 except in membrane orientation. The assay must thus be topologically specific and highly sensitive. The first and simpler assay is to detect the N-terminal tail of the type III CD38 exposed to the outside of live cells using specific antibodies. The integrity of the plasma membrane serves to ensure the antibodies can only target the tails exposed to outside. This approach assumes that the type III CD38 is expressed not only in the ER but also in the plasma membrane (Fig. 1 and 2).

Anti-tail immunofluorescence. Highly specific monoclonal and polyclonal antibodies (Ab, Fig. 2) have been raised against the N-terminal tail of CD38, which sequence is quite unique (47). Immunofluorescence staining of live human HL-60 cells following retinoic acid induction showed increased signals of the tail on the surface, suggesting an increased expression
of the endogenous type III CD38 concomitant with the granulocytic differentiation of the cells. Similar results were seen in U937 cells, a human monocytic cell line, after activation by interferon gamma. A population of type III CD38 positive cells in human peripheral blood mononuclear cells was also detected by immunostaining with anti-tail (47).

Likewise, endogenous type III CD38 was detected in mouse spleen cells and the LAK cells using mouse anti-tail (49). It was found to be non-glycosylated, consistent with the results obtained from the constructed human type III CD38 described above (47,48). Together, these results show that type III CD38 is naturally present in both human and mouse cells, and that its expression is physiologically regulated.

The DepID assay. The main limitation of the anti-tail approach is that it can only detect type III CD38 expressed in the plasma membrane. As described above, most of the constructed type III CD38 is expressed in the ER instead (47,48). A novel method called DepID (Dual epitopes protein IDentification) has been devised to detect the type III CD38 from inside live cells (50). It is based on the use of nanobodies (Nbs), single domain antibodies that can be functionally expressed inside cells. Nbs have been widely used to modulate the functions of intracellular proteins (51-53) and, when tagged with a fluorescent protein (chromobody), to visualize their dynamic changes in live cells (54,55).

A series of Nbs targeting three separate epitopes on the catalytic domain of CD38 has been generated (56). For DepID, two of them, Nb1053 and Nb551 (Fig. 3), were each fused with a segment of a luciferase, LucN and LucC (Fig. 2). Separately, the segments are inactive. But when the constructs bind to the two epitopes on the same type III CD38, the segments are brought together to reconstitute the active luciferase and produce luminescence. The assay has low background, since the unbound probes are inactive, and high sensitivity, because of the activity of the reconstituted luciferase. The specificity of DepID is also much higher than standard immunoassay as it requires the target protein to have both epitopes on the same molecule.

The DepID assay was verified using HEK293 cells expressing the constructed type III CD38 described above as control (50). To detect endogenous type III CD38, the DepID assay was applied to two human multiple myeloma cell lines, LP-1 and OPM2. Strong luminescence signal was seen in LP-1 cells but not when CD38 was ablated by using the CRISPR/Cas9 technique. Even stronger signals were seen in OPM2 cells, but only when both DepID probes were expressed in them (50). Thus, two totally distinct assays, anti-tail and DepID, both document the natural existence of type III CD38 in the plasma membrane and internally in human and mouse cells.

Regulation of type III CD38. That the catalytic domain of the endogenous type III CD38 is facing the cytosol raises the possibility that it is interacting with a variety of regulators present in the cytosol, allowing elucidation of its regulation. In turn, the identification of these cytosolic factors can further substantiate the membrane topology of the natural type III CD38.

CIB1 regulates cellular cADPR levels. Using the yeast two hybrid technique with the catalytic domain of CD38 as bait, a number of interacting regulators have been identified, including a cytosolic Ca\(^{2+}\)-binding protein, CIB1 (50). It has four Ca\(^{2+}\) binding sites and two of them are of high affinity with the EF-hand motif (57) (Fig. 1). CIB1 is a multi-functional regulator involved in a
wide range of cellular activities, including modulation of the IP3-receptor Ca\(^{2+}\)-release channel (58).

That CIB1 indeed binds to the type III CD38 was verified in vitro using recombinant proteins, in cell lysates using immuno-precipitation and in live cells using the bimolecular fluorescence complementation technique (BiFC) (50). Mutational analyses further identified the N-terminus of CIB1 being involved in interacting with the catalytic domain of type III CD38 (50). The functional consequence of the interaction was assessed using shRNA to knock down and Cas9/guide RNA to knock out CIB1. A direct correlation between the cellular cADPR and CIB1 levels was demonstrated. These results indicate that the type III CD38 is functionally active in producing cellular cADPR and that the activity is specifically modulated through interaction with cytosolic CIB1 (50).

Chaperones regulate folding. As described above, the catalytic domain of the type III CD38 is highly unusual in that, when expressed in the cytosol, can readily form its six disulfides needed for its enzymatic activities (35,47) (Fig. 1). It is possible that the information for the disulfide formation is encoded in its primary sequence, such that during its translation and folding, the microenvironment and proximity of appropriate cysteines facilitate the disulfide formation. Indeed, it is documented that the local microenvironment of protein can greatly influence the ionization and reactivity of the thiolate groups of cysteines at physiological pH (59). Specific binding of cytosolic chaperones can further help the disulfide formation. This may well be the general process that allows not only type III CD38 but also the hundreds of cytosolic proteins to form and retain disulfide bonds (45,46).

Consistently, it was shown using the yeast two hybrid technique that ST13, a co-chaperone also called Hip, is another cytosolic protein associated with the catalytic domain of CD38 (60). In addition, a complex of chaperones, including also Hsp90, Hsc70, DnaJA1 and DnaJA2, was identified by mass spectrometry in the immuno-precipitates of the C-domain of CD38 (60). Knockdown of ST13 or Hsp90 resulted in reduction of the type III CD38, suggesting that both chaperones are involved in the correct folding of type III CD38, including its disulfide formation. Their knockdown may thus increase mis-folding and degradation (Fig. 4A).

Chaperones regulate degradation. In contrast, knockdown of Hsc70, DnaJA1 or DnaJA2 caused large increase in type III CD38 levels by as much as two to three folds. Likewise, an inhibitor of Hsp70 and Hsc70, VER-155008, dramatically increased the levels of type III CD38 and concomitantly raised the cellular cADPR levels (60), further documenting that type III CD38 is responsible for cellular cADPR. The results suggest that these chaperones are facilitating the degradation of type III CD38 and their knockdown reduces degradation, allowing the proteins to accumulate. This was verified by chase experiments using cycloheximide to block translation and subsequent degradation of type III CD38 was expectedly slowed by the knock down of Hsc70 (60).

It is known that Hsc70 facilitates protein degradation via chaperone-mediated autophagy (CMA) (61). Consistently, blocking the lysosomal degradation with bafilomycin increased the levels of type III CD38, while MG-132, an inhibitor of the proteasome degradation pathway for soluble proteins, had no effect. The import of the type III CD38 into lysosomes for degradation was via the lysosomal receptor,
Lamp2A (61), and its knockdown increased the level of type III CD38 by 2-3 folds (60) (Fig 4A). Immuno-staining showed significant co-localization of type III CD38 and Lamp2A, which could be further increased by geldanamycin, an inducer of CMA degradation of the ryanodine receptor (62). The binding region on the type III CD38 for Lamp2A was identified in the last 19 residues from the carboxyl terminus, which was also the region where Hsc70 and DnaJA2 bound to the type III CD38 (Fig. 4A) (60).

CMA of soluble proteins has been most commonly studied. But type III CD38 is not the only membrane protein degraded through CMA (63). Others include the ryanodine receptor (62), a calcium channel in the ER targeted by cADPR, and the EGF-receptor (64) are also known to be degraded via CMA as well. Exactly how membrane proteins are transported to the lysosomes has not been elucidated. It can involve p97/Cdc48-mediated protein dislocation or intramembrane proteolysis extracting transmembrane segments and transporting to lysosomes, as has been proposed (65).

Nox4 regulates disulfide formation. Type III CD38 in mouse has been characterized and is shown to be a non-glycosylated protein of about 36 kDa (49), similar to the constructed human type III CD38 (47,48). Its disulfide within the last 19 carboxyl residues is not formed, in contrast to type II CD38. A monoclonal antibody, M19, raised against this part of the molecule can thus specifically recognize the type III CD38 (49).

As described in the introduction section, mouse CD38 is shown to mediate the interleukin-8 signaling in LAK cells (33,49). A recent study further identifies type III CD38 being responsible for mediating the Ca\textsuperscript{2+}-changes induced by reactive oxygen species in these cells (49). Inhibiting NADPH oxidase 4 (Nox4) in the cells using either specific inhibitor, shRNA or gene knockout, inhibited the interleukin-8 activated cADPR production and Ca\textsuperscript{2+} changes (49). Nox4 makes reactive oxygen species/superoxide, which can oxidize cysteines, leading to disulfide bond formation. In the cell lysates, Nox4 was shown to activate the cADPR synthesizing activity, which also required the presence of the phosphorylated p22phox, an accessory component of Nox4 (49). H\textsubscript{2}O\textsubscript{2}, product of Nox4, could directly stimulate the activity. The results suggest that Nox4 and the phosphorylated p22phox bind to and activate type III CD38 to produce cADPR through oxidization of its cysteines. That disulfides are important for cADPR-synthesizing activity of type III CD38 is consistent with that seen in human type III CD38 (47). Immuno-precipitation of type III CD38 using M19 shows that Nox4 and p22phox were indeed associated with the type III CD38. Mutating individual cysteines of CD38 identified cysteine164 as the Nox4 target, which was oxidized to form a disulfide with cysteine177 (Fig. 3B) (49). Enzymes such as Nox4 and/or disulfide isomerases (66), in association with chaperones may well be responsible for the disulfide formation of type III CD38 during its translation and folding. Taken together, these results further establish that the type III CD38 is a regulated Ca\textsuperscript{2+}-signaling enzyme.

**Potential mechanisms for regulating the expression of type III CD38.**

The existence of type III CD38 with its catalytic domain facing the cytosol readily resolves the topological enigma of Ca\textsuperscript{2+}-signaling mediated by cADPR. It is thus of clear interest to survey and explore what known mechanisms may be consistent with and may involve in regulating the
expression of type III CD38 in particular and single pass membrane proteins in general. This is not a trivial issue since, theoretically, the existence and co-expression of an alternate membrane orientation for any membrane protein can potentially endow it with entirely different functions.

That type III CD38 can be constructed by altering the charges in the N-terminal tail of CD38 indicates that the “positive inside” rule governs its expression. The molecular basis of the rule is depicted in Figure 4 (left panel). Protein synthesis generally begins in the cytosolic ribosomes. Binding of the signal recognition particle (SRP, Fig. 4) to the nascent polypeptide then directs the continuation of the translation process of membrane protein to the ER. The electrostatic repulsion of the positive charge of the nascent polypeptide with the translocation complex leads to its folding and exiting through the side channel of the complex, and inserting into the membrane in the type II orientation (67).

Phosphorylation of the tail. In the case of CD38, the charge difference between the two sides spanning the transmembrane segment is small and the charge repulsion of its two sides with the translocation complex is more similar. The folding of the nascent polypeptide may thus become stochastic, but still with a preference for the type II orientation. Consistent with this notion, cells transfected with the wildtype CD38 expressed a significant portion in the type III orientation as detected by DepID (50), even though the major portion was type II.

Increasing the negative charges at the N-terminal side of the nascent polypeptide should increase the charge attraction with the translocation complex and reduce its folding inside the complex (middle panel, Fig. 4), which can result in a graded increase in the portion of the expressed proteins being in the type III orientation. This is the case. With the four positive residues changed to negative, essentially all the expressed CD38 were found to be in the type III orientation (47,48). Similar results were seen in mouse CD38 (49), attesting to species generality of the rule.

A possible and natural mean to increase the negative charge in the tail is phosphorylation. There are three serines in the N-terminal tail of CD38. Changing them to aspartates to simulate phosphorylation indeed increased the expression of type III CD38 (48). Naturally, co-translational phosphorylation of the nascent polypeptide has been observed (68-70) (middle panel, Fig. 4). In particular, mTORC2 kinase has been shown to be associated with a large ribosomal subunit protein at the tunnel exit and can phosphorylate the nascent polypeptide on both serine (69) or threonine (70). Whether a similar phosphorylation of the serines in N-terminal tail during its translation can lead to increase in the expression of type III CD38 is a worthwhile investigation.

Lipid modulation. Another factor that can affect the orientation of the membrane proteins is the lipid content of the membrane. In bacteria, lactose permease exhibited a mixture of topological conformations from one orientation to a complete inversion of the N-terminal helical bundle depending on the phosphatidylethanolamine content. The topological orientations were even interconvertible by post-assembly synthesis or dilution of the lipid in vivo (71). Using fluorescence energy transfer, the rate of the protein flipping in the lipid bilayer in both directions triggered by a change in the proteoliposome lipid composition occurred rapidly in the range of seconds (72). The possibility that lipid binding to type II CD38 might reduce the energy barrier sufficiently
to allow orientation flipping is an intriguing possibility.

**SRP-independent ER insertion.** Although the SRP-dependent insertion of protein into the ER membrane (Left and middle panels, Fig. 4) is generally thought to be the dominant process, the SRP-independent insertion has also been described (73,74). In yeast, nascent polypeptides with signal sequences not recognized efficiently by SRP are bound by soluble chaperone Ssa1, targeting the nascent chains to the Sec 63 complex (right panel, Fig. 4), which in turn interacts and transfers the nascent chains to the translocation complex. The movement and folding of the nascent chains through the channel of the complex is facilitated by Kar2, a luminal chaperone (74).

In mammalian cells, it is believed that Hsp70s serves similar function as Ssa1. Hsc70, a member of Hsp70 chaperone family, is associated with the type III CD38 (60). Its sequence, however, only have minimal homology with Ssa1. It remains a possibility that binding of an appropriate chaperone to the hydrophobic segment of CD38 may prevent aggregation and direct the nascent polypeptide to the translocation complex (right panel, Fig. 4), or allow the entire protein to be translated in the cytosol. Subsequent insertion into the ER membrane in the type III orientation can be mediated by this SRP-independent pathway.

**Type II CD38**

The discovery of type III CD38, with its cytosolic orientation and its efficient production of intracellular cADPR, solves one long-standing CD38 mystery. But another mystery remains: Though CD38 is capable of making both cADPR and NAADP, the optimal pHs for the two reactions are substantially different, meaning that type III CD38 cannot make NAADP in the cytosol. So how then does NAADP get made? Solving this mystery requires returning to earlier proposals of CD38 trafficking.

Type II CD38 on cell surface is the dominant form of CD38 found in many cells, especially in blood cells, and is enzymatically active. Its catalytic domain facing the extra-cytosolic compartment is topologically inappropriate to contribute to production of cellular cADPR. However, the topological barrier of the bilayer membrane, in principle, can be specifically and selectively relaxed by membrane transporters. Among them, connexin 43 is permeant to NAD (75), while nucleoside transporters can mediate both concentrative and equilibrative movement of cADPR (76). It has thus been proposed that the surface type II CD38 is internalized through endocytosis together with surface transporters into the endo-lysosomal compartment (16,77-79). The cytosolic NAD can enter via connexin 43 and get converted to cADPR inside the compartment, which is then transported out by the nucleoside transporter back to the cytosol to target the ER Ca^{2+}-stores.

**Biogenesis of NAADP.** The endo-lysosomal compartment is highly acidic and not conducive for the cyclization reaction of CD38 to produce cADPR, even if NAD is made available by the transporters. It is, however, well suited for the base-exchange reaction to produce NAADP by CD38, which has an optimum at pH 4 (22,80). This is because of the acidic pH neutralizes the charge repulsion between the negative Glu146 at the catalytic site with the base-exchange substrate, nicotinic acid (80). Ever since the first documentation of this peculiar acidic requirement of the NAADP synthesis (22), the endo-lysosomes have been suggested to be a special cellular compartment for the biogenesis of NAADP.
Topology of CD38/cADPR/NAADP-signaling

The idea has been made more substantive with the findings that the endo-lysosomes are functional Ca\textsuperscript{2+}-stores (81,82) and are targeted by NAADP, which specifically activates the endo-lysosomal two pore channels (9,10) to effect Ca\textsuperscript{2+} release from the stores in response to various physiological stimuli. It is a novel notion that the two forms of CD38 are topologically segregated into two compartments so as to produce two structurally and functionally distinct Ca\textsuperscript{2+} messengers.

The first test for this proposal is to identify a stimulus for endocytosis and determine if the surface type II CD38 can indeed be internalized into the endo-lysosomes via endocytosis. Nanobody (Nb) against the catalytic domain of CD38 is such a stimulus. Phenomenologically, it is very effective in inducing the endocytosis of the endogenous type II CD38 expressed on the surface of human myeloma LP-1 cells, as well as that expressed in Hela and HEK293 cells transfected with CD38 (83). The Nb can be fluorescently labeled and imaging showed that the endocytosed complex reached the endo-lysosomes. Concomitant with the Nb-activated endocytosis, the cellular NAADP level in LP-1 cells progressively increased in the presence of exogenous nicotinic acid (NA) (83). These results show that endocytosis can indeed deliver surface type II CD38 to the endo-lysosomes and that the internalized CD38 remains enzymatically active for at least a period of time, even though it was eventually degraded (84). This was further substantiated by constructing a lysosomal CD38 specifically directed to express in the lysosomes by splicing the catalytic domain of CD38 to the C-terminal segment of LAMP1 for lysosomal retention. The construct, likewise, efficiently elevated the cellular NAADP levels in the presence of NA (83).

The requirement for NA, however, indicates that the access of substrate to the extra-cytosolic compartment of the endo-lysosomes is still limited. This tight compartmentation suggests the possibility of a novel mechanism for signaling. Type II CD38 is constitutively active and the regulation is not on CD38 itself but on the access of substrates through activation of specific membrane transporters. For example, the permeability of connexin 43 was shown to be regulated by phosphorylation by protein kinase C (85). Whether connexin 43, or other transporters, is mediating the movement of NADP and NA, substrates for the base-exchange reaction of CD38, remains to be investigated.

**Biogenesis of cADPR.** Evidence described above suggests that the type II CD38 may contribute to the production of NAADP, but intriguing results indicate that production of cellular cADPR is likely to be tightly compartmentalized in the cytosol and that type II CD38 may not contribute substantially to the cellular cADPR synthesis. This was first seen in cells transfected with wildtype human CD38 or with charge mutants with increasing number of the N-terminal positive residues changed. Normalizing the cellular cADPR contents to the amounts of CD38 expressed, it was seen that cells expressing CD38 with all four N-terminal positive charges mutated, and thus in the type III orientation, had the highest normalized cADPR contents, higher than the cells expressing wildtype CD38 (47,48). As most of the wildtype was expressed as type II on the surface and only a small amounts as type III inside the cells (50), the result is the first indication that type II CD38 may not contribute much to the production of cytosolic cADPR.

Consistently, in mouse LAK cells, the type II CD38 is constitutively active, but the
activity cannot be stimulated by interleukin-8. Type III CD38, in contrast, has low basal activity but can be stimulated by the cytokine (49). The elevation of the cellular cADPR levels activated by the cytokine is thus mediated by the type III and not the type II CD38.

To directly address whether the surface type II CD38 contributes to the production of cellular cADPR, a crosslinking technique was developed to identify possible regulators of surface CD38. The results show that the tranferrin receptor, CD71, specifically interacts with surface CD38 and that its knockdown reduces type II CD38 expression (84). To simulate this regulatory effect of CD71, a construct was made by splicing the amino tail of CD71 to a nanobody against CD38. As described above, binding of the nanobody to surface CD38 induced endocytosis. Its anchoring to the surface by the amino tail of CD71 greatly increased its effectiveness and resulted in more than eighty percent reduction of type II CD38 via endocytosis, which was then degraded in the lysosomes (84). Remarkably, the cellular cADPR levels did not decrease correspondingly. Bafilomycin-mediated blockade of lysosomal degradation greatly elevated active type II CD38 by trapping it in the lysosomes, but also did not increase cADPR levels. Retention of type II CD38 in the ER by expressing an ER construct that prevented its transport to the cell surface, likewise did not change cADPR levels (84). These results provide the first and direct evidence that cADPR biogenesis occurs in the cytosol and is catalyzed mainly by type III CD38, and that type II CD38, compartmentalized in the ER, lysosomes, or cell surface, contributes only minimally to cADPR biogenesis.

A new enzyme emerges

As described in the introduction section, CD38 has been the only fully characterized enzyme for synthesizing cADPR and NAADP. It has, however, been observed that ablation of the CD38 gene in mice resulted in large decreases of cADPR contents in many tissues except in the brain (86), suggesting the possibility of another enzyme for synthesizing cADPR in the brain. However, searches in the mouse genome data base have produced no promising candidates.

It is thus surprising that Sterile alpha and Toll/interleukin-1 receptor motif-containing 1 (SARM1), a protein unrelated and has no sequence similarity with CD38, has recently been shown to be able to catalyze the synthesis of cADPR (87,88). This unexpected discovery is a convergence of two different and independent lines of investigation.

SARM1 is known to play an important role in axonal degeneration underlying several neurological disorders. Its activation triggered axon degeneration locally and induced NAD depletion (89), suggesting that it may have NAD hydrolyzing activity. Consistently, the Toll/interleukin-1 receptor (TIR) domain of SARM1, recombinantly produced and purified, was shown to be an effective NADase (87,88). Surprisingly, analysis of its hydrolysis products by HPLC showed, in addition to ADP-ribose, a small peak corresponding to cADPR (87,88).

That SARM1 is indeed an endogenous and regulated enzyme capable of producing cADPR in live cells was demonstrated through an entirely separate effort, which was aimed at developing specific inhibitors of CD38. One of the most potent found was sulfo-araF-NMN (CZ48), an catalysis based inhibitor and a cell-permeant mimetic of nicotinamide mononucleotide (NMN) (90). It blocked CD38 by forming a covalent bond with its catalytic residue, Glu226, as

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revealed by crystallography (90). It is thus expected that CZ48 should inhibit CD38 and cause reduction in cellular cADPR.

Most surprisingly, when CZ48 was applied to HEK293 cells, it effectively raised the cellular level of cADPR instead (88). CRISPR/Cas9 deletion of the CD38 gene did not prevent the induced cADPR increase, indicating the production was not mediated by CD38, the hitherto only known enzyme that can increase cellular cADPR. Deletion of SARM1, however, eliminated the cADPR increase while overexpressing SARM1 enhanced it, indicating the target of CZ48 is SARM1 (88).

As CZ48 is a mimetic of NMN, it suggests that SARM1 may be stimulated by NMN as well. This is the case. Both NMN and CZ48 were equally effective in activating SARM1 to produce cADPR from NAD with essentially identical half-maximal effective concentrations (88). Consistently, knockout of NMN adenylyltransferase 1, the main NMN utilizing enzyme in cells, elevated cellular NMN and activated cADPR production, indicating that NMN is an endogenous activator of SARM1.

In contrast to CZ48, NMN cannot activate cADPR increases when applied to live cells. Structure-function studies indicate that the sulfur attached to the phosphate group of CZ48 is important for its cell permeability. A wide range of human and mouse cells are responsive to CZ48, including primary mouse dorsal root ganglions (88). The extents of cADPR increases in response to CZ48 correlated well with the cellular SARM1 levels. CZ48 is clearly a cell permeant, versatile, useful and specific activator of cellular cADPR production via SARM1.

Unlike CD38, SARM1 possesses multiple domains, including a N-terminal domain with multiple armadillo repeat motifs (ARMs), two tandem sterile alpha motif (SAM) domains, and a catalytic C-terminal TIR domain. The amino terminal segment is believed to be associated with the mitochondria, while the catalytic TIR domain is facing the cytosol (91,92). The ARM domain is auto-regulatory since its deletion removed the regulatory response of SARM1 to CZ48 and NMN. The remaining SAM-TIR portion became constitutive active in producing cADPR both in vitro and in vivo (88). The results suggest that CZ48 activates SARM1 by inducing a conformational change (Fig. 5) to release the catalytic TIR-domain from the sequestration of the ARM-domain. In addition, CZ48 also induced dimerization of the TIR-domain as part of the process of activating the enzymatic activity of SARM1 (Fig. 5) (88).

SARM1 and CD38 are clearly two totally different proteins with no sequence and structural similarity. It is thus surprising that they are virtually identical catalytically (88). Both enzymes catalyze NAD hydrolysis and cyclization of NAD to cADPR, as well as a base-exchange reaction using NADP and nicotinic acid as substrates to produce NAADP. Similar catalysis of the two enzymes would suggest similarity in their active sites. This seems not to be the case, as CZ48 formed covalent bond with the catalytic residue of CD38 and irreversibly inhibited its enzymatic activity, while it activated SARM1 reversibly without forming a covalent linkage (88). Another notable difference was that SARM1 had much less cADPR hydrolyzing activity than CD38, making SARM1 more efficient for cADPR synthesis. It is thus catalytically more similar to Aplysia ADP-ribosyl cyclase
Studies on SARM1 have hitherto mostly focused on its role in axonal degeneration. This is mainly because it is highly expressed in neuronal cells. In non-neuronal HEK-293 cells overexpressing SARM1, CZ48 did induce non-apoptotic cell death with elevation of cADPR and depletion NAD, similar to that observed during axonal degeneration (88). However, SARM1 is a ubiquitous enzyme endogenously present in a wide range of non-neuronal cells as well. In these cells, it was also fully efficient in elevating cADPR in response to CZ48 activation, but without inducing cell death (88). It is thus certain that SARM1 is not just a mediator of cell death. What physiological roles SARM1 plays in these cells are undoubtedly of wide interests. As cADPR is documented as a second messenger for mediating a variety of physiological functions, SARM1 may well be a regulable enzyme involved in some or all of these signaling processes.

Summary
Evidence amassed in the past thirty years firmly establishes the wide range of signaling functions of cADPR and NAADP in cells spanning three biological kingdoms. Much is now known about their Ca^{2+}-signaling mechanisms and the stores the two nucleotide messengers target. Despite the fact that both messengers were unknown molecules that had never been described before, their synthesis enzymes, the cyclase and CD38, were soon identified and their catalysis elucidated. Contrasting to these advances is the conspicuous weakness in the understanding of the regulation of CD38. The conceptual bottleneck is centered around the topological issues described in this review. It is puzzling how a type II enzyme with its catalytic domain facing outside can be regulated by the cell.

The discovery of the type III CD38 expressed both in the ER and the plasma membrane as summarized in Figure 5 has now begun to clarify the issue. With the catalytic domain facing the cytosol, type III CD38 can readily use the cytosolic NAD as substrate and produce cADPR (Fig. 5), which targets the ryanodine receptor in the ER. Regulators of type III CD38 have since been identified, including CIB1 chaperones and Nos4. The type II CD38 is expressed on the cell surface and can be internalized via endocytosis, together with specific transporters. The acidic lumen of the endo-lysosomes is conducive for the type II CD38 to catalyze the base-exchange reaction to produce NAADP, which is then transported out via the transporters to target the two-pore channel.

The groundbreaking finding that SARM1 can catalyze the production of cADPR and NAADP same as CD38 should usher in a new frontier. It is an auto-regulated enzyme. NMN, or its mimetic CZ48, activates dimerization of SARM1, releasing the catalytic TIR-domain from the inhibition of the ARM-domain (Fig. 5). SARM1 is known to regulate axonal degeneration. That it may involve in regulating other functions and function as a Ca^{2+}-signaling enzyme are intriguing and should warrant further investigation.

Another advance derived from the study of type III CD38 that may well have widespread application is the novel approach of the DepID assay specifically designed to detect the topology of cellular proteins. Although it is developed for CD38, it is, in fact, quite versatile and can readily applied to any intracellular protein. A variation in the format of ELISA has also been used as a diagnostic test to precisely quantify the levels of soluble CD38 in the plasma of...
multiple myeloma patients. Myeloma cells are known to overexpress surface CD38 and show increased shedding as soluble CD38. Using the DepID as an ultra-sensitive and specific assay, it was shown that the plasma levels of soluble CD38 were significantly higher than those from healthy donors and were correlated with the progress of the disease (93). DepID is thus a topologically selective, highly sensitive and specific approach for \textit{in vivo} and \textit{in vitro} monitoring of proteins.

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Figure 1. **Type II and III CD38.** The structure of the catalytic C-domain is based on crystallography (PDB, 1YH3) and is shown with NAD at the active site. Its six disulfides are indicated and colored cyan. The N-terminal side of the transmembrane segment (orange helix) of type II CD38 has 3 positive arginines (red R), one more than the C-terminal side. Changing these positive residues to aspartates (green D) convert type II to type III with the catalytic domain facing the cytosol. CIB1 (PDB, 1XO5) has four Ca²⁺-binding sites and it interacts with the cytosolic C-domain of the type III CD38.
Figure 2. Immunoassay and Dual epitopes protein IDentification (DepID) for detecting type III CD38 inside cells. Immunoassay uses externally applied antibody against the N-terminal tail to detect the tail of the type III CD38 exposed to the outside. DepID probes are constructed with nanobodies (Nb, blue and cyan) against two different epitopes on the catalytic domain of type III CD38 and each is fused with a luciferase fragment (LucN and LucC, semicircle). Binding of the Nbs to the epitopes on the type III CD38 (PDB, 3RAJ) brings the luciferase fragment together, reconstituting the luciferase and produces luminescence (light green).
Figure 3. Regulators of the type III CD38. A. CIB1 interacts with type III CD38 and modulates its cADPR producing activity. The correct folding and disulfide formation of type III CD38 are assisted by chaperones Hsp90 and ST13. On the other hand, chaperones Hsc70, DnaJA1 and DnaJA2 mediate degradation of type III CD38 via the lysosomal receptor Lamp2A. B. NADPH oxidase 4 (Nox4) and its associated component p22phox activates mouse type III CD38 (PDB, 2EG9) by oxidizing cysteine164 to form disulfide with cysteine177.
Figure 4. Potential mechanisms for expression of type III CD38. **Left panel.** Translation begins in the cytosolic ribosomes. Binding of the signal recognition particles (SRP) to the signal sequence of the nascent polypeptide directs it to the translocation complex in the ER. The positive charges of the nascent interact with the complex and result in folding of the polypeptide. Subsequent exit from the side channel of the complex results in type II insertion into the ER membrane (left). **Middle panel.** Kinases are known to associate with the ribosomes and the phosphorylation of the nascent polypeptide can reduce its charge interaction with the translocation complex and thus the folding of the polypeptide, resulting in the type III orientation. The lipid content of the ER can also modulate the membrane orientation of CD38, as is observed in bacteria. **Right panel.** Chaperones such as Hsp70 or Ssa1 can bind to the hydrophobic segment of the nascent polypeptide and, through the Sec63 complex, direct its insertion into the ER membrane in the type III orientation via the translocation complex.
Figure 5. The CD38/cADPR/NAADP-signaling pathway. Type III CD38, present in the ER and plasma membrane, cyclizes cytosolic NAD to produce cellular cADPR (PDB, 2O3Q), which targets the ryanodine receptor in the ER and releases Ca\(^{2+}\) from the store. The activity of the type III CD38 is modulated by CIB1, a Ca\(^{2+}\)-binding regulator. Type II CD38 is expressed on the cell surface and can be internalized through endocytosis to the endo-lysosomes, which acidic lumen is conducive for NAADP production. Transporters internalized together with the type II CD38 facilitate the movement of NAADP (PDB, 4F45) into the cytosol, where it targets Ca\(^{2+}\)-channels, such as the two-pores channel, in the acidic Ca\(^{2+}\)-stores. CZ48 (PDB, 3ROM) is a mimetic of NMN and is cell permeant. It induces dimerization and conformational changes in SARM1, releasing the catalytic TIR domain from the auto-inhibition of the ARM domain. The amino segment of SARM1 is associated with the mitochondria, while the TIR is facing the cytosol. The activated SARM1 cyclizes NAD to produce cADPR in a manner similar to CD38.
Resolving the topological enigma in Ca2+-signaling by cyclic ADP-ribose and NAADP
Hon Cheung Lee and Yong Juan Zhao

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