Several members of tumor necrosis factor receptor (TNFR) superfamily that these members activate caspase-8 from death-inducing signaling complex (DISC) in TNF ligand-receptor signal transduction have been identified. In the extrinsic pathway, apoptotic signal transduction is induced in death domain (DD) superfamily; it consists of a hexahelical bundle that contains 80 amino acids. The DD superfamily includes about 100 members that belong to four subfamilies: death domain (DD), caspase recruitment domain (CARD), pyrin domain (PYD), and death effector domain (DED). This superfamily contains key building blocks; with these blocks, multimeric complexes are formed through homotypic interactions. Furthermore, each DD-binding event occurs exclusively. The DD superfamily regulates the balance between death and survival of cells. In this study, the structures, functions, and unique features of DD superfamily members are compared with their complexes. By elucidating structural insights of DD superfamily members, we investigate the interaction mechanisms of DD domains; these domains are involved in TNF ligand-receptor signaling. These DD superfamily members play a pivotal role in the development of more specific treatments of cancer. [BMB Reports 2016; 49(3): 159-166]
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Fig. 1. Cell signaling pathway through which DD complex elicits a balance between survival and programmed cell death (9).

...receptors and TNF receptors. They constitute apoptotic signaling platforms of extrinsic pathway (5, 10, 11). Apoptotic signal transduction is induced through a homology domain containing a hexahelical bundle of 80 amino acids. With this process, DD superfamily members are produced (12-14). Furthermore, DDs construct key building blocks that are involved in the formation of multimeric complexes; these complexes are associated with death signaling cascades. In this study, we summarize recent findings that elucidate three dimensional structures of TNF ligand-receptor superfamily. They provide molecular and functional characterization of homotypic DD interaction motifs, which are associated with programmed cell death.

THE DEATH-FOLD INTERACTIONS

When the signal of Fas receptor is activated, the Fas-associated death domain protein (FADD) develops homotypic interaction motifs, such as DD and death effector domain (DED). With these motifs, caspase-8 can be recruited to the docking site. In this process, Fas and caspase-8 interact simultaneously via DD and DED (9). The aspartate-specific cysteine proteases (Caspases) are primary executioners of non-inflammatory cell death. Effector caspases cleave regulatory enzymes, such as poly (ADP-ribose) polymerase (PARP). They also cleave activating endonucleases, such as caspase-activated deoxyribonuclease (CAD) (15). Biologically, caspases are broadly categorized into initiator and effector caspases. The initiator caspases have death-fold motifs, such as DED or caspase recruitment domain (CARD). They trigger non-inflammatory cell death by activating effector caspases. The activation of effectors is conducted by initiator caspases, which act at cytoplasmic sites, nucleus, and other organelles. Many death-fold interactions are formed by specific adaptor molecules, such as CARDS, DEDs, DDs, and pyrin domains (PYDs): they belong to the superfamily of DD (16).

An apoptotic protease cascade develops when cytochrome c influences dATP-dependent interaction of apoptotic protease activating factor (Apaf-1) and caspase-9. Consequently, the apoptosome is generated via a CARD-CARD homotypic interaction (Fig. 2A-C) (17). When cytochrome c and Apaf-1 are bound to each other, the signaling complex is activated. Consequently, there is autoactivation and recruitment of procaspase 9 (18). The active apoptosome complex contains several Apaf-1, procaspase-9 proteins, and general complex forms; the molecular weight of these components is approximately 700 kDa (19).

THE DD SUPERFAMILY

Many proteins have significantly different amino acid sequences. However, functional and structural characteristics indicate that a common ancestor is probably involved in superfamilies (20). The DD superfamily, which is characterized by the presence of conserved homotypic interaction motifs, has emerged as the principal mediator in the transduction of a cell signal (21). About 100 members of DD, DED, CARD, and PYD subfamilies have been identified till date (12-14). Despite the appreciable divergence in the overall sequence of each member, the structure of each DD is hallmarkd. The DD superfamily constitutes six amphipathic α-helices that are folded in an antiparallel α-helical bundle. The atomic structures of DD superfamily are determined by either nuclear magnetic reso-
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Fig. 2. (A) Ribbon representations of Apaf-1 (blue) and caspase-9 (green). CARDs complex is shown (PDB ID: 3YGS). (B) Carbon stick representation of CARDs complex is shown. Each important amino acid in CARD interaction is shown in yellow. (C) The Apaf-1 and caspase-9 complex, which undergoes a 180° rotation along the horizontal axis, is shown as a surface representation. (D) Amino acid sequence alignment of DED superfamily members. The secondary structural elements of DED superfamily members, as indicated. Alpha helices are presented as red ellipses, while loop regions are presented as gray lines. The negatively charged amino acids are conserved through species, while highly conserved motifs are presented in red. (E-H). Three dimensional structures of different DD subfamilies (23). (E) The DD of FADD (PDB: 1E3Y). (F) The CARD of Apaf-1 (PDB: 1CY5). (G) The pyrin domain of apoptosis repressor with CARD protein (PDB: 1UCP). (H) The second DED of v-FLIP (PDB: 2BBR). The DD superfamily structures are presented as flat ribbon. Each specific region is illustrated (yellow).

nance (NMR) or X-ray crystallography (Fig. 2E-H) (22-26).
Recently, many studies have been conducted on DD superfamily proteins. In these studies, it is proved that DD superfamily proteins contain many different motifs; the features of these different motifs have significant differences. The differences exist even within the members of subfamilies. These differences are caused by the alterations in the size and arrangement of α-helices. Furthermore, these differences are also caused by the varying distribution of charged and hydrophobic residues along the surface (23-26). As a result, all the superfamily members are included in the homotypic interaction proteins of every kind. Furthermore, they do not cross the boundaries of subgroup (27). Compared to respective structural features, it is likely that DDs have an exposed third α-helix. This α-helix is more pliable than other subfamilies (Fig. 2E-H) (22-26). The CARD is another DD subfamily consisting of members with a highly conserved structure. There is striking similarity between members of CARD and the amino-terminal domains of DD-containing proteins: CRADD and Apaf-1 (24, 28). The most significant difference between CARD and other subfamilies is as follows: the first α-helix (α1) leads to the disruption of hydrogen bonds between nitrogen atoms of amide and oxygen atoms in the carbonyl group of CARD. The helix in CARD is composed of two short helices in α-helix: the first part is α1a and the second part is α1b. Both the parts are connected by a linker region. Owing to bending, residues present at the C-terminal of helix α1 in Apaf-1 are closer to the hydrophobic core; they are far away from the corresponding residues in RAIDD CARD (24, 28). This helical shift is propagated in the remaining portion of the molecule. Consequently, the positions and boundaries of all other helices and surface loops are altered (24, 28-30). The PYD subfamily members have a hydrophobically stabilized loop region between helices α2 and α3. The connecting loop of PYD is longer than that of other subfamilies, whereas the exposed helix α3 in PYD is shortest in the DD superfamily (25, 31). The DED subfamily has exposed hydrophobic sites, which are smaller than DDs in the helix region. Moreover, a conserved motif is present between

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helices α5 and α6 (26, 32, 33). This specific motif contains amino acids, such as aspartic acid and arginine, in helix α6. They mediate protein folding and ensure the functional conformation of DEDs. Negatively charged amino acids (glutamic acid, aspartic acid, or asparagine) are conserved at the N-terminal. Furthermore, there is an Arg-x-Asp-Leu (RxDL; ‘x’ for any amino acid) motif in nearly all DEDs (33). Most single and tandem DED proteins have RxDL motifs that are consistent in the highly conserved region; however, this motif is not present in other DD subfamilies (Fig. 2D) (26, 33, 34). When the three-dimensional structure of MC159 (v-FLIP) was elucidated, a hydrogen-bonded triad was revealed on the surface of DED1 and DED2. This triad contributes to the highly charged features on one face of the structure (32). Previous studies prove that RxDL motif mediates molluscum contagiosum virus (MCV) protein MC159 and block apoptosis (33, 35). Conversely, the charged amino acids in caspase-8 and caspase-10 are either greatly altered or missing in DEDs. The DED2 domain in caspase-8 has three charged amino acids: Glu, Lys, and Ser (36). By converting arginine to lysine, the hydrogen bonding potential decreases and there is interaction with only one negatively charged amino acid. In caspase-10, RxDL motif is missing in both DED1 and DED2. Therefore, we deduce that this motif does not play an important role in the recruitment of caspase. However, it is important in other functions of proteins containing DED (32).

THE DEATH INDUCIBLE SIGNALING COMPLEX

NMR spectroscopy is used to develop three-dimensional models of FADD, DED, and DD (32, 37). Members of DD superfamily include Fas and FADD; they share helix α6 strands with Greek key topology (32, 27). It has been shown that this topology governs the process of protein folding. This process is initiated by the formation of a hydrophobic pocket, which contains almost all the conserved sites of DD superfamily (38). The DD domains regulate the assembly of death inducing, multimeric signaling complexes exhibiting the activity of critical effectors (9). However, very limited information is available about the structural features of DDs complexes till date. By performing X-ray crystallography, models of cell death have been identified; these models induced Fas/FADD-DISC (39). The complex activates procaspase-8, which forms an integral part of programmed cell death pathway. It is triggered by binding between Fas-receptor and FasL. When TNFR-1 and TNF-α (40, 41), FasL is expressed in activated T lymphocytes or natural killer (NK) cells. They cause the destruction of target cells, such as virus-infected or damaged cells (42). The Fas/FADD-DISC crystal structure exists in the closed form owing to lack of a stimulus. This is followed by multimerization of its factors; they form an active platform (39). In FasL-treated cells, Fas and FADD proteins form signaling complexes via modular interaction in their C-terminal DDs (39, 42). Furthermore, FADD recruits caspase-8 through DED, promoting their enzymatic activity and stimulating the expression of apoptosis-related proteins (38, 39, 43).

In Fas receptor signaling, the central event involves the formation of DISC; it is comprised of procaspase-8, procaspase-10, cellular FLICE-like inhibitory proteins (c-FLIP), and adapter molecule FADD (44). The procaspase-mediated apoptosis is crucial to Fas/FADD-DISC signaling study. All procaspase-8 isoforms have two N-terminal DEDs that are similar to the isoforms of c-FLIP protein. In this study, we identified several DED isoforms of procaspase and c-FLIP at DISC. There were two procaspase-8 isoforms: (procaspase-8a [p55] and procaspase-8b [p53]). In addition, there were three c-FLIP isoforms, such as c-FLIP long (c-FLIPL), c-FLIP short (c-FLIPS), and c-FLIP Raji (c-FLIPR) (Fig. 3) (26, 44-47). Recently, Schleicht et al. demonstrated that FADD, procaspase, and c-FLIP were the DED members forming a signaling complex in their DED domains (48). The new paradigm of DISC proved that procaspase-8/10 is activated via the assembly of DED chains assembly (48, 49). These chains enable the homodimerization of procaspase-8, which plays an important role in the activation of procaspase-8 in DISC. For example, owing to the mutations of some key binding amino acids in procaspase-8, DED2 interrupts the formation of DED chain in cells. Moreover, these specific binding sites are conserved in nearly all DED domains; these domains interact with another DED via hydrophobic interaction motif of phenylalanine/leucine (FL motif) (Figs. 2D and H) (22-26, 49, 50). After the formation of DED chain, procaspase-8a/b homodimer processing involves two sequential cleavage steps at the DISC. The first cleavage step takes place at the aspartic acid 374, which lies between p18 and p10 caspase subunits. The second cleavage step occurs in two aspartic acids: Asp216 and Asp384. Consequently, large protease subunit p18, small protease subunit p10, and the domain p26/p24 are produced. Owing to DED chain processing, the active caspase-8 heterotetramer, p18-p10, is subsequently released into cytosol and it triggers an apoptotic signal cascade (51, 52). The procaspase-8-dependent apoptotic death is interrupted by c-FLIP; however, it is also activated when DED interacts at DISC. The shorter c-FLIP isoforms (c-FLIPL and c-FLIPS) consist of only tandem DEDs that block cell death and procaspase-8 cleavage. However, the long splice variant of c-FLIP isomorf (c-FLIPII) either accelerates or blocks cell death and procaspase-8 cleavage; the interaction of c-FLIP isomorf depends on its density (53, 54). The DED chain model helps us in elucidating the molecular mechanisms of DD superfamily.

SIGNAL TRANSDUCTION OF LIGANDS AND RECEPTORS ASSOCIATED WITH TNF SUPERFAMILY

Two different forms of TNF-α and lymphotoxin-α were identified in activated macrophages and T cells (55, 56). Several members of TNF superfamily and their receptors are identified in many tissues and organs of animals (57). Each factor of TNF...
superfamily interacts with at least one receptor of TNFR superfamily. Furthermore, some TNF factors bind with several receptors (5, 8, 21). TNF-α is a cytokine that exhibits inflammatory activity; a member of a group of cytokines is included in the TNF-ligand superfamily. The TNF-ligand superfamily plays an important role in regulating cellular life and death (57, 58).

TNF-α plays an important role in the regulation of immune cells; however, it can also induce apoptotic cell death and inhibit tumorigenesis, metastasis, bone resorption, and viral replication (59-61). In addition, TNF-α plays an important role in the activation of two distinct cell surface receptors: TNFR-1 and TNFR-2 (6, 59, 60). The expression of TNFR-2 is manifested in the immune system and endothelial cells, whereas the expression of TNFR-1 is detected in many tissues. In TNFR-1, the extracellular position has four characteristic cysteine-rich domains (CRDs) for the purpose of interaction; these CRDs bind directly with the TNF-α trimer (6, 57-59). Previous studies show that owing to the interaction between TNF-α and TNFR-1, activation of proteins is mediated in the following components: tumor necrosis factor receptor type 1-associated death domain (TRADD), receptor-interacting serine/threonine-protein kinase 1 (RIPK1), and TNF-R-associated factor 2 (TRAF2). These events play a crucial role in the cascade of TNF signal transduction pathway (59-61).

**DD SUPERFAMILY-MEDIATED CELL SIGNAL CASCADE**

In most cell lines, nearly all death receptors mediate apoptosis; moreover, TNFR has several signal transduction pathways, controlling a broad spectrum of life and death in cells (Fig. 1) (9, 57, 58). With an activated TNFR-1 signal cascade, the lysine 63 polyubiquitination of RIPK1 is promoted (62, 63). Protein kinase RIPK1 functions as a kinase depending on the extent to which it undergoes ubiquitination. It either regulates prosurvival transcription factor NF-κB or promotes apoptotic and necrotic cell death under the effect of TRADD, FADD, caspase-8, and RIPK3 (62-65). The assembly of RIPK1, TNFR-1, and TRADD complex (complex I) was found to be inhibited by cellular inhibitor of apoptosis protein 1 (cIAP1) and cellular inhibitor of apoptosis protein 2 (cIAP2) in cytosol (62, 66, 67). Furthermore, with the interaction of these two inhibitor proteins, the polyubiquitin chains of RIPK1 were formed. They act as a crucial binding site in TAK1/TAB2/3 complex and NEMO/IKK complex. NF-κB is translocated in the nucleus by these two complexes. Thereafter, there is expression of cylindromatosis (CYLD), tumor necrosis factor alpha-induced protein 3 (TNFAIP3), and other NF-κB related factors (64, 66, 68).

Recent studies prove the intracellular signaling kinase RIPK1 acts as a key switch in the regulation of cell fate. Depending on the cellular context, RIPK1 determines whether the proinflammatory cytokine TNF induces activation, apoptosis, or programmed necrosis of NF-κB (69, 70). RIPK1 is involved in the regulation of cell death when there is internalization of death receptor. RIPK1 is dissociated from TNFR-1 of TNF signal transduction complex, while lysine 63 deubiquitination of RIPK1 is mediated by the following deubiquitination enzymes: CYLD and A20. These enzymes inhibit the expression of prosurvival transcription factor, NF-κB (5, 62, 63). Following deubiquitination, RIPK1 interacts with TRADD, FADD, RIPK3,
generating TRADD-dependent complex (complex IIA) and RIPK1-dependent complex (complex IIB) (63, 70-72). Complex IIA necessitates a TRADD-FADD scaffold to recruit caspase-8, which has DED and DD sites. Thus, an apoptotic pathway is initiated (5, 21, 71).

When TRADD does not exist, RIPK1 is formed and apoptosis signal pathway is initiated. With the formation of complex IIB, there is FADD-mediated recruitment and activation of caspase-8 at the cleavage of RIPK1 and RIPK3. Owing to the interaction between these two RIPK superfamily proteins, there is increase in metabolic activities. In this process, the activities of glutamate dehydrogenase 1, glutamate ammonia ligase, and glycogen phosphorylase may an important regulatory role. This results in programmed necrotic cell death (63, 65, 69-71).

When X-ray crystallographic and NMR studies were performed on the members of DD superfamily, a great deal of information was obtained. These studies prove that its members play a pivotal role in the assembly and regulation of complexes, which are crucial for cell survival and programmed cell death signaling. They ensure the recruitment of kinase proteins through an interaction with homotypic motifs. Despite conservation in topology, members of DD superfamily have three dimensional structures with significant difference. Among DD superfamily proteins, the major structural difference is the presence of diverse twisting angles between each α-helix bundle. These angles are formed by several helices in DD, and they exhibit the biggest difference in three dimensional structure. Owing to the structural differences in DD superfamily, none of the superfamily members undergo cross interactions among subgroups. Owing to the interaction between TNF-ligand and TNF-receptor superfamily, there is mediation in many DD proteins. These proteins create new signal transduction complexes, such as apoptosome, DISC, and other multimers of DD. The apoptosis that is induced by TNF receptor is a biologically relevant form of cell death; it occurs in the presence of sufficient DISC activity. In the extrinsic pathway, Fas/FADD oligomerization regulates apoptotic signal cascade. Depending on the homotypic interaction with DD superfamily, the transition occurs between programmed cell death and survival (71). Previous studies prove that the expression of DD superfamily increases in cancer cells and tissues, which are treated with anticancer agents. This indicates that DD superfamily participates in the action of these agents (73, 74). Moreover, many extrinsic signaling pathways stimulate tumor cells to induce programmed cell death; these events are independent of tumor-suppressor p53 (75). Thus, DD receptors might be a useful target in cancer research studies. In this study, we summarized the structural characterization and function of homotypic interaction motifs and their complexes. The findings of this study play an important role in the development of more specific treatments that are used for curing inflammation and cancer.

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