DAP12 signaling: from immune cells to bone modeling and brain myelination

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DAP12 (also called KARAP) is a transmembrane adapter well known for its role in transducing activation signals for an extended array of receptors in NK cells, granulocytes, monocytes/macrophages, and DCs (1, 2). In this issue of the JCI, Takai and colleagues unveil that DAP12-deficient mice exhibit osteopetrosis and demyelination of the CNS (3). Human studies have concurrently shown that genetic defects of DAP12 result in a rare syndrome characterized by bone cysts and presenile dementia (4, 5). Thus, DAP12 appears to be critically involved in bone modeling and brain myelination. If so, how can DAP12 regulate both immune and non-immune functions?

DAP12-expressing cells in the immune system, bones, and brain

DAP12 was originally identified as a transmembrane adapter molecule that pairs with activating NK receptors, such as KIR2DS and NKG2D, and myeloid cell receptors, including SIRP-1β, MDL-1, TREM-1, -2, and -3 (1, 2). The cytoplasmic domain of DAP12 contains a tyrosine-based motif, which functions as a docking site for p72syk and ZAP70 tyrosine kinases. These promote recruitment and activation of PI3K, the phospholipase Cγ1, and the p44/p42 extracellular signal-regulated kinase (ERK) pathways (6). In agreement with the role of DAP12 in NK and myeloid cell activation, the initial characterization on DAP12-deficient mice revealed lack of function of activating NK cell receptors (7, 8), accumulation of DCs in peripheral tissues (7) and impairment of Th1 responses (8), possibly due to a reduced ability of DCs to migrate to lymph nodes and prime T cell responses.

This view of DAP12 as an immune signaling mediator was subsequently challenged by the observation that rare genetic mutations of human DAP12 cause no obvious immune defects but result in a syndrome characterized by bone cysts and presenile dementia called Nasu-Hakola disease (NHD), or polycystic lipomembranous osteodysplasia with sclerosing leuкоencephalopathy (PLOSL) (4). Since DAP12 is expressed in cells of myeloid origin, it was suggested that DAP12 may regulate the function of osteoclasts and microglial cells, which share a myeloid origin and are critical for bone modeling and brain function, respectively. Prompted by this hypothesis, Takai and colleagues generated DAP12-deficient mice and found that they exhibit functional defects of osteoclasts and oligodendrocytes that were overlooked in previous studies (3).

DAP12 as a critical regulator of osteoclasts and oligodendrocytes

The study by Takai et al shows that DAP12 protein is expressed in normal osteoclasts (3). These are polykaryons of the monocyte/macrophage lineage that specialize in bone resorption (9). Normal osteoclasts derive from the fusion of mononuclear myeloid precursors in the presence of two cytokines: MCSF and a tumor necrosis factor-related protein defined as the receptor activator of the NF-κB ligand (RANKL) (also known as TRANCE, OPGL, and ODF) (10). However, DAP12-deficient bone marrow precursors do not differentiate in vitro into mature osteoclasts with bone resorptive function, and DAP12-deficient mice develop osteopetrosis in vivo (3).

Even though the clinical manifestations of DAP12-deficient mice resemble those of NHD patients, there are some significant differences. While DAP12-deficient mice exhibit osteopetrosis, NHD patients develop bone cysts filled with lipids, mainly in bones of the extremities. In DAP12-deficient mice demyelination is concentrated in the medial thalamus. In contrast, NHD patients exhibit leukodystrophy, accumulation of lipidic material, sclerosing leuкоencephalopathy, axonal loss, and massive gliosis predominant.
2/DAP12 may favor osteoclast and oligodendrocyte differentiation, whereas inflammatory conditions may promote differentiation of macrophages, DCs and microglial cells through engagement of other DAP12-associated receptors (Figure 1). The TREM-2/DAP12 signaling pathway may also trigger changes in actin polymerization and cytoskeleton organization, which are required for the fusion of osteoclast precursors and to ensure that the cellular processes of oligodendrocytes contact and wrap around the neuronal axons. The discovery of the ligands of TREM-2 and other DAP12-associated receptors will be critical to verify these hypotheses and to precisely characterize the multiple contributions of DAP12 to immune and non-immune functions.


Figure 1
Hypothetical role of DAP12 in myeloid cell and oligodendrocyte development. In homeostatic conditions, the TREM-2 receptor may trigger the DAP12 signaling pathway in immature oligodendrocytes and osteoclasts, promoting their differentiation into mature cells capable of myelination and bone resorption, respectively. During inflammatory conditions, DAP12 may be solicited by different receptors, leading to differentiation and/or activation of microglial cells, macrophages, and DCs. While oligodendrocytes are thought to derive from a neuroepithelial progenitor, it is possible that DAP12-expressing oligodendrocytes may be of myeloid origin.

The functions of DAP12-associated receptors and their ligands
As DAP12 couples with a variety of cell surface receptors, it is important to identify the specific receptor(s) involved in the pathogenesis of the osteoclast- and oligodendrocyte-derived defects. Recent analysis of NHD patients with an intact DAP12 gene and normal expression levels of DAP12 revealed loss of function mutations in TREM-2 (13). In addition, TREM-2 transcripts have been detected in human osteoclasts (13), human brain (13), and murine microglial cells (14). Thus, TREM-2 may be the DAP12-associated receptor that predominantly regulates bone modeling and CNS myelination. Future studies need to test how TREM-2/DAP12 influences osteoclast and oligodendrocyte functions. Current data suggest that TREM-2/DAP12 may guide differentiation of precursors into osteoclasts, oligodendrocytes and microglial cells, possibly by modulating responsiveness to certain cytokines or reinforcing signaling pathways triggered by cytokine receptors or integrins (Figure 1). Under homeostatic conditions, engagement of TREM-