

Original article

A comprehensive manually curated reaction map of RANKL/RANK-signaling pathway

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Receptor activator of nuclear factor-kappa B ligand (RANKL) is a member of tumor necrosis factor (TNF) superfamily that plays a key role in the regulation of differentiation, activation and survival of osteoclasts and also in tumor cell migration and bone metastasis. Osteoclast activation induced by RANKL regulates hematopoietic stem cell mobilization as part of homeostasis and host defense mechanisms thereby linking regulation of hematopoiesis with bone remodeling. Binding of RANKL to its receptor, Receptor activator of nuclear factor-kappa B (RANK) activates molecules such as NF-kappa B, mitogen activated protein kinase (MAPK), nuclear factor of activated T cells (NFAT) and phosphatidylyl 3-kinase (PI3K). Although the molecular and cellular roles of these molecules have been reported previously, a systematic cataloging of the molecular events induced by RANKL/RANK interaction has not been attempted. Here, we present a comprehensive reaction map of the RANKL/RANK-signaling pathway based on an extensive manual curation of the published literature. We hope that the curated RANKL/RANK-signaling pathway model would enable new biomedical discoveries, which can provide novel insights into disease processes and development of novel therapeutic interventions.

Database URL: http://www.netpath.org/pathways?path_id=NetPath_21

Introduction

The paracrine system involved in many tissues includes numerous cytokines and growth factors necessary for the differentiation and maturation of cells of which amongst tumor necrosis factor (TNF) family assumes a vital role. The TNF superfamily consists of numerous ligand/receptor proteins involved in various cellular processes such as development, homeostasis and apoptosis. RANKL, also known as TNFSF11, osteoprotegerin ligand, TNF-related activation-induced cytokine (TRANCE) or osteoclast differentiation

factor, is a member of the TNF superfamily (1). The membrane bound form of RANKL is a type II transmembrane glycoprotein with an extracellular region, a transmembrane domain and an intracellular cytoplasmic region (2). RANKL also exist in a soluble form derived either through an alternative splicing event or by the proteolytic cleavage of the membrane anchoring domain of RANKL by a membrane-associated metalloprotease, such as TNF converting enzyme, matrix metalloproteinase 14, matrix metalloproteinase 7 or Disintegrin and metalloproteinase

domain-containing protein 19 (3–6). As a member of TNF family, RANK is expressed mainly on cells of monocyte/macrophage lineage such as pro-osteoclasts and osteoclasts, whereas the RANKL is synthesized by and expressed on cell membrane of bone marrow stromal cell and osteoblasts. Expression of RANKL has also been detected in a wide variety of tissues and specific cells including spleen, placenta, heart, stomach, thyroid gland, lung, brain, thymus, lymph nodes, osteoclasts and peripheral blood leukocytes (7–10). In bone metabolism, osteoclasts play primary role by involving in bone resorption. In brain, RANKL is expressed in the lateral septal nucleus and RANK is expressed specifically in the neurons and astrocytes of the preoptic area and medial septal nucleus in the hypothalamus. Hanada *et al.* (11) have shown a novel function of RANKL/RANK signaling in brain that plays a significant role in the regulation of body temperature and fever response in inflammation. Prolactin, progesterone and parathyroid hormone-like related protein stimulate RANKL expression on mammary gland epithelial cells whereas RANK is constitutively expressed in these cells. Knockout studies in mice have shown that RANKL and RANK are required for the development, proliferation and survival of lactating mammary gland during pregnancy. RANKL/RANK-signaling system is critical for the induction of lactation by mediating lobulo-alveolar morphogenesis. Kim *et al.*, have also shown that RANK activation by RANKL is crucial to control proliferation of mammary epithelial cells (12,13). Furthermore, RANKL is reported to be upregulated in primary malignant bone tumors including osteosarcoma, chondrosarcoma, giant cell tumor (14) and multiple myeloma (15). RANKL has also been shown to be overexpressed in >90% of metastatic tumor cells in adenocarcinoma lesions of prostate, breast, lung and thyroid origin (16) and also in breast cancer stromal cells (17).

RANKL binds to type I transmembrane protein known as RANK (TNFRSF11A) to form a functional hexamer complex containing trimeric RANKL and RANK (18–20). The RANKL/RANK interaction is essential to osteoclastogenesis as it stimulates the differentiation, maturation and survival of osteoclasts (21–25). In the immune system, RANKL expressed by activated T cells interacts with RANK expressed on dendritic cells in order to regulate survival and function of dendritic cells (10,26–28). RANKL/RANK system plays an important role in controlling the development of AIRE⁺ thymic medullary epithelial cells in the thymus (29) and in lymph node organogenesis (30). Osteoprotegerin (OPG), also known as (TNFRSF11B) or osteoclastogenesis inhibitory factor, is a decoy receptor that modulates signaling by RANKL during osteoclastogenesis. OPG binds to RANKL and neutralizes its function thereby negatively regulating osteoclast differentiation, maturation and survival (31,32). Undifferentiated pre-osteoblastic bone marrow stromal cells were known to express high levels of RANKL along

with relatively low levels of OPG. The ratio of RANKL/OPG increases during their activation and differentiation into osteoclasts. This ratio declines during differentiation of undifferentiated pre-osteoblastic cells into mature osteoblasts during which their osteoclastogenic activity is suppressed (33). Thus the RANKL/OPG ratio plays a significant role in determining bone mass and skeletal integrity. RANK, being a member of TNF receptor superfamily, lacks any catalytic activity and therefore recruits adaptor molecules such as TNFR-associated factors (TRAFs) to transduce signals.

Several pathways have been shown to be activated by RANKL/RANK-signaling system. These pathways include, NF-kappa B, mitogen activated protein kinases (MAPK's), protein kinase C (PKC), Ca²⁺/Calcineurin/nuclear factor of activated T cells (NFAT) and phosphoinositide-3-kinase (PI3K). The functional importance of NF-kappa B (34–40), JNK (41–44), SRC-PI3K-AKT (45,46) and p38MAPK (47) pathways in osteoclast differentiation and osteoclastogenesis have been extensively studied. Although the RANKL/RANK pathway has been well studied, a comprehensive map of this signaling pathway is not available in any of the publicly available signal transduction pathway resources. Our group has previously developed a signaling pathway resource called NetPath (<http://www.netpath.org>), which currently comprises of 20 human signaling pathways involved in immune- and cancer-signaling pathways (48). With the goal of creating manually curated individual pathways relevant to molecular biologists and clinicians alike, we describe here the development of a comprehensive resource of RANKL/RANK-signaling pathway. Similar approaches were previously undertaken by various groups to provide comprehensive maps of epidermal growth factor receptor (49), interleukin-1 receptor (50) and toll-like receptor (51) signaling pathways. RANKL/RANK-signaling pathway resource is a NetPath module freely available at http://www.netpath.org/pathways?path_id=NetPath_21.

Curation of RANKL pathway reactions

An extensive curation of scientific literature was carried out to catalog RANKL/RANK stimulated reactions. The reactions in NetPath collectively represent individual biochemical events such as protein–protein interactions (PPIs), enzyme–substrate reactions or protein translocation events. We have cataloged these reactions which are proved to be induced or enhanced by the RANKL–RANK interaction. The curation of RANKL pathway reactions was based on the following criteria: (i) reactions should be stimulated by RANKL/RANK system, (ii) reactions must be either induced or enhanced *in vivo*, (iii) proteins involved in reactions should be from human system, however other mammalian

proteins were annotated if they were not reported from human system (The mammalian proteins should have orthologous proteins identified in human), (iv) reactions if stimulated by multiple ligands were not considered and (v) only experiments carried out in cell lines of mammalian origin were considered. We used PathBuilder (52), a curation tool previously developed by our group, to annotate pathway reactions captured in this curation process.

PPIs

PPIs are important mediators and regulators of signaling. We have curated PPIs reported in literature for RANKL/RANK-signaling system as direct or complex interactions. Direct interactions represent interaction between two molecules whereas complex interactions represent the existence of more than two molecules as a complex. We have also curated various features pertaining to these PPIs including domains and motifs involved in the interaction, dependence of the reactions on various post-translational modifications (PTMs), subcellular location of the interaction and also the cell type in which the reactions were reported.

Enzyme–substrate reactions

PTMs are known to regulate signaling cascades by altering functional properties and recruiting proteins to subcellular compartments. We have captured several events of phosphorylation, dephosphorylation, ubiquitination and proteolytic cleavage in RANKL/RANK-signaling pathway. We could not capture any study specific to other PTMs such as sumoylation, desumoylation, deubiquitination, acetylation, deacetylation, methylation, demethylation, palmitoylation and glycosylation reactions in RANKL/RANK-signaling pathway. We cataloged these reactions as either direct or indirect catalytic events. Whenever the upstream enzymes responsible for modification were reported in the literature, PTMs are curated as direct. However, induced/indirect represents PTMs on proteins for which no report is available on the upstream enzyme responsible for the modification. We documented the enzyme–substrate reactions responsible for direct catalysis events also as PPIs. Whenever available in the literature, we have also cataloged the site and residue information of the PTMs and mapped these on to the respective protein sequences provided in RefSeq database. The molecules which are reported to be activated or inhibited by various assays such as GTPase assay or kinase activity assay but their PTM status has not been reported in RANKL signaling are cataloged into activation/inhibition reactions.

Protein translocation events

During signal transduction, proteins translocate across subcellular compartments or are secreted into the extracellular

space, a fate determined by interaction partners, various PTMs, or a specific regulatory event. We have curated translocation events in RANKL pathway along with their PTM dependence and description of the experimental system of observation. We used Gene Ontology (GO) to define subcellular localization (53).

Gene regulation

We have cataloged transcriptionally regulated genes by RANKL/RANK signaling from normal human cells. The cataloging involves the identification and curation of genes differentially regulated by RANKL signaling reported in literature as demonstrated using diverse experimental strategies including DNA microarrays, northern blot, serial analysis of gene expression and quantitative RT-PCR. The cells/cell types in which the genes are identified to be differentially regulated are also curated. In addition, we have also documented transcriptional regulators of the differentially regulated genes.

Summary of RANKL/RANK pathway curation

Our manual literature survey on RANKL/RANK-signaling pathway has cataloged 88 molecules involved in 73 PPIs, 59 enzyme–substrate reactions and 10 protein translocation events. PPIs include 58 binary interactions and 15 protein complexes. The 59 enzyme–substrate reactions comprise of 14 direct reactions which were also considered as direct PPIs and 45 RANKL induced catalytic reactions. We have also cataloged 74 genes which are differentially regulated at the mRNA level by RANKL signaling in various cell types in human system. The interaction domains or motifs for 14 proteins in this pathway have also been documented. The site and residue information for PTMs were captured for 17 proteins. We annotated transcriptional regulators for eight differentially regulated genes from our literature survey pertaining to studies on RANKL pathway. Each and every pathway reactions annotated in NetPath were linked to scientific articles. Six PPIs which were not available in HPRD were submitted to HPRD (54). All the RANK receptor activated reactions were reviewed by a pathway authority (http://www.netpath.org/pathway_authority?path_id=NetPath_21). We have captured several molecules associated with RANKL signaling but not included in the review articles on this pathway to the best of our knowledge. These molecules include FHL2, LYN, FYN, PTK2, ATP6V1E1, ETV5, CYLD, SQSTM1 and LIMD1. We hope that future review articles will consider the addition of these proteins while describing RANKL/RANK signaling.

Protein complexes in RANKL/RANK-signaling pathway

Multiprotein complexes, not individual proteins, are increasingly being recognized as the molecular basis of cellular fluxes of signals. Thus, the identification of dynamic protein complexes is of major interest to the experimental as well as computational biologists. The current effort on RANKL/RANK-signaling system has cataloged several such protein complexes. These complexes are enlisted in Table 1. There were seven protein complexes which were associated with RANK and seven complexes with TRAF6 as a constituent.

Data formats and availability

The RANKL pathway data described in this work is integrated into NetPath which is publicly available at (<http://www.netpath.org>). NetPath RANKL pathway page provides a summary of the pathway, statistics of the number of molecules and total reactions curated under different data categories such as physical interactions, enzyme catalysis, transport and gene regulation. An overview of the RANKL/RANK pathway page in NetPath with the curated information on PPIs, enzyme catalysis and PTMs, protein translocation reactions and gene regulation is provided in Figure 1. The annotated data can be downloaded in various community standard data exchange formats. These include Proteomics Standards Initiative for Molecular Interaction (PSI-MI version 2.5), Biological PATHway eXchange (BioPAX level 3) and Systems Biology Markup Language (SBML level 2.1). PSI-MI is a community standard for

molecular interaction data which facilitates data comparison, exchange and verification (55,56). BioPAX is an emerging standard community language that enables integration, exchange, visualization and analysis of biological pathway data which also incorporates information that can be stored in the PSI-MI format (57). SBML is a machine-readable format for representing models which facilitates mathematical simulation (58). An excel file is also provided to download the list of upregulated and downregulated genes.

Visualization of the RANKL/RANK-signaling pathway

RANKL pathway data which are made available using different community standards allows interoperability with other data visualization and analysis software such as Cytoscape (59), Ingenuity pathway analysis, Visualization and layout services for BioPAX pathway models (VISIBIOweb) (60) or Chisio BioPAX Editor (ChiBE) (61). Moreover, various software tools are currently under development which converts user-defined formats into these standard formats. For example, conversion between GenMAPP Pathway Markup Language (GPML) and standard pathway exchange formats such as SBML and BioPAX are under development using Paxtools (62,63). A signaling network of RANKL pathway can be created using cytoscape by importing the BioPAX file. The number of steps which leads to any reaction from the stimulated receptor currently depends on the interconnectivity of molecules and their reactions reported in a pathway. Thus, we hope that the RANKL pathway data available in various data formats will help the computational as well as basic biologist to visualize their own custom maps using different pathway visualization tools available in the community.

However, taking into account the heterogeneity and the necessity for experimental validation of independent pathway reactions, we have applied a set of stringent criteria to select pathway reactions from the curated data to be depicted in a pathway map. The selection of such reactions stimulated by RANK was based on any of the following criteria: (i) availability of two or more research articles which established a reaction, (ii) availability of two or more different cell types in which a particular reaction was demonstrated, (iii) proof for a PPIs as component of known protein complexes, (iv) Whether reactions reported using high-throughput experiments have a low-throughput experimental support and (v) whether reactions were represented in expert review articles with their biological significance. A basic network model of the RANKL pathway applying the above criteria was manually generated using PathVisio software (Figure 2). A .gpml file with

Table 1. Protein complexes curated in RANKL/RANK signaling

Protein complex	References
TNFRSF11A–TRAF6–TAB2–MAP3K7–TAB1	Mizukami <i>et al.</i> , 2002 (67)
TNFRSF11A–TRAF1–TRAF3–TRAF6	Wong <i>et al.</i> , 1999 (68)
RUNX1–TRAF6–FHL2	Bai <i>et al.</i> , 2008 (69)
TNFRSF11A–SRC–TRAF6	Wong <i>et al.</i> , 1999 (68)
TNFRSF11A–CBL–PIK3R1–PIK3R2–TRAF6	Arron <i>et al.</i> , 2001 (70)
TNFRSF11A–TAB1–MAP3K7	Mizukami <i>et al.</i> , 2002 (67)
TNFRSF11A–TAB2–TRAF6	Mizukami <i>et al.</i> , 2002 (67)
SQSTM1–TRAF6–CYLD	Jin <i>et al.</i> , 2008 (71)
LYN–TNFRSF11A–PTPN6–GAB2	Kim <i>et al.</i> , 2009 (72)
IKBKG–IKBKG–IKBKG	Darwech <i>et al.</i> , 2009 (73)

The table contains a list of proteins which are reported to form complexes when RANK (TNFRSF11A) is stimulated by RANKL (TNFSF11). Five of the protein complexes which contained RANK also had TRAF6 as a constituent molecule, indicating a central role of TRAF6 in RANKL/RANK signaling.

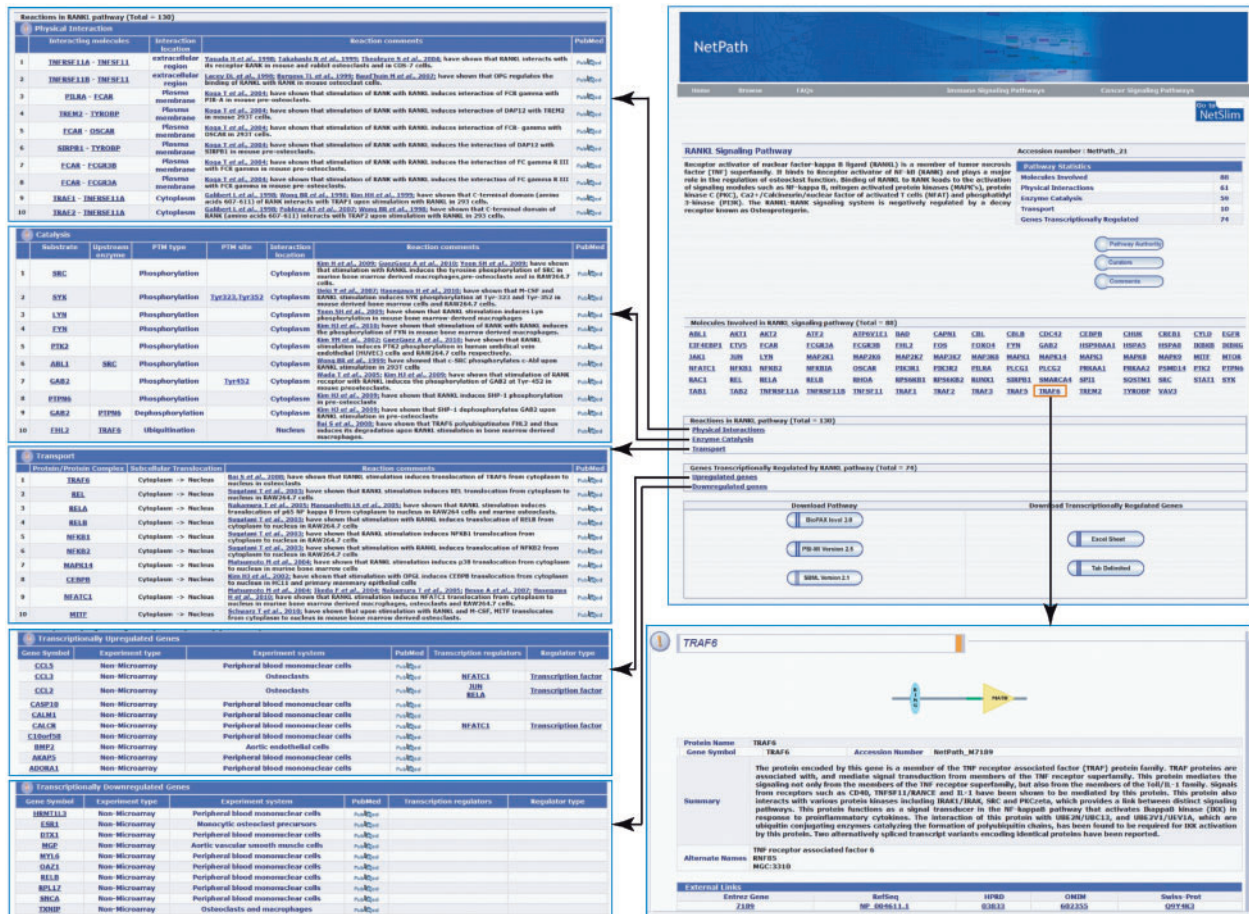


Figure 1. An overview of the RANKL pathway page in NetPath. RANKL pathway page in NetPath hosts the information on the number of molecules curated for RANKL pathway, statistics of the total number of molecules, link to RANKL pathway reactions and the list of genes which are differentially upregulated by RANKL/RANK pathway. Every molecule in the pathway page is linked to the corresponding NetPath molecule page which is further linked to Entrez gene, HPRD, OMIM and Swiss-Prot identifiers. The reaction page of the RANKL pathway contains the list of each type of reactions such as physical interactions, enzyme catalysis and transport with a brief description about the reactions with their PTM dependence or interacting regions/domains/motifs whenever it was available in literature. The list of curators and reviewers are provided in the RANKL pathway page with the details of the pathway authority. A comments tab is provided in the pathway page to invite the queries and suggestions from the community so as to update and improve RANKL pathway alike other pathways in NetPath.

citation containing description of each of the reactions, PTM status, site and residue of modification mapped to specific RefSeq accession and PubMed identifiers are also available to download at http://www.netpath.org/netpath/rankl_pathway.html. A full view of RANKL pathway image in html format is also available at the above link with each node and edge hyperlinked to their corresponding NetPath molecule page and literature evidences respectively.

Update process

With the involvement of the pathway authority, the RANKL/RANK pathway will be updated periodically as

more reactions become available. The proteins and the microRNAs which are differentially regulated by activated RANK will be curated in the future. We hope to get the active participation from the scientific community to update and improve the RANKL pathway reactions.

Conclusions

Curated and organized data pertaining to signaling pathways are crucial to understand biological processes and gene regulation. This is especially true in the wake of increasing use of high-throughput technologies to study gene and protein expression as well as PTMs (64–66). Combining such a pathway map with high-throughput

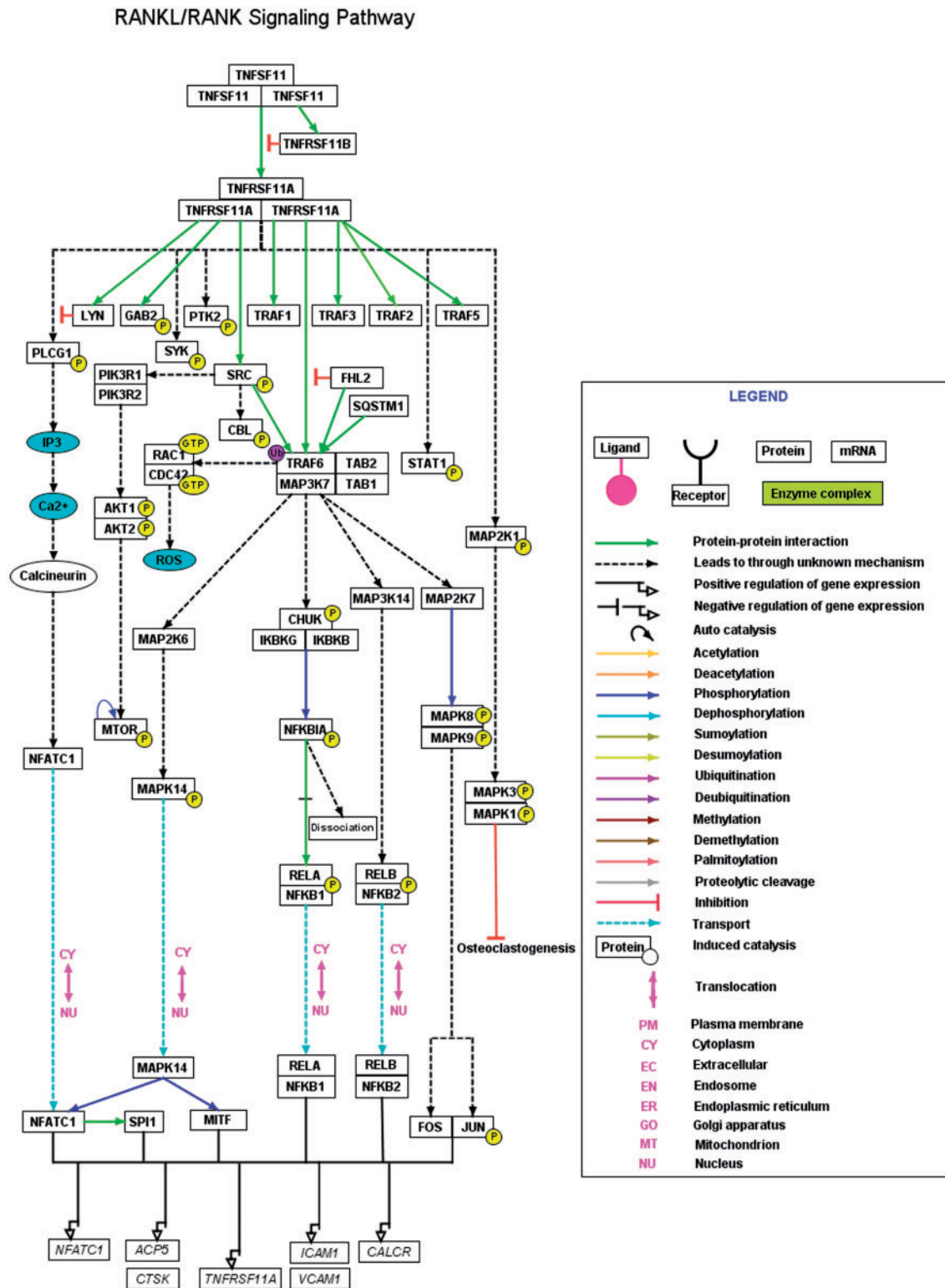


Figure 2. A high-confidence RANKL/RANK pathway reaction map using PathVisio. The RANKL/RANK pathway reaction map represents the high-confidence molecular reactions induced by the binding of RANKL homotrimer to its homotrimer receptor complex. The presented map of RANKL/RANK contains 43 proteins involved in 17 molecular associations, 24 enzyme catalysis reactions and four translocation events. The nodes and edges represent the molecules and their reactions respectively. A detailed legend representative of the information of different types of edges distinguished with various colors is provided. A downloadable version of this map is available with the description for each of the reactions and the inclusion criteria for selection of the reaction in the map at http://www.netpath.org/netslim/rankl_pathway.html.

data can be used to study perturbations using systems biology approaches. We anticipate that the RANKL/RANK signaling pathway resource described here will help accelerate research in both normal physiology as well as disease mechanisms. The RANKL/RANK pathway reactions are available to download at (http://www.netpath.org/pathways?path_id=NetPath_21). We encourage interested members of the scientific community to help us improve the quality of the information in NetPath by submitting their suggestions and critical comments through <http://www.netpath.org/comments>.

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