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(54) HETEROLOGOUS POLYPEPTIDE OF THE TNF FAMILY
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## ABSTRACT

Anewly identified heteromeric ligand of the Tumor Necrosis Factor (TNF)-family, referred to hereinafter as "APBF" has been identified.
MPASSPFLLAPKGPPQNMGGPVREPALSVALWSWGAALGAVAGAMALLT 50
QQTELQSLRREVSRLQGTGGPSQNGEGYPWQSLPEQSSDALEAWENGERS 100
LSAPRIL
RKRRAVLTQKQKKQHSVLHLVPINATSKDDSDVTEVMWQPALRRGRGLQA 150
QGYGVRIQDAGVYLLYSQVLFQDVIFIMGQVVSREGQGRQETLFRCIRSM 200
PSHPDRAYNSCYSAGVFHLHQDILSVIIPRARAKLNLSPHGTFLGFVKL 250
FIG. 1a

## 1 <br> NO:


SEQ ID NO: 2

SEQ ID NO： 3
CAGAGAGCCA TGGGGGCTGT CAAAGCCTAA SYDSYYDYJ ATGTCCTGGA วองษอルปษปอ AGTTAACATT CAGTACTTAG

 AGAGAAACTC L甘甘D甘LDJOL つYOLYLlyLy CCGCATGGAA GTGGGGATTT CAAGAACAGC TTTCCCGCTC

 U皆 GAGGGCCTTC


 U ACATTGTACG



 TAGCCTTTCT TATAAAGGGG AAGGAACGGT
 GGCCACATGG CGAGGCTCCA GTTGAGTTGG TGATCCAACA CAGCGGAGTG TTGCTCTTTG gTCGCACTAC GAGCCGGCTG CCTCTGGGAG
 GAATTTCGGCA GACTTGTGCT GGCGGGAGGT GGAGAGCGCC
-

AATCTCGGAG
 GACAGAGGTG GCCCAGGGAG TAGTCAGGTC

 YLLLYOLLLLD

 GACAAAGGAC
CTGTTGTTCC
 AGCCTGGAAG AGAAGCACAA ACCTCCAAGG GCGTGGGAGA GGCCCTGGAG ACTGGAATTT CACAATGGGT TATTCCGATG AGCTGCTACA TGTCAAAATT CATTCCTGGG CCCATTCCAA TCTCCATGGC TCTCCATGGC CCACTATCTG TGTTTATCTC CCAAAAA FIG．1d tatcagang GTGCAGGTGT CCACGGGCAA GTTTGTGAAA AAACTGGCTA TTTGCCTTGA GGCTTTGACT ATCTGCTCTA CAGGTGGTAT GATGGGGCGA GAAGAAGCAC ACTCTGACGT GGCCCTGGAG
SEQ ID NO: 4

FIG. $1 e$
ACATCAACAA ACACAGATAA CCCCAACCTT AGCAGTCACG AAGGAGTGTG CTCCAAAGAC CTTGCTGCCT GACCTGGCCA GCCAGCAGGA TCACCGCGGG TCCAGTCAGA AGTCACTCAA TACAAAAAGG GGAAGTGCCC TTACTHTLTT TGGGACATCT AGTCTGGTGA CAATAATTCC AACTCCAACT GATGTCACAT GTCTGTAGCT TAACTGAAAA ACATCAACAA
ATTCTAAAGG
ACAGAAAGGG
AATGAAACTG
CTGTCCGATC GCACTGCTGT CCTGCAAGGG CGGAGAAGCT GCTCCAGCTG AGAAGGCAAC CAGAAGAAAC ACACCAACTA CTMTAAAAGG AAGAAACTGG ACCTACGCCA GGATGAATTG AAACACTACC GAAGGAGATG ACTGGATGGA CTTACACCAT AGAAAGAATC

## TGCACGCAGG

 GTGGTCACTM GGATGACTCC AAAGAGAAGAGTTCAGGGTC AGACAGTGAA皆 ATATTGGTCA TACTGATAAG ATGTCTTITGG AATATGCCTG AAAACTGGAA
 TGGCTGCAAC TCTTTCTACC AGAGCTGCAG TTTGAACCAC TAAGCGTGCC AACTGATTGC TTTGTTCCAT AGAGAATAAA AGGTMTTATA AAGAAGGTCC ATGTATTCAA CTGGCATIGC AGAGAAAATG ATTGAAACTG CTTTCTCTGT TGCCAAGCCC CAGGAAATGA CAAAGTTCAA CCTMACTTCT TITCCATCCT GGAAAGCTGC CACGGTGGTG GCCTCCGGGC GCAGGAGCCC ACTGAAAATC ACAGCAGAAA GACTGCTTGC ATCTTACACA TAGAAGAAAA ATATATGGTC AATTCAGAGG CITIGTTICG TGCTATICAG TGCAATACCA TITITGGTGC ATMTTCCTCC TA ID.
1
51
101
151
201
251
301
351
401
451
501
551
601
651
701
751
801
851
901
951

FIG. 2b

## ID NO． 7



FIG. 2d



FIG. 4

## HETEROLOGOUS POLYPEPTIDE OF THE TNF FAMILY

## RELATED APPLICATIONS

[0001] This is a continuation of PCT/US01/04121, filed on Feb. 8, 2001, which claims priority from U.S. provisional application Ser. No. 60/181,670 filed on Feb. 11, 2000.

## TECHNICAL FIELD

[0002] This invention relates, in part, to a newly identified heteromeric ligand of the Tumor Necrosis Factor (TNF)family, referred to hereinafter as "APBF", its variants, derivatives, agonists and antagonists; and uses thereof. In particular, the invention relates to an APBF having a TNFfamily member APRIL subunit linked non-covalently to a TNF-family member BAFF subunit.

## BACKGROUND OF THE INVENTION

[0003] Tumor Necrosis Factor (TNF)-family members can best be described as master switches in the immune system controlling both cell survival and differentiation. Given the current progress in manipulating members of the TNFfamily for therapeutic benefit, including anti-tumor activity as well as immune regulation and inflammation, it is likely that members of this family will provide unique means to control disease. The medical utility of the TNF ligands and antagonists to the ligands has been shown for several systems. Most notable is TNF. TNF controls a wide array of immune processes, including inducing acute inflammatory reactions, as well as maintaining lymphoid tissue homeostasis. Because of the dual role this cytokine can play in various pathological settings, both agonist and antagonist reagents have been developed as modifers of disease. For example TNF and LT $\alpha$ (which also signals through the TNF receptors) have been used as a treatment for cancers, especially those residing in peripheral sites, such as limb sarcomas. In this setting direct signaling by the cytokine through the receptor induces tumor cell death (Aggarwal and Natarajan, 1996. Eur Cytokine Netw 7:93-124). In immunological settings agents which block TNF receptor signaling (eg., anti-TNF mAb , soluble TNF-R fusion proteins) have been used to treat diseases like rheumatoid arthritis and inflammatory bowel disease. In these pathologies, TNF is acting to induce cell proliferation and effector function, thereby exacerbating autoimmune disease. In this setting blocking TNF binding to its receptor(s) has therapeutic benefit (Beutler, 1999. J Rheumatol 26 Suppl 57:16-21).
[0004] A more recently discovered ligand/receptor system appears amenable to similar manipulations. Lymphotoxin beta (LT $\beta$ ), a TNF family member which forms heterotrimers with LT $\alpha$, binds to the LT $\beta$-R. Some adenocarcinoma tumor cells which express LT $\beta$-R can be killed or differentiated when treated with an agonistic anti-LT $\beta-\mathrm{R} \mathrm{mAb}$ (Browning et al., 1996. J Exp Med 183: 867-878). In immunological settings it has been shown that anti-LT $\alpha$ mAb or soluble receptor fusion protein LT $\beta$-R-Ig can block the development of inflammatory bowel diseases, possibly by influencing dendritic cell and T cell interaction (Mackay et al., 1998. Gastroenterology 115:1464-1475).
[0005] In addition to the TNFR and LT $\beta$-R systems, manipulation of the TRAIL (Gura, 1997. Science 277: 768) and OPG (Simonet et al. 1997. Cell 89: 309-319) pathways
may be therapeutically beneficial in treating cancer and bone loss, respectively. Recently, through database searches, there has been a number of newly described members of the TNF family of ligands and receptors. In addition to the number of new members, the complexity of the ligand/receptor interactions has also increased. It is now apparent that the TNF and LT systems are not unique in the ability of the ligand to interact with more than one receptor. Among the ligands reported to bind more than one receptor or receptor decoy are FasL, TRAIL, RANKL, and LIGHT.
[0006] Thus, there is a clear need to identify and characterize additional molecules which are members of the TNF family thereby providing additional means of controlling disease and manipulating the immune system.,

## SUMMARY OF THE INVENTION

[0007] The present invention relates to the identification of a newly discovered heteromer in the TNF-family, APBF, its nucleotide sequences, its protein sequences and resulting polynucleotides, polypeptides as well as to its soluble form; receptor to the APBF and antibodies specific for APBF and its receptor; and uses therefrom.
[0008] The invention relates to an isolated polypeptide comprising an APRIL subunit linked via a non-covalent interaction to a BAFF subunit. In one aspect the invention is directed to an isolated polypeptide comprising an APRIL subunit selected from the group consisting of human APRIL, partial human APRIL, murine APRIL or partial murine APRIL, or amino acid substitution variants thereof; linked via non-covalent interaction to a BAFF subunit selected from the group consisting of human BAFF, partial human BAFF, murine BAFF or partial murine BAFF, or amino acid substitution variants thereof. In preferred embodiments, the partial BAFF or APRIL polypeptides are soluble portions of the polypeptides.
[0009] In preferred embodiments of the invention, the heterologous polypeptide comprises more than one APRIL subunit, and more preferably two APRIL subunits, linked non-covalently to a BAFF subunit. In alternative embodiments, the heterologous polypeptide comprises more than one BAFF subunit, and more preferably two BAFF subunits, linked 10 non-covalently to an APRIL subunit. Thus, in preferred embodiments, the present invention is directed to heterologous polypeptide trimers of BAFF and APRIL subunits, in which the ratio of APRIL to BAFF subunits is 2:1, or alternatively $1: 2$.
[0010] The present invention also relates to therapeutic methods utilizing the heteromers of the invention. One aspect of the invention relates to methods of inhibiting B-cell, T-cell or tumor cell growth in an animal by administering a therapeutically effective amount of a composition selected from the group consisting of an isolated APBF molecule or active fragment thereof, a recombinant APBF molecule or active fragment thereof, and an antibody specific for APBF or an active fragment thereof. Another aspect of the invention relates to methods of stimulating B-cell or T-cell growth in an animal by administering a therapeutically effective amount of a composition selected from the group consisting of an isolated APBF molecule or active fragment thereof, a recombinant APBF molecule or active fragment thereof, and an antibody specific for APBF or an active fragment thereof.

## BRIEF DESCRIPTION OF THE DRAWINGS

[0011] The following drawings depicts certain embodiments of the invention. They are illustrative only and do not limit the invention otherwise disclosed herein.
[0012] FIG. $1 a$ shows the amino acid sequence of human APRIL (SEQ ID NO: 2). The predicted transmembrane region (TM, boxed), the potential N-linked glycosylation site (star) and the N -terminus of the recombinant soluble APRIL sequences are indicated. FIG. $1 b$ shows the DNA sequence encoding human APRIL (SEQ ID NO.: 1), the amino acid sequence of human APRIL (SEQ ID NO.: 2) is shown in FIG. 1c, the DNA sequence encoding mouse APRIL (SEQ ID NO.: 3) is shown in FIG. $1 d$, and the amino acid sequence of mouse APRIL (SEQ ID NO.: 4) is shown in FIG. $1 e$.
[0013] FIG. $2 a$ shows the DNA sequence encoding human BAFF (SEQ ID NO.: 5), the amino acid sequence of human BAFF (SEQ ID NO.: 6) is shown in FIG. 2b, the DNA sequence encoding mouse BAFF (SEQ ID NO.: 7) is shown in FIG. $2 c-2 d$, and the amino acid sequence of mouse BAFF (SEQ ID NO.: 8) is shown in FIG. 2d. Amino acids 1 to 46 from SEQ ID NO.: 6 represent the intracellular domain, amino acids 47 to 72 from SEQ ID NO.: 6 represent the transmembrane domain and amino acids 73 to 285 from SEQ ID NO.: 6 represent the extracellular domain.
[0014] FIG. 3 shows a comparison of two western blots from cells co-transfected with various APRIL and BAFF encoding plasmids. The detection reagent used in Panel A is an anti-FLAG antibody. The detection reagent used in Panel $B$ is an anti-BAFF antibody.
[0015] FIG. 4 shows a western blot of the immunoprecipiations of conditioned media from cells co-transfected with plasmids encoding various soluble APRIL and soluble BAFF proteins and immunoprecipitated with an anti-FLAGtagged antibody. The detection reagent for the western blot is an anti-myc tagged antibody.

## DETAILED DESCRIPTION

## [0016] Definitions

[0017] The term "APBF" or "APBF ligand" when used herein encompasses any native or recombinantly produced polypeptide having an APRIL subunit linked via a noncovalent interaction to a BAFF subunit. APBF may be isolated from a variety of sources, such as from murine or human tissue types or from other sources, or prepared by recombinant or synthetic methods. A large number of analytical biochemistry methods, known to those of skill in the art, can be utilized to determine the stoichiometry of APBF, its variants and derivatives. For example, cation exchange chromatography can be used to determine which of the various stoichiometric forms are present in the preparation derived from affinity columns. Also gel chromatography of the purified fractions will show the molecular weights of each form. The molecular weights of APRIL and BAFF are known. For example, the molecular weight of full length human BAFF, amino acids $1-285$, is predicted to be 34.2 kDa for each polypeptide. The molecular weight of soluble human BAFF, amino acids A132-285, is predicted to be 18.2 kDa per polypeptide. The molecular weight of full length human APRIL, amino acids 1-250, is predicted to be 30.0 kDa for each polypeptide. The molecular weight of soluble
human APRIL, amino acids A105-250, is predicted to be 17.5 kDa per polypeptide. Stoichiometric combinations contemplated in the present invention include the following formula, X APRIL: Y BAFF, where X and Y are integers greater than or equal to one. It is contemplated that the heteromer may exist as a soluble molecule, wherein all subunits are of soluble APRIL or BAFF polypeptides. It is further contemplated that the heteromer may exist as a cell associated molecule, wherein at least one of the subunits is the full length molecule containing a transmembrane domain and the other subunit(s) may contain either full length or soluble forms of APRIL or BAFF.
[0018] The term "APRIL subunit" when used herein encompasses any native or recombinantly produced APRIL polypeptide. The APRIL subunit may be isolated from a variety of sources, such as from murine or human tissue types or from other sources, or prepared by recombinant or synthetic methods. For example, an APRIL subunit can have an amino acid sequence encoded by human APRIL (SEQ ID NO.: 1) or murine APRIL (SEQ ID NO.: 3) and variants, derivatives and unique fragments thereof. Specifically contemplated are human and murine soluble construct forms of APRIL (see above, and SEQ ID NOs.: 2 and SEQ ID NO.: 4) and variants, derivatives and unique fragments thereof.
[0019] The term "BAFF subunit" when used herein encompasses any native or recombinantly produced BAFF polypeptide. The BAFF subunit may be isolated from a variety of sources, such as from murine or human tissue types or from other sources, or prepared by recombinant or synthetic methods. For example, a BAFF subunit can have an amino acid sequence encoded by human BAFF (SEQ ID NO.: 5) or murine BAFF (SEQ ID NO.: 7) and variants, derivatives and unique fragments thereof. Specifically contemplated are human and murine soluble construct forms of BAFF (see above, and SEQ ID NO.: 6 and SEQ ID NOs.: 8) and variants, derivatives and unique fragments thereof.
[0020] As defined herein, a "unique fragment" of a protein or nucleic acid is a peptide or oligonucleotide of sufficient length to have a sequence unique to a particular gene or polypeptide, i.e., a sequence not shared by related or unrelated genes or polypeptides. Thus, for example, a unique nucleic acid fragment typically will have at least 16 nucleotide residues, and a unique polypeptide fragment typically will have at least 6 amino acid residues. Preferably, to ensure substantially unique occurrence in a typical higher eukaryotic genome, a unique nucleic acid fragment should have at least 20 nucleotide residues, and a unique polypeptide fragment should have at least 8 amino acid residues.
[0021] An "isolated" polypeptide, polynucleotide, protein, antibody, or other substance refers to a preparation of the substance devoid of at least some of the other components that may also be present where the substance or a similar substance naturally occurs or is initially obtained from. Thus, for example, an isolated substance may be prepared by using a purification technique to enrich it from a source mixture. Enrichment can be measured on an absolute basis, such as weight per volume of solution, or it can be measured in relation to a second, potentially interfering substance present in the source mixture. Increasing enrichments of the embodiments of this invention are increasingly more preferred. Thus, for example, a 2 -fold enrichment is preferred, 10 -fold enrichment is more preferred, 100 -fold enrichment
is more preferred, 1000 -fold enrichment is even more preferred. A substance can also be provided in an isolated state by a process of artificial assembly, such as by chemical synthesis or recombinant expression. "Hybridization" is the noncovalent, antiparallel bonding of complementary nucleic acid strands, in which Watson-Crick base pairing is established. To ensure specificity, hybridization should be carried out under stringent conditions, defined herein as conditions of time, temperature, probe length, probe and/or target concentration, osmotic strength, pH , detergent, carrier nucleic acid, etc. that permit no more than an occasional base-pairing mismatch within a probe/target duplex. Highly stringent conditions exclude all but about one base pair mismatch per kb of target sequence. Exemplary highly stringent conditions involve hybridization to membrane immobilized target nucleic acid at a temperature of $65^{\circ} \mathrm{C}$. in the presence of $0.5 \mathrm{~m} \mathrm{NaHPO}, 7 \%$ SDS, ImM EDTA, followed by washing at $68^{\circ} \mathrm{C}$. in the presence of $0.1 \times \mathrm{SSC}$, $0.1 \%$ SDS. Current Protocols in Molecular Biology (1989), Ausubel et al., eds, Greene Publishing and Wiley Interscience, New York, N.Y. In circumstances where relatively infrequent mismatches, e.g., up to about ten mismatches per kb of target, can be tolerated, moderately stringent conditions may be used. For moderate stringency, probe/target hybrids formed under the above conditions are washed at $42^{\circ} \mathrm{C}$. in the presence of $0.2 \times \mathrm{SSC}, 0.1 \%$ SDS.
[0022] An "individual" is a vertebrate, preferably a mammal, more preferably a human. Mammals include, but are not limited to, farm animals, sport animals, pets, primates, mice and rats.
[0023] An "effective amount" is an amount sufficient to effect beneficial or desired clinical results. An effective amount can be administered in one or more administrations. For purposes of this invention, an effective amount is an amount of APBF, variants and derivatives of APBF and agonists and antagonists of APBF that is sufficient to ameliorate, stabilize, or delay development of a disease state associated with APBF. Particularly APBF-associated tumors. Detection and measurement of these indicators of efficacy are discussed below.
[0024] As used herein, "treatment" is an approach for obtaining beneficial or desired clinical results. For purposes of this invention, beneficial or desired clinical results include, but are not limited to, one or more of the following: alleviation of symptoms, diminishment of extent of disease, stabilized (i.e., not worsening) state of disease, preventing spread (i.e., metastasis) of disease, preventing occurrence or recurrence of disease, delay or slowing of disease progression, amelioration of the disease state, and remission (whether partial or total). Also encompassed by "treatment" is a reduction of pathological consequences of an APBFassociated tumor(s).
[0025] As used herein, the term "cancer" refers to any neoplastic disorder, including such cellular disorders as, for example, renal cell cancer, Kaposi's sarcoma, chronic leukemia, breast cancer, sarcoma, ovarian carcinoma, rectal cancer, throat cancer, melanoma, colon cancer, bladder cancer, mastocytoma, lung cancer, mammary adenocarcinoma, pharyngeal squamous cell carcinoma, and gastrointestinal or stomach cancer. Preferably, the cancer is leukemia, mastocytoma, melanoma, lymphoma, mammary adenocarcinoma, and pharyngeal squamous cell carcinoma.
[0026] To determine the "percent homology" of two amino acid sequences or of two nucleic acids, the sequences are aligned for optimal comparison purposes (e.g., gaps can be introduced in the sequence of a first amino acid or nucleic acid sequence for optimal alignment with a second amino acid or nucleic acid sequence). The amino acid residues or nucleotides at corresponding amino acid positions or nucleotide positions are then compared. When a position in the first sequence is occupied by the same amino acid residue or nucleotide as the corresponding position in the second sequence, then the molecules are homologous at that position (i.e., as used herein amino acid or nucleic acid "homology" is equivalent to amino acid or nucleic acid "identity"). The percent homology between the two sequences is a function of the number of identical positions shared by the sequences (i.e., $\%$ homology=\# of identical positions/total \# of positions $\times 100$ ). The determination of percent homology between two sequences can be accomplished using a mathematical algorithim. A preferred, non-limiting example of a mathematical algorithim utilized for the comparison of two sequences is the algorithm of Karlin and Altschul (1990) Proc. Natl. Acad. Sci. USA 87:2264-68, modified as in Karlin and Altschul (1993) Proc. Natl. Acad. Sci. USA 90:5873-77.
[0027] The invention encompasses all nucleic acids, peptides, polynucleotides, polypeptides and proteins of the present invention that can be produced, expressed, and/or manipulated by conventional molecular engineering techniques such as the techniques set forth in Current Protocols in Molecular Cloning, Ausubel et al., eds. (1989), Greene Publishing and Wiley Interscience, New York, N.Y. and in Sambrook et al. (1989), Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y., and the teachings described and referenced in Watson et al. (1992), Recombinant DNA 2nd ed., Scientific American Books and W. H. Freeman \& Co., New York, N.Y.

## DESCRIPTION OF THE INVENTION

[0028] The present invention relates to a newly identified heteromeric member of the TNF-family, APBF, wherein APBF comprises an APRIL subunit linked non-covalently to a BAFF subunit.
[0029] APRIL, a TNF ligand known to have a role in inducing tumor cell proliferation is described in detail in PCT publications WO99/12965, WO97/33902, WO99/ 50416 and U.S. Provisional Application Ser. No. 60/106, 976, each incorporated by reference herein. It has been shown that high levels of APRIL mRNA are detected in several tumor cell lines, as well as in colon carcinomas, metastatic lymphomas and thyroid tumors. Moreover, it has been shown that the in vitro addition of recombinant APRIL stimulates the proliferation of various cell lines. It is also recognized that in addition to inducing tumor cell proliferation that APRIL may modulate a variety of functions of immune system cells in vitro and in vivo (Hahne et al., (1998) J. Exp. Med. 188 :1185-1190).
[0030] The second component of APBF, BAFF, has been shown to have a role in inducing the proliferation of naive B cells and is described in detail in PCT publications WO98/18921, WO98/27114, and WO99/12964, each incorporated by reference herein. Like APRIL, BAFF has also
been shown to modulate a variety of functions of immune system cells in vitro (Schneider et al., (1999) J. Exp. Med. 189: 1747-1756) and in vivo (Mackay et al., (1999) J. Exp. Med.190:1697-1710; Moore et al., (1999) Science.285: 260263).
[0031] To date, all known TNF-family members, with the exception of the lymphotoxins, form homomers. It was therefore a surprising discovery, as a result of the work described herein, to identify a heteromeric polypeptide having an APRIL subunit linked non-covalently to a BAFF subunit FIGS. 1 and 2 provides the full length and partial nucleic acid and amino acid sequences of mammalian APRIL and mammalian BAFF, respectively. The intracellular, transmembrane, and extracellular domains are identified, and a protease cleavage site is marked. N-terminal amino acid sequence analysis of APRIL secreted into the media of EBNA293 cells transfected with the full length murine APRIL cDNA plasmid identified alanine at position 87 as the first amino acid in the secreted form. Similar analysis of human BAFF overexpressed in EBNA293 cells showed that alanine at position 134 (numbering of amino acids corresponds to the naturally occuring human BAFF sequence, as found for example, in Schneider et al. 1999 J. Exp. Med 189: 1747-1756) was the first amino acid of the secreted form through amino acid 285.
[0032] In one embodiment, APBF comprises an APRIL subunit derived from a mammalian APRIL linked via a non-covalent interaction to a BAFF subunit derived from a mammalian BAFF. It is contemplated that subunits of APRIL and/or BAFF may remain cell membrane bound via their transmembrane domains, and comprise part of a mem-brane-bound APBF. Alternatively, the APBF may consist of the natively cleaved forms of APRIL and BAFF extracellular domains, or fragments derived from the natively cleaved forms. As illustrated in Example 1, when FLAG-tagged soluble APRIL is co-expressed with full-length BAFF, the soluble heteromeric complex is formed. This shows that the full-length BAFF is readily cleaved and complexes with the artificially generated soluble APRIL molecule. Alternatively, Example 2 demonstrates that the complex can be formed when both APRIL and BAFF are expressed as soluble molecules. This indicates that the region between the transmembrane and receptor binding domain (stalk) is not required for association. However, if one or more subunits remains uncleaved then the complex will remain tethered to the cell surface. Alternatively, the complex will be secreted. Since additional modification may take place after proteolytic cleavage from the cell surface, other subunit forms are envisioned, for example, one or more subunits may consist of a portion of the extracellular domain, as when the stalk portion (before the first beta sheet) is shortened. Also, as APRIL and BAFF contain glycosylation sites it is conceivable that one or more subunits may be a glycosylated or differentially glycosylated. Such modifications may depend on the cell in which the heteromer is expressed.
[0033] As a result of the work described herein, in which we identified APBF by co-expression and differentially tagging (see Examples 1 and 2), we are able to produce and isolate APBF by any of a number of techniques known to those of skill in the art, including for example, affinity methods, as described for example, in Example 3. Another example of a known method for isolation of proteins include ion exchange chromatography. For example, APBF may
easily be separated by ion exchange chromatography based on the widely ranging pI values for soluble human APRIL (ie: hAPRIL pI=9.81) and soluble human BAFF (hBAFF $\mathrm{pI}=4.75$ ). Heterocomplexes commonly have pI values that are additive in nature. Stoichiometrically different combinations of BAFF and APRIL proteins would be expected to bind DEAE and S-sepharose columns with significantly different affinities at a given pH . Visualization of these heterocomplexes can also be done by isoelectic focusing (IEF), followed by blotting and detection with antibodies. IEF can also be used to isolate small amounts of proteins. Since native IEF generally does not disrupt protein function it may well present itself as a useful way to assay the protein binding affinities cells and receptors and to evaluate transfections for the level of the various heteromers produced Such methods are particularly useful in separating different subunit stoichiometries which may be present after cotransfection.
[0034] The invention further provides degenerate variant nucleic acids that encode the SEQ ID NOS.: 2, 4, 6 and 8 polypeptides or a unique fragment thereof. In yet further embodiments, the invention provides nucleic acids encoding variant APBF polypeptides, comprising amino acid sequences sharing at least $75 \%$ sequence similarity with the SEQ ID NOs.: 2, 4, 6 and/or 8. Preferably, these nucleic acids encode polypeptides sharing at least $80 \%, 85 \%, 90 \%$ or more preferably $95 \%$ amino acid sequence similarity with SEQ ID NOs.: 2, 4, 6 and/or 8. The encoded variant polypeptides comprise amino acid mutations (substitutions, deletions and/or insertions) distributed in any random or non-random frequency within SEQ ID NOs.: $2,4,6$ and/or 8. "Similarity" as used herein refers to the sum of aligned amino acid residues that are identical to the residues of corresponding SEQ ID NOs.: 2, 4, 6 and 8 and those that are allowed point mutations therefor. Moderate gaps and/or insertions (e.g., less than about 50, preferably less than about 15 , more preferably less than about 5 amino acid residues) in the aligned sequence are ignored for similarity calculation purposes. Allowed point mutations are substitutions by amino acid residues that are physically and/or functionally similar to the corresponding aligned residues of SEQ ID NOs.: 2, 4, 6 and/or 8 , e.g., that have similar size, shape, hydrophilic or hydrophobic character, charge and chemical properties.
[0035] It should be understood that the present invention provides oligonucleotides that hybridize to any of the foregoing variant APBF nucleic acids, i.e., to nucleic acids that encode polypeptides comprising amino acid sequences that share at least $75 \%$ sequence -similarity with SEQ ID NOs: 2, 4, 6 and/or 8 . More particularly, the invention provides olignucleotides that hybridize to one or more unique fragments of nucleic acids encoding APBF. For therapeutic purposes and/or for PCR investigative or diagnostic purposes, the present oligonucleotides hybridize to a unique fragment comprising 5 ' untranslated sequence, a transcription initiation site, ORF or polypeptide coding sequence, intron-exon boundary, polyadenylation site or $3^{\prime}$ untranslated region of the present APBF nucleic acids.
[0036] The invention also relates to heteromers formed with partial sequences of human and murine APRIL and human and murine BAFF. Preferably, these partial sequences comprise soluble forms of BAFF and APRIL. Preferred partial human APRIL molecules include amino
acids A105 to L250, K110 to L250 and H115 to L250 of the full-length human APRIL sequence. Preferred partial murine APRIL molecules include amino acids A87 to L233 of the full-length murine APRIL sequence. Preferred partial human BAFF sequences include amino acids A134 to L285 and Q136 to L285 of the full-length human BAFF sequence (see, Schneider et al. 1999, J.Exp. Med. 189:1747-1756, incorporated by reference herein).
[0037] Preferred partial sequences of human and murine APRIL and BAFF also include splice variants of APRIL and BAFF. A preferred partial human APRIL sequence includes a splice variant that is the complete APRIL human sequence missing amino acids $\mathbf{1 1 3}$ to $\mathbf{1 2 8}$ (see, Kelly et al. 2000, Can. Res. 60: 1021-1027, incorporated herein by reference). Preferred partial human BAFF sequences include a splice variant which is the full-length BAFF sequence missing amino acids $\mathbf{1 4 2}$ to 160 (see, WO 00/50597). Preferred partial murine BAFF sequences include a splice variant which is full-length BAFF sequence missing amino acids 166 to 184.
[0038] The invention also encompasses soluble secreted forms of APBF. See Example 2. To create a soluble secreted form of APBF, one would use techniques known to those of skill in the art, including for example removing at the DNA level the N -terminus transmembrane regions encoding either the APRIL and/or BAFF N-terminus transmembrane regions, and some portion of the corresponding stalk region, and replace these regions with a type I leader or alternatively a type II leader sequence that will allow efficient proteolytic cleavage in the chosen expression system. A skilled artisan could vary the amount of the stalk region retained in the secretion expression construct to optimize both receptor binding properties and secretion efficiency. For example, the constructs containing all possible stalk lengths, i.e. N-terminal truncations, could be prepared such that proteins starting at amino acids 105-135, for APRIL and 134-164 for BAFF would result. The optimal length stalk sequence would result from this type of analysis.
[0039] Isolated APBF can be used for a number of purposes, such as the production of monoclonal or polyclonal antibodies, and identification of novel modulators affecting biological function (e.g., inhibitors), and identification of receptors interacting with APBF.
[0040] As a result of the work described herein, antibodies (polyclonal or monoclonal) specific for the identified APBF can be produced, using known methods (See, for example, Antibodies: A Laboratory Manual ed. by Harlow and Lane (Cold Spring Harbor Press: 1988)). Such antibodies and host cells (i.e. hybridoma cells) producing the antibodies are also the subject of the present invention.
[0041] Antibody production involves administration of one or more immunogenic doses of an APBF polypeptide preparation (whether isolated or incorporated in a cell membrane) to an appropriate non-human animal, such as a mouse, rat, rabbit, guinea pig, turkey, goat, sheep, pig, or horse. To enhance immunogenicity, the preparation can be emulsified with a conventional adjuvant, such as Freund's complete or incomplete adjuvant. Routine monitoring of serum immunoglobulins, using peripheral blood samples withdrawn at appropriate intervals (e.g., seven to ten days) after an initial or subsequent immunization, can be used to detect the onset and/or maturation of a humoral immune
response. Detection and, optionally, quantitation, of immunoglobulins selectively reactive with an APBF epitope can be achieved through any conventional technique, such as ELISA, radioimmunoassay, Western blotting, or the like.
[0042] An immunoglobulin "selectively reactive with an APBF epitope" has binding specificity for the recognized epitope such that an antibody/epitope complex forms under conditions generally permissive of the formation of such complexes (e.g., under conditions of time, temperature, ionic strength, pH ionic or nonionic detergent, carrier protein, etc.). Serial dilution (titration) analysis by standard techniques is useful to estimate the avidity of antibodies in the immune serum sample for one or more epitopes unique to APBF. As defined herein, an "epitope unique to APBF" is a unique, immunogenic fragment of the full-length APBF polypeptide. A unique linear epitope typically ranges in size from about ten to about twenty-five amino acid residues, and frequently is about twelve to eighteen residues in length.
[0043] Immune serum having a high titer generally is preferred herein. Serum having a half maximal avidity for a unique APBF epitope of at least about $1: 1000$, preferably at least about $1: 10,000$, can be harvested in bulk for use as a source of polyclonal antibody useful in the detection and/or quantitation of APBF. Polyclonal immunoglobulins can, if desired, be enriched by conventional fractionation of such serum, or can be isolated by conventional immunoadsorbent techniques, e.g., using a Protein A or Protein G chromatography resin. Immune, high titer murine, rat, hamster or guinea pig serum alternatively is preferred herein for the production and screening of hybridomas secreting monoclonal antibodies selectively reactive with APBF. The present hybridomas can be produced according to wellknown, standard techniques. The present monoclonal antibodies can be obtained from hybridoma culture supernatant, or from conventionally produced ascites fluid, and optionally isolated via immunoadsorbent chromatography or another suitable separation technique prior to use as agents to detect and/or quantitate APBF.
[0044] A preferred antibody, whether polyclonal or monoclonal, is selectively reactive with a unique APBF epitope that is displayed on the surface of or secreted from APBF expressing cells. The preferred antibody accordingly can be used to detect and, if desired, quantitate APBF expressing cells, e.g., normal or transformed cells in a mammalian body tissue or biopsy sample thereof. Specifically, the preferred antibody can be used to detect APBF expressing cells whether such cells are host cells or mammalian body tissue cells that aberrantly express APBF. Advantageously, intact, e.g., living, cells that display a unique APBF epitope can be detected by standard immunohistochemical, radiometric imaging or flow cytometry techniques. The present antibody can be used to detect and/or monitor APBF polypeptide production. Thus, the antibody can be used to assess the natural tissue-specific production of APBF, and thus to assess tissues likely to give rise to carcinomas or sarcomas. In addition, the present antibody can be used to monitor tumor biopsy samples to provide information relevant to selecting or revising a course of disease management, or to diagnosis, prognostication and/or staging of any disease associated with an abnormality affecting APBF. Furthermore, the present antibody can be used in a cell-sorting procedure or other cell isolation procedure to generate a substantially pure preparation of APBF expressing cells, or
a cell population substantially depleted of APBF expressing cells. Each of the foregoing can be achieved through routine practice or modification of well-known techniques, including but not limited to the conjugation of a detectable moiety (e.g., a radionuclide, fluorophore, chromophore, binding pair member, or enzyme) to the APBF reactive antibody.
[0045] A hybridoma secreting an APBF reactive monoclonal antibody of the present invention additionally provides a suitable source of nucleic acid for the routine construction of a fusion polypeptide comprising an antigenbinding fragment derived from the APBF reactive antibody. The present fusion polypeptide can be prepared by routine adaptation of conventional techniques described in Deeley et al. (1996), U.S. Pat. No. 5,489,519 (incorporated herein by reference). The fusion polypeptide can be a truncated immunoglobulin, an immunoglobulin having a desired constant region (e.g., IgG in lieu of $\operatorname{IgM}$ ), or a "humanized" immunoglobulin having an APBF reactive Fc region fused to a framework region of human origin. Additional fusion polypeptides can comprise, in addition to an $A P B F$ reactive antigen-binding fragment, a non-immunoglobulin polypeptide such as a cytotoxic polypeptide (e.g., diphtheria toxin, ricin) or a chemoattractant polypeptide that stimulates immune effector cells (cytotoxic T cells, natural killer cells, macrophages) to kill cells that display APBF. Standard techniques well-known in the art can be used to produce appropriate immunoglobulin fusion polypeptides of the present invention.
[0046] Various forms of antibodies can also be made using standard recombinant DNA techniques. For example, humanization techniques have been developed that render non-human Mabs less antigenic in humans. Methods for humanizing Mabs by chimerisation procedures are described in EP0120694, EP0125023, EP-A-0 171496, EP-A-0173 494 and WO $86 / 01533$, each incorporated by reference herein. Chimerisation procedures generally involve preparing antibody having the variable domains from a mouse MAb and the constant domains from a human immunoglobulin. Alternatively, methods for humanizing Mabs by CDR-grafting are described in EP-A-0239400 (Winter), WO90/07861 (Queen), WO91/09967 (CellTech), and WO91/09967 (CellTech), incorporated by reference herein. CDR-grafting generally involves grafting the complementarity determining regions (CDRs) of a mouse MAb onto the framework regions of the variable domains of a human immunoglobulin by site directed mutagenesis using long oligonucleotides. In WO91/09967, the preparation of humanized CDR-grafted antibody products which have specificity for TNF-alpha is described. In particular WO91/ 09967 describes in Example 5, preparation of specific humanized CDR grafted antibodies to human TNF-alpha derived from murine anti-human TNF-alpha Mabs. Using any of these known methods, therefore, antibodies specific for APBF can be produced and isolated.
[0047] The polypeptides and methods disclosed herein enable the identification of receptors which specifically interact with APBF or fragments thereof. For example, the APBF receptor can be cloned using any of the techniques known to those of skill in the art, including for example, one or more of the following approaches.
[0048] For example, one can identify an APBF receptor using expression cloning in mammalian cells. Specifically, a
cDNA expression library is generated from a cell line or cell population shown to express the highest level of the receptor to the protein of interest, i.e. APBF. This approach was shown for the leptin receptor (Tartaglia et al., 1995 Cell 83: 1263-1271). The cDNA library DNA is made as pools of $2-3,000 \mathrm{cDNAs}$ and transfected into an appropriate' cell line which does not express the receptor. A plate assay format may be used to detect expression of the receptor on the surface of the receptor negative cell line using purified APBF. An antibody to one of the subunits or to an epitope tag is used to detect the bound protein of interest, i.e. APBF and an alkaline phosphatase conjugated secondary antibody and alkaline phosphatase substrates are used to visualize the positive cells. The plate wells are screened using a microscope. The complexity of the cDNA pools from the positive wells are reduced and then screened again. The screening continues until transfection of a single cDNA produces a positive signal. The DNA of the cDNA is sequenced and the predicted amino acid sequence analyzed for motifs and structure consistent with members of the TNF receptor family. Other expression cloning formats are available, for example, by panning on ligand coating plates, or by sorting with tagged ligand in FACS analyses.
[0049] In another approach, one can identify an APBF receptor using direct DNA sequence analysis. Specifically, a directional cDNA library is generated from a receptor positive cell line and the $5^{\prime}$ ends sequenced using ABI automated DNA sequencing technology to determine the open reading frame. Programs to look at the cysteine residue spacing, signal and transmembrane sequences can be employed to identify potential TNF receptor family members. Full-length clones will then be isolated, expressed and examined for the ability to bind APBF. The library can also be subtracted with a APBF receptor negative cell line to reduce the complexity of the library.
[0050] In another approach, one can identify an APBF receptor by examining known or orphan receptors. Specifically, purified APBF can be used in FACS, immunoprecipitation, ELISA or Biacore assays against a panel of orphan receptors and those that have known ligands. A receptor for APBF will be positive in all of these assays.
[0051] In yet another approach, one can identify an APBF receptor using protein sequence analysis methods. Specifically, APBF can be cross-linked to the surface of cells determined to be receptor positive using standard reagents. The cross-linked complex can be immunoprecipitated using an antibody to one of the subunits or an epitope tag on the subunit. The complex can be separated on a SDS polyacrylamide, blotted to a membrane and subjected to amino acid analysis. Once amino acid sequencing reveals information about the receptor, degenerate oligonucleotide probes can be synthesized and used to screen a cDNA library made from a receptor positive cell line.
[0052] Alternatively, one can identify an APBF receptor using known biochemical approaches in combination with mass spectometry to specifically identify masses and sequences. Three illustrative examples are set forth below:
[0053] Strategy 1: Binding of [125I]-labelled APBF to different cell lines. After choosing the appropriate cell line, crosslinking of [125]I-APBF to the cells followed by polyacrylamide gel electrophoresis (SDS-PAGE) and autoradiography will reveal a receptor protein that will be analyze by mass spectrometry.
[0054] Strategy 2: Crosslinking of [125I]-labelled APBF to cell lines, followed by immunoprecipitation using specific antibodies and polyacrylamide gel electrophoresis.
[0055] Strategy 3. Choosing appropriate cell lines that bind APBF. Adding purified APBF would allow the heteromer binding to the receptor. Immunoprecipitation using specific antibodies to the heteromer followed by SDS-PAGE and autoradiography will reveal specific bands that will be isolated and analyzed by mass spectrometry to identify the receptor/s.
[0056] The foregoing compositions can be used for a number of purposes, including the assessment (e.g., for diagnostic purposes) of abnormalities in the structure and/or expression of a cellular APBF gene as well as diagnosis of conditions involving abnormally high or low expression of APBF activities. For example, in WO/99/12965, it was shown that while transcript of APRIL are of low abundance in normal tissues, high levels of mRNA are detected in several tumor cells lines. The expression and growth stimulating effect of APRIL on tumor cells suggested a role for APRIL in tumorigenesis. Similarly, the monitoring of APBF transcript or polypeptide production, or gene expression level, or fluctuations therein, in one or more tumor biopsy samples is expected to provide information relevant to diagnosis, prognostication and/or staging of neoplastic disease in a cancer sufferer.
[0057] Any suitable means for detecting APBF transcript or polypeptide production or stabilization, or gene expression level, can be applied for the present diagnostic purposes. Appropriate methods are described in Sambrook et al. (1989), Molecular Cloning: A Laboratory Manual, 2nd ed., Cold Spring Harbor Laboratory Press, Cold Spring Harbor, N.Y.
[0058] Standard methods of analysis allow the detection of activity by cells in response to ligand binding. For example preparations of the APBF heteromer will be useful in analyses of cellular proliferation, differentiation, and apoptosis. Numerous cell types can be rapidly screened in such a manner using standard methods such as radioactivethymidine incorporation, cell cycle analysis, and MTT uptake and conversion (detailed in Celis et al., Cell Biology, A Laboratory Handbook, Volume One, Academic Press, San Diego, Calif. (1997). Other methods of analysis that can be used to assess activity include protein phosphorylation analysis, for example, of Nuclear Factor $\kappa B$ transcription factor ( NF KB ) or c-Jun N-terminal Kinase (JNK) (eg., Mackay et al., J. Biol. Chem. 271: 24934-24938 (1996); Wong et al., J. Biol. Chem. 272: 25190-25194 (1997)). Other readily accessable assays include measurements of cytokine secretion (eg. II-8: Chicheportiche et al., J. Biol. Chem. 272: 32401-32410 (1997)), calcium flux, pH change, cell/cell adhesion, etc (with references).
[0059] In addition to these readouts, analyses of upregulated or downregulated genes are readily done, for, by example, Northern blot, targeted array, or gene array analyses ( eg. Teague et al., Proc. Natl. Acad. Sci. USA 96 :12691-12696 (1999); Lockhart et al., Nat. Biotechnol. 14: 1675-1680 (1996)). Such differential gene expression studies identify specific sets of genes which respond to ligand activity, and can provide detailed profiles of ligand function (eg. Jiang et al., Oncogene 11: 1179-1189 (1995)). Particularly sensitive to such analyses are modifiers of cell growth,
eg. growth hormone receptor genes, transcription factors, genes whose proteins induce or block cell death, and cell cycle mediators, among many others.
[0060] The present invention is useful for diagnosis or treatment of various immune system-related disorders in mammals, preferably humans. Such disorders include but are not limited to cancer, including, but not limited to, cellular disorders as, for example, renal cell cancer, Kaposi's sarcoma, chronic leukemia, breast cancer, sarcoma, ovarian carcinoma, rectal cancer, throat cancer, melanoma, colon cancer, bladder cancer, mastocytoma, lung cancer, mammary adenocarcinoma, pharyngeal squamous cell carcinoma, and gastrointestinal or stomach cancer. Additionally, the present invention is useful for the treatment of proliferative conditions that are not considered to be tumors, i.e. cellular hyperproliferation (hyperplasia), such as, for example, scleroderma, pannus formation in rheumatoid arthritis, postsurgical scarring and lung, liver and uterine fibrosis. In addition, the present invention is useful for the treatment of immunodeficiencies, inflammatory diseases, lymphadenopathy, autoimmune diseases, and graft versus host disease.
[0061] In one embodiment, conditions caused by a decrease in the normal level of APBF activity in an individual can be treated by administration of APBF or an agonist to APBF, where an agonist to APBF refers to any natural or synthetic composition which potentiates function, where function refers to any measurable effect of APBF interaction with cells, tissues or organisms, as measured in any known in vitro or in vivo assays, which is mediated by APBF. In one embodiment, APBF is in soluble form. The invention also provides a method of treatment of disorders caused by an increase in the normal level of APBF activity in an individual by administration of an antagonist to APBF, where an antagonist to APBF refers to any natural or synthetic composition that blocks function, where function refers to any measurable effect of APBF interaction with cells, tissues or organisms, as measured in any known in vitro or in vivo assays, which is mediated by APBF heteromers.
[0062] Pharmaceutical compositions of the invention may comprise a therapeutically effective amount of APBF, or its receptor, or fragments or mimetics thereof, and, optionally may include pharmaceutically acceptable carriers. Accordingly, this invention provides methods for treatment of cancer, and methods of stimulating, or in certain instances, inhibiting the immune system, or parts thereof by administering a pharmaceutically effective amount of a compound of the invention or its pharmaceutically acceptable salts or derivatives. In certain preferred embodiments, the invention relates to methods for inhibiting B-cell growth, T-cell growth or tumor cell growth by administering a therapeutically effective amount of an isolated APBF polypeptide or active fragment thereof, or a recombinant APBF molecule or active fragment thereof, or an antibody specific for APBF or active fragment thereof. In the context of this invention "inhibition" relates to any and all mechanisms for reducing or ameliorating activity, including inducing cell death (apoptosis). It should of course by understood that the compositions and methods of this invention can be used in combination with other therapies for various treatments.
[0063] The compositions can be formulated for a variety of routes of administration, including systemic, topical or
localized administration. For systemic administration, injection is preferred, including intramuscular, intravenous, intraperitoneal, and subcutaneous for injection, the compositions of the invention can be formulated in liquid solutions, preferably in physiologically compatible buffers such as Hank's solution or Ringer's solution. In addition, the compositions may be formulated in solid form and, optionally, redissolved or suspended immediately prior to use. Lyophilized forms are also included in the invention.
[0064] The compositions can be administered orally, or by transmucosal or transdermal means. For transmucosal or transdermal administration, penetrants appropriate to the barrier to be permeated are used in the formulation. Such penetrants are known in the art, and include, for example, for transmucosal administration, bile salts, fusidic acid derivatives, and detergents. Transmucosal administration may be through nasal sprays or using suppositories. For oral administration, the compositions are formulated into conventional oral administration forms such as capsules, tablets, and tonics. For topical administration, the compositions of the invention are formulated into ointments, salves, gels, or creams as known in the art.
[0065] The dose and dosing regimen will depend on the type of disease, the patient and the patient's history. In one embodiment the disease is cancer. The amount must be effective to treat, suppress, or alter the progression of the disease. The doses may be single doses or multiple doses. If multiple doses are employed, as preferred, the frequency of administration will depend, for example, on the type of host and and type of disease, dosage amounts etc. For some types of cancers or cancer lines, daily administration will be effective, whereas for others, administration every other day or every third day will be effective. The amount of active compound administered at one time or over the course of treatment will depend on many factors. For example, the age and size of the subject, the severity and course of the disease being treated, the manner and form of administration, and the judgments of the treating physician. However, an effective dose may be in the range of from about 0.005 to about $5 \mathrm{mg} / \mathrm{kg} /$ day, preferably about 0.05 to about $0.5 \mathrm{mg} / \mathrm{kg} /$ day . The dosage amount which will be most effective will be one which results in no tumor appearance or complete regression of the tumor, and is not toxic to the patient. One skilled in the art will recognize that lower and higher doses may also be useful.

## EXAMPLES

## Example 1

[0066] This example describes the detection of APRIL and BAFF heteromers by immunoprecipitation following cotransfection into mammalian cells.
[0067] Methods:
[0068] The plasmids encoding FLAG-tagged human soluble APRIL, beginning with residue A105 (LT033) or K110 (PL448), soluble FLAG-tagged human TWEAK beginning at A106 (PS288) or soluble FLAG-tagged human EDA beginning at A242 (PS548) or empty vector (CH269) were co-transfected with a full-length human BAFF construct (PS544) into 293T cells using lipofectamine (Life Technologies). At 48 hrs. post-transfection, conditioned media was collected and used for immunoprecipitation
experiments. The immunoprecipitation samples contained $200 \mu \mathrm{l}$ of conditioned media, $5 \mu \mathrm{~g} / \mathrm{ml}$ of the anti-FLAG antibody M2 (Sigma) and $800 \mu \mathrm{l}$ of DMEM containing $10 \%$ FCS, glutamine, Pen-Strep, G418 and sodium azide and were incubated at $4^{\circ} \mathrm{C}$. for 1 hour, with agitation. Then, 30 $\mu \mathrm{l}$ of ProteinA-Sepharose beads (Pharmacia) was added to the samples and the mixture was incubated overnight at $4^{\circ}$ C. with agitation. The beads were collected by centrifugation and then washed one time with the DMEM media described above and then 3 times with PBS. The final pellet containing the beads was then suspended in $2 \times$ SDS nonreducing sample buffer and boiled for 5 minutes. The beads were spun out and $25 \mu \mathrm{l}$ of the supernatant was loaded onto 2 separate $4-20 \%$ SDS-PAGE gradient gels (Novex). In order to examine the level of ligand expression, non-immunoprecipitated conditioned media from the co-transfected cells were also loaded. These samples were diluted two fold with $2 \times$ non-reducing sample buffer, boiled for 2 min . and then $25 \mu \mathrm{l}$ was loaded into each lane. Each gel contained a set of immunoprecipitations and a set of non-immunoprecipitated conditioned media. After the gels were transferred to Immobilon (Millipore) filters using a BioRad apparatus, the filters were blocked in 5\% non-fat dry milk diluted in TBST for 1 hr at room temperature. The filters were then separated and one was incubated with $5 \mu \mathrm{~g} / \mathrm{ml}$ of the biotinylated anti-FLAG antibody, M2 and the other was incubated with $1 \mu \mathrm{~g} / \mathrm{ml}$ of an anti-human BAFF antibody 53.14 (rat IgM) for about 1 hr at room temperature. The filters were washed with 3 changes of TBST and then incubated in a 1:3000 dilution of streptavidin-HRP or antirat IgM-HRP (Jackson ImmunoResearach) for 30 min . at room temperature. The filters were again washed 3 times and then detected using ECL reagents (Amersham). The filters were exposed to x-ray film for various lengths of time.
[0069] The results of the co-expression experiment are shown in FIG. 3. Panel A, lanes 1-5, show western blots of straight conditioned media from cells co-transfected with various human soluble TNF family ligands and human full length BAFF encoding plasmids. Panel A, lanes 7-12, show western blots of the straight supernatant after immunoprecipitation with an anti-FLAG antibody. The detection reagent used in Panel A is an anti-FLAG antibody. Lane 1: FLAG-tagged human soluble APRIL A105 (beginning at residue A105)+human full length BAFF; Lane 2: FLAGtagged human soluble APRIL K110 (beginning at residue K110)+human full length BAFF; Lane 3: FLAG-tagged human soluble TWEAK+human full length BAFF; Lane 4: FLAG-tagged human EDA+human full length BAFF; Lane 5: empty control vector; Lane 6: molecular weight standards (Benchmark, LifeTechnologies) in $\mathrm{kDa}, 185,119,85,62$, $51,38.2,26.0,20.2,14.5,9.1$; Lane $7-11$ correspond to Lanes 1-5, respectively, after immunoprecipitation with an anti-FLAG antibody; Lane 12: purified human FLAG-BAFF Q136, 5 ng .
[0070] Panel B, lanes 1-5, show western blots of straight supernatant from cells co-transfected with various APRIL and BAFF encoding plasmids. The detection reagent is an anti-BAFF antibody. Panel B, lanes 7-11 show western blots of immunoprecipitates in which cells were co-transfected with various APRIL and BAFF encoding plasmids and immunoprecipitated with an anti-FLAG antibody. The detection reagent is an anti-BAFF antibody. Lane 1: FLAGtagged human soluble APRIL A105 (beginning at residue A105)+human full length BAFF; Lane 2: FLAG-tagged
human soluble APRIL K110 (beginning at residue K110)+ human full length BAFF; Lane 3: FLAG-tagged human soluble TWEAK A106+human full length BAFF; Lane 4: FLAG-tagged human EDA A242+human full length BAFF; Lane 5: empty control vector; Lane 6: molecular weight standards (Benchmark, LifeTechnologies) in kDa, 185, 119, $85,62,51,38.2,26.0,20.2,14.5,9.1$; Lane 7-11 correspond to Lanes 1-5, respectively, after immunoprecipitation with an anti-FLAG antibody; Lane 12: purified human FLAGBAFF Q136, 5 ng .
[0071] In panel A, the detection with the anti-FLAG antibody, M2 shows that all the FLAG-epitope tagged soluble ligands are expressed and secreted into the cell culture of the transfected cells. The two APRIL constructs shown in lanes $\mathbf{1}$ and $\mathbf{2}$ are expressed about 5 fold lower than TWEAK (lane 3) or EDA (lane 4). No protein is visible in lane 5, which is the control vector lane. Lanes 7-11 represent the FLAG-tagged proteins after they are immunoprecipitated. Here both APRIL proteins (lanes 7 and 8 ), TWEAK (lane $\mathbf{9}$ ) and EDA (lane 10) are precipitated and detected by M2. Lane $\mathbf{1 2}$ is approximately 5 ng of FLAG-BAFF for a standard.
[0072] In panel B, lanes $\mathbf{1 - 5}$ represent the detection of BAFF in the co-transfected cell culture media. BAFF is expressed in combination with all the FLAG-epitope tagged soluble ligands and is slightly higher in the APRIL cotransfections (lanes 1 and 2). On the right side of the blot in panel B are the M2 immunoprecipitations detected with the anti-BAFF antibody, 53.14. Here, BAFF is only immunoprecipitated in combination with APRIL (lanes 7 and 8 ) and not TWEAK or EDA. The standard in lane 12 indicates the size of a soluble FLAG-BAFF molecule expressed in 293T cells, which is approximately the molecular weight of the naturally cleaved molecule. This demonstrates that BAFF and APRIL form a heteromeric complex. The co-immunoprecipitations have also been evaluated in the presence of 1 M NaCl and the results are the same.

## Example 2

[0073] This example describes the detection of APBF heteromers by immunoprecipitation following co-transfection of two soluble constructs into mammalian cells.
[0074] Methods:
[0075] Plasmids encoding the following human soluble TNF family ligands were constructed with the indicated N -terminal epitope tags beginning at the ligand amino acid residue indicated in a PCR3 based mammalian cell expression vector: FLAG-APRIL, beginning with residue A105 (plasmid \#LT033) or H115 (plasmid \#LT038), FLAGTWEAK A106 (plasmid \#PS288), myc-APRIL A105 (plasmid \#JST557), and myc-BAFF Q136 (plasmid \#JST556). Various constructs encoding FLAG-tagged ligands, full length murine APRIL(plasmid \#LT022), or empty vector control (plasmid \#CH269) were each co-transfected with the myc-BAFF Q136 construct into 293T cells using lipofectamine (Life Technologies, Gaithersburg, Md.). At 48 hrs. post-transfection, conditioned media was collected and used for immunoprecipitation experiments. The immunoprecipitation samples contained $100 \mu \mathrm{l}$ of conditioned media, 5 $\mu \mathrm{g} / \mathrm{ml}$ of the anti-FLAG antibody M2 (Sigma, St Louis, Mo.) and $900 \mu \mathrm{l}$ of DMEM containing $10 \%$ FCS, glutamine, Pen-Strep, G418 and sodium azide and were incubated at $4^{\circ}$
C. for 1 hour, with agitation. Then, $30 \mu \mathrm{l}$ of ProteinASepharose beads (Amersham Pharmacia, Piscataway, N.J.) was added to the samples and the mixture was incubated overnight at $4^{\circ} \mathrm{C}$. with agitation. The beads were collected by centrifugation and then washed one time with the DMEM media described above and then 3 times with PBS. The final pellet containing the beads was then suspended in $2 \times$ SDS non-reducing sample buffer and boiled for 5 minutes. The beads were spun out and $25 \mu \mathrm{l}$ of the supernatant was loaded onto a $4-20 \%$ SDS-PAGE gradient gel (Novex, San Diego, Calif.). After the gel was transferred to Immobilon (Millipore, Bedford, Mass.) using a BioRad apparatus, the filters were blocked in $5 \%$ non-fat dry milk diluted in TBST for 1 hr at room temperature. The filter was then incubated with $1 \mu \mathrm{~g} / \mathrm{ml}$ of anti-myc antibody 9E10. The filter was washed with 3 changes of TBST and then incubated in a $1: 3000$ dilution of anti-mouse IgG-HRP (Jackson ImmunoResearch, West Grove, Pa.) for 30 min . at room temperature. The filter was again washed 3 times and then detected using ECL reagents (Amersham Pharmacia, Piscataway, N.J.). The filter was exposed to x-ray film for various lengths of time.
[0076] The results shown in FIG. 4 show a western blot of the immunoprecipiations of conditioned media from cells co-transfected with plasmids encoding various soluble APRIL and soluble BAFF proteins and immunoprecipitated with an anti-FLAG-tagged antibody. The detection reagent for the western blot is an anti-myc antibody, 9E10.
[0077] Lanes 1-6, show western blots of straight conditioned media from cells co-transfected with plasmids encoding various human soluble TNF family ligands and human soluble myc-BAFF Q136. Lane 7 is a molecular weight marker. Lanes $\mathbf{8 - 1 2}$, show western blots of the conditioned media after immunoprecipitation with an anti-FLAG antibody. The detection reagent used an anti-MYC antibody, 9E10. Lane 1: FLAG-tagged human soluble TWEAK A106+human soluble myc-BAFF Q136; Lane 2: FLAGtagged human soluble APRIL HI 15+human soluble mycBAFF Q136; Lane 3: MYC-tagged human soluble APRIL A105+human soluble myc-BAFF Q136; Lane 4: FLAGtagged human soluble APRIL A105+human soluble mycBAFF Q136; Lane 5: full length murine APRIL+human soluble myc-BAFF Q136; Lane 6: empty vector control+ human soluble myc-BAFF Q136; Lane 7, molecular weight standards (Benchmark, LifeTechnologies) in $\mathrm{kDa}, 38.2$, 26.0, 20.2, 14.5; Lanes 8-12 correspond to Lanes 1-5, respectively, after immunoprecipitation with an anti-FLAG antibody.
[0078] In lanes 1-6, the western blot of the conditioned media as detected by anti-MYC antibody 9E10 shows that all co-transfected 293T cells express and secrete myc-Baff Q136 into the cell culture media in nearly equal amounts except for the FLAG-TWEAK+myc-Baff Q136 (lane 1) which shows significantly lower amounts of myc-BAFF. Lanes 8-12 show immunoprecipitation of conditioned media with anti-FLAG antibody followed by detection on western blot with anti-myc antibody 9E10. Lanes $\mathbf{1 0}$ and $\mathbf{1 2}$ show conditioned media of myc-BAFF co-transfected with mycAPRILA105 or full length murine APRIL, respectively, and serve as negative controls in that neither APRIL construct contains the flag epitope and therefore were not immunoprecipitated by the anti-flag antibody. The FLAG-TWEAK co-transfection shows no band corresponding to myc-BAFF Q136, even upon overexposure, and therefore does not
interact with MYC-BAFF. Only lanes 9 and 11, those containing myc-Baff Q136 co-expressed with FLAG-APRIL molecules H115 and A105 respectively, show the myc-baff band after anti-FLAG immunoprecipitation. Bands of approximately 18 kDa , the predicted size of myc-BAFF Q136, are observed in each lane. The intensity of the band co-expressed with FLAG-APRIL A87 (lane 3) greater than that co-expressed with FLAG-APRIL H97 (lane 4). This indicates that only soluble FLAG-APRIL ligands were able to interact with soluble MYC-BAFF to form heteromeric complexes.
[0079] This demonstrates that soluble forms of BAFF and APRIL have the ability to form a heteromeric complex, and that no cell associated form appears to be required for heteromer formation.

## Example 3 <br> Production and isolation of APBF by Affinity Methods

[0080] Plasmids encoding the following human soluble TNF family ligands were constructed with N -terminal FLAG or $6 \times$ His epitope tags beginning at the amino acid residue indicated in a PCR3 based mammalian cell expression vector: FLAG-APRIL, beginning with residue A87 (plasmid Lf133) and RGS(H)6-BAFF Q134. These plasmids are then co-transfected into 293 T cells using lipofectamine (Life Technologies, Gaithersburg, Md.) and at 48 hrs. posttransfection, conditioned media is collected. The conditioned media is dialyzed against $50 \mathrm{mM} \mathrm{NaH} 2 \mathrm{PO} 4, \mathrm{pH} 8.0$; $300 \mathrm{mM} \mathrm{NaCl} ; 10 \mathrm{mM}$ imidazole and run over a Ni - NTA Superflow column (Qiagen, Valencia, Calif.). Homomers and heteromers containing the $6 \times$ His tagged BAFF subunit bind to the Ni column; homomeric FLAG-APRIL molecules flow through. The column is washed with 50 mM NaH 2 PO 4 , pH8.0; $300 \mathrm{mM} \mathrm{NaCl} ; 20 \mathrm{mM}$ imidazole with $5-10$ column volumes. The column is eluted with 5 column volumes with $50 \mathrm{mM} \mathrm{NaH} 2 \mathrm{PO} 4, \mathrm{pH} 8.0 ; 300 \mathrm{mM} \mathrm{NaCl} ; 250 \mathrm{mM}$ imidazole. The eluted material includes RGS(H)6-BAFF Q134 homomers and heteromers with FLAG-APRIL. This eluted material is applied to an M1 or M2 anti-FLAG Ab affinity column (Sigma, St. Louis, Mo.). A buffer exchange to 150 $\mathrm{mM} \mathrm{NaCl}-50 \mathrm{mM}$ Tris pH 7.0 is performed, and the buffer
adjusted to 2 mM CaCl 2 if using the M 1 column (this Ab requires Ca for binding). The column is washed with 150 $\mathrm{mM} \mathrm{NaCl}-50 \mathrm{mM}$ Tris pH 7.0 (with 2 mM CaCl 2 for M1). The M1 column is eluted by incubating the column with 150 $\mathrm{mM} \mathrm{NaCl}-50 \mathrm{mM}$ Tris $\mathrm{pH} 7.0-2 \mathrm{mM}$ EDTA for 30 minutes, followed by aliquots of $150 \mathrm{mM} \mathrm{NaCl}-50 \mathrm{mM}$ Tris pH7.0-2 mM EDTA, 10 min incubations, 6 times. Alternatively, both the M1 and M2 columns can be eluted by competition with FLAG peptide by allowing the column to drain completely and eluting 5 times with one column volume each of to $150 \mathrm{mM} \mathrm{NaCl}-50 \mathrm{mM}$ Tris pH 7.0 containing $100 \mathrm{vg} / \mathrm{ml}$ FLAG peptide. Eluted material will contain only native FLAG-APRIL:: RGS(H)6-BAFF Q134 heteromers.
[0081] Alternatively, cell lines or cells transfected, as above, with plasmids encoding full length or untagged soluble APRIL and BAFF constructs could be used as a source to isolate APBF complexes with anti-peptide antibodies raised against regions of the extracellular domains for APRIL and BAFF. These antibodies could be coupled to a resin by conventional means. Conditioned media or cell extracts of such cells could be run first over a column containing the coupled antibody (s) against one of the ligands, for example anti-BAFF antibodiy (s). In this instance, only homomers and heteromers containing a BAFF subunit bind to the anti-BAFF column; homomeric APRIL molecules flow through. After washing the column in 150 $\mathrm{mM} \mathrm{NaCl}-50 \mathrm{mM}$ Tris pH 7.0 , the bound molecules could be eluted off by competition with the same BAFF peptide(s) used to raise the anti-BAFF antibody(s) by allowing the column to drain completely and eluting 5 times with one column volume each of to $150 \mathrm{mM} \mathrm{NaCl}-50 \mathrm{mM}$ Tris pH 7.0 containing $100 \mu \mathrm{~g} / \mathrm{ml}$ or greater the peptide(s). This eluate could be dialyzed to remove the peptide(s) and then similarly run over a column containing anti-peptide antibody(s) raised against the other ligand, in this example APRIL. ). In this instance, homomeric BAFF molecules would not bind to the column and flow through. Only the remaining APBF heteromers bind to the anti-BAFF column. These APBF heteromers could be similarly eluted by competition with the same APRIL peptide(s) used to generate the anti-APRIL antibody(s).

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| $\begin{gathered} \text { Ile } G \ln G \ln T h \\ 35 \end{gathered}$ |  |  |  |  |  |  |
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1-16. (canceled)
17. An antibody specific for APBF.

