Fas ligand is not only expressed in immune privileged human organs but is also coexpressed with Fas in various epithelial tissues

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Abstract

Aims—To confirm the recent data obtained in mice, showing that the Fas ligand (FasL) is involved in the phenomenon of “immune privilege” (the apparent defect of the immune system in specific anatomical sites) and to extend this finding to humans.

Methods—The expression of FasL was analysed in a panel of histologically normal human tissues by reverse transcriptase polymerase chain reaction and Western blotting. The tissues sampled were brain, breast, bone marrow, oesophagus, kidney, liver, lung, lymph node, ovary, pancreas, pituitary gland, prostate, spleen, stomach (antrum and fundus), striated muscle, testis, thyroid, and uterus. These were obtained from patients with various neoplastic and non-neoplastic disorders; placental tissue was obtained after normal obstetric delivery, and spontaneous or voluntary abortion.

Results—Strong FasL expression was detected in testis and placenta. FasL expression was also detectable, although it was seen to a lesser extent, in oesophagus, prostate, lung, and uterus, which also coexpressed variable amounts of Fas mRNA or protein or both. The other organs tested for FasL expression were all negative.

Conclusions—FasL in humans is expressed predominantly in immune “sanctuaries” such as testis and placenta, suggesting that, similar to mice, this expression may contribute to the immune privileged status of these organs, by preventing dangerous inflammatory responses. The coexpression of FasL and Fas in particular epithelia suggests that the physiological cell turnover of some tissues may be regulated by the Fas-FasL apoptotic pathway.

Keywords: Fas/Fas ligand; Western blot; reverse transcriptase polymerase chain reaction; immune privilege

Fas/CD95 and its ligand (FasL) are members of two superfamilies of complementary receptors and ligands, respectively, that play important roles in immune regulation. Fas, also called APO-1, is a 45 kDa type I membrane protein belonging to the tumour necrosis factor receptor (TNFR) superfamily that includes the p55 and p75 TNFRs, the low affinity nerve growth factor receptor, CD27, CD30, CD40, OX40, and 4-1BB. Among these receptors, Fas and the p55 TNFR share a 70 amino acid intracellular “death domain” that transduces signals for cell death. Fas mediates apoptosis when cross-linked with agonistic anti-Fas antibodies. FasL is a 40 kDa type II membrane protein expressed as a membrane bound form that is proteolytically processed into a soluble cytokine retaining biological activity. There is increasing evidence from lpr and gld mice models that CD95/CD95 ligand mediated apoptosis is a crucial regulatory process which limits immune response and provides a safeguard against emergence of autoreactive lymphocytes. lpr mutant mice harbour either an early transposable element or a mutation in the Fas gene, which either reduces severely the expression of a full size Fas transcript, or abolishes its ability to transduce an apoptotic signal. gld mice harbour a point mutation in the extracellular domain of FasL; this mutation abolishes the binding between FasL and its receptor. Mice carrying a homozygous mutation for either lpr or gld develop similar phenotypes, characterised by massive lymphadenopathy, splenomegaly, B cell activation, and hypergammaglobulinemia. Lymphadenopathy in lpr and gld mice results from the expansion of a non-neoplastic subset of chronically activated cells that are TCR+, CD8+, CD4+, and CD8 (dubiously negative T cells). The major immune defect in lpr and gld mice appears to be the defective TCR induced death of mature T cells, which results in a failure in peripheral, rather than thymic, T cell deletion.

The human counterpart of lpr mice is the recently described auto-immune lymphoproliferative syndrome, caused by dominant interfering Fas mutations. Affected children have a defect in apoptosis of phenotypically normal mature T cells, and present with accumulation of doubly negative T cells in spleen and lymph nodes.

Because of its capacity to induce apoptosis of T cells, FasL has been thought to be involved in the phenomenon of “immune privilege”. Immunologically privileged sites are regions of the body, such as the eye and brain, where the immune system does not appear to function. Testis also acts as a sanctuary in which infectious organisms, tumour cells, or allogeneic grafts neither elicit destructive nor protective immunity. The concept of immune privilege
can be extended to the observed lack of immune response of mothers against embryos during pregnancy, which remains an enigma. FasL was shown recently to be a key effector of immune privilege in the mouse; thus, it is expressed abundantly in testes and eyes, where it can induce the apoptosis of migrating immune and inflammatory cells.

In order to confirm and extend these data to humans, we analysed the expression of FasL in human tissues by combining reverse transcriptase polymerase chain reaction (RT-PCR) amplification and Western blot analysis.

**Methods**

**TISSUE SAMPLING**

Histologically normal specimens were surgically removed from various organs—brain, breast, bone marrow, oesophagus, kidney, liver, lung, lymph node, ovary, pancreas, pituitary gland, prostate, spleen, stomach (antrum and fundus), striated muscle, testis, thyroid, and uterus. Some of the patients presented with a benign tumour located in the same organ as the tissue sampled, but at a distance from the normal tissue sample (breast, lung, pituitary gland, prostate, thyroid). Other patients suffered from non-neoplastic diseases (kidney, lymph node, ovary, spleen, stomach, uterus). Bone marrow, liver, pancreas, and striated muscle were taken from patients with a malignant tumour in another organ. Only the brain and testis samples corresponded to histologically normal areas located at a distance from a malignant tumour in the same organ. The placental specimens were obtained after normal obstetric delivery, and spontaneous or voluntary abortion. RNA could be obtained from at least one sample of each normal tissue. The remaining tissues were processed for immunohistochemistry or Western blot analysis.

**CELLS AND ACTIVATION CONDITIONS**

The Jurkat T-cell line and human peripheral blood lymphocytes (PBL) from a healthy donor were used as positive controls for FasL mRNA expression. Jurkat cells were stimulated with phytohaemagglutinin (PHA, Sigma, Dorset, UK) (10 µg/ml) for one or two hours. PBL were activated with 2.5 µg/ml of PHA and 30 U/ml of recombinant human interleukin-2 (IL-2) (Peprotech, Chiron, France), then grown for four days in medium containing 20 ng/ml IL-2. Cells were restimulated with 1 ng/ml of phorbol myristic acetate (PMA, Sigma) and 500 ng/ml of concanavalin A (ConA, Sigma) for 24 hours as described previously prior to poly(A)+ RNA preparation.

**RT-PCR ANALYSIS OF FASL AND FAS EXPRESSION**

Total RNAs were prepared from frozen tissues by lysis in guanidium isothiocyanate. Reverse transcription was performed as described previously, using 1 µg of poly(A)+ RNA from control cells, and 1 µg of total RNA from tissues. One quarter of the cDNA preparation was used for FasL amplification using a 20-mer sense oligonucleotide (5'-CTACAGGAC TGAG AAGAAGT-3') upstream of the initiation codon, and a 22-mer antisense oligonucleotide (5'-CAACATTCTCGGTGCTGTA AC-3') downstream of the termination codon. Oligonucleotide primers for Fas (sense primer, 5'-GTAAGG TAATGATTTTGAGTGCAG-3' (nucleotide positions 1718-1744), and antisense primer, 5'-CATT TTGGGGGGTGG-3' (positions 2175-2191)) were in the non-coding region; they both amplify mature mRNAs. Integrity of RNA samples had been proved by means of electrophoresis of an aliquot on denaturing agarose/formaldehyde gel, and was further checked by performing control amplification for human β actin (sense primer, 5'-TACCATCTGGCAGT GATGGACT-3', and antisense primer 5'-TCCATTGTCAGT TTGTCGGCATA-3'). The cDNA was mixed with 50 µl of a PCR mixture containing the reaction buffer and 2.5 U of Taq polymerase (Perkin Elmer Cetus, New Jersey, USA). The PCR cycle profile was as follows: denaturation (94°C for one minute (two minutes for the first cycle)), annealing (55°C for two minutes), and extension (72°C for three minutes (10 minutes for the last cycle)); 30 cycles of amplification were used. PCR products were visualised in ethidium bromide stained 1.5% agarose gels.

**SOUTHERN BLOT ANALYSIS**

PCR products separated by means of gel electrophoresis were transferred to Hybond N+ membranes (Amersham, Buckinghamshire, UK) as recommended by the manufacturer. Membranes were prehybridised for two hours at 45°C in 5× SSC containing 0.1% laurylsarcosin, 0.02% SDS and 0.5% blocking reagent (Boehringer Mannheim, France). They were hybridised overnight at 45°C in the same buffer containing a digoxigenin labelled internal oligonucleotide: an 18-mer (CAATA AGG GCCACCCCATG, spanning the 3' end of exon 2 and the 5' end of exon 3) for FasL, and a 24-mer (5'-ACATGTGATGAACCCCATGTTTGA-3' (nucleotide positions 2118-2142)) for Fas, respectively. Oligonucleotide probes were labelled with a 3' tailing kit (Boehringer Mannheim) according to the supplier's recommendations. After hybridisation, filters were washed twice for 10 minutes at room temperature in 2× SSC containing 0.1% SDS, and twice for 10 minutes at 55°C in 0.1× SSC containing 0.1% SDS, then visualised using a chemiluminescence detection kit (Boehringer Mannheim).

**WESTERN BLOT ANALYSIS**

Frozen tissues, crushed in liquid nitrogen, were thrown in lysis buffer (1% SDS, 1 mM sodium vanadate, 10 mM Tris HCl, pH 7.4) and heated in a microwave oven for 10–15 seconds. Proteins from lysates were separated by 7.5% SDS-PAGE and transferred to Immobilon-P membranes (Millipore, Massachusetts, USA) as recommended by the manufacturer. Detection of FasL was performed with a mouse IgG3 monoclonal antibody (MoAb) (Transduction Laboratories, Kentucky, USA), raised against an 18 kDa polypeptide corresponding to amino...
Fas ligand in human tissues

Results

FASL AND FAS PROTEIN EXPRESSION

The Fas-FasL apoptotic pathway has been shown to play a key role in the immune system by triggering the activation induced suicide of T cells. Resting T cells normally express low amounts of Fas; this expression increases significantly within hours of engagement of their antigen receptor. Several days after the initial activation, this pathway becomes effective, and T cells undergo apoptosis via cross-linking of the FasL concomitantly up-regulated in activated T cells. Fas is also expressed in non-lymphoid tissues as diverse as liver, heart, ovary, and lung, where its physio-

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Immunodetection of Fas was performed in frozen sections as described previously using the MoAb CH-11 (Immunotech, Marseille, France). Reactive lymph nodes were used as control.

Figure 1. Expression of the Fas ligand in human tissues. Upper panel: RT-PCR detection of FasL transcripts in normal human tissues: (1) negative control, (2) positive control (PHA activated Jurkat cells), (3) placenta, (4 and 16) DNA molecular weight markers, (5) lung, (6) heart, (7) testis, (8) liver, (9) breast (sample a), (10) kidney, (11) oesophagus, (12) stomach (antrum), (13) striated muscle, (14) thyroid gland, (15) breast (sample b), (17) ovary, (18) spleen, (19) uterus, (20) cerebellum, (21) pituitary gland, (22) brain (frontal lobe), (23) total bone marrow, (24) lymph node. The amplified FasL product is an 840 base pair band (arrow), detected in the positive control (lane 2) and in placenta (lane 3), lung (lane 5), and testis (lane 7). Middle panel: Ethidium bromide stained agarose gel containing aliquots of cDNA amplified with the β actin primers and arranged in the same order as the upper panel. The β actin PCR product is a 460 base pair band. Lower panel: Same as upper panel after Southern blotting and hybridisation with a FasL specific digoxigenin-labelled probe, which revealed a weak positivity in oesophagus (lane 11) and uterus (lane 19). Other tissues were negative.

acids 116–277 of human FasL. Immunoreactive material was visualised by enhanced chemiluminescence (Amersham). Stimulated Jurkat cells served as positive controls. Negative controls were an ovalbumin/bovine serum albumin solution and unstimulated Jurkat cells. To confirm the specificity of the PCR data, a representative panel of negative and positive samples from neoplastic and normal tissues were analysed.

Discussion

Western blot analysis of FasL expression correlated with the PCR data. The expected 37 kDa band was detected in the placenta, whereas PCR negative samples such as lymph node were negative (figure 2). The Fas protein was detected by immunohistochemistry in lung, prostate, and ovary, whereas three placenta samples were negative (data not shown).

The rela-
Table 1  Expression of Fas and Fas ligand in human tissues

<table>
<thead>
<tr>
<th>Tissue</th>
<th>RT-PCR positivity*</th>
<th>Fas ligand</th>
<th>Fas</th>
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<td>1/1 (+)</td>
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<tr>
<td>Prostate</td>
<td>1/1 (++)</td>
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*Number of positive samples /number of independent samples analysed. The intensity of PCR bands is given in parentheses: +++, strong band on ethidium bromide stained gel; +++, weak band on ethidium bromide stained gel; +, band detected after Southern blot hybridisation; +/-, weak band detected after Southern blot hybridisation. ND, not done for technical reasons.

Figure 2  Western blot analysis of Fas ligand expression in a panel of normal tissues. The 37 KDa FasL protein was detected in PCR positive samples such as placenta and stimulated Jurkat cells. A faint signal was also detected in a PCR negative spleen sample, which may reflect a low level of FasL protein expression, more easily detectable than the corresponding transcript.

The logical function is unclear, as these tissues show no developmental abnormalities in lpr and gld mice. It is also expressed in tumours, especially lymphomas. In contrast to its receptor, FasL expression appears tightly regulated, except for a pathological situation such as seen in gld mice that display a constitutive expression of a mutated FasL. In mice, FasL is expressed in lymph nodes and spleen but it is usually undetectable by Northern blot in non-lymphoid tissues such as brain, heart, liver, kidney, ovary, and skin. Some investigators have reported variable amounts of FasL expression in the testis and eye, and very low levels in small intestine and uterus. These results suggest that the physiological role of FasL extends beyond that exerted on the immune response, and that it may be the basis of the particular immune status of the eye and testis known as “immune privilege”.

To determine whether FasL expression could represent the basis of immune privilege in humans, we studied FasL mRNA expression and localisation by using RT-PCR to analyse various organs, including those known to act as immune sanctuaries. We found that FasL expression was confined to a few specific sites, in good correlation with earlier findings in mice.

Expression of FasL mRNA has been reported in rodent but not in human testis. Testis has long been known to be a remarkably immune privileged site; rat testis, when placed in the abdominal cavity, allows the successful transplantation of both allografts and xenografts. The expression of FasL in human testis reported here suggests that similar mechanism(s) could underlie immune privilege in mice and humans. Testicular FasL might induce apoptosis of T cells activated in response to tumour antigens, thereby favouring the growth of neoplastic cells. Such a hypothesis may help to explain particular clinical observations, like the aggressive behaviour of primary testicular lymphomas which frequently relapse in the contralateral testis. It may be that one of the purposes of immune privilege in the testis could be to arrest the spread of dangerous inflammatory responses, potentially able to destroy germinal cells and thus threaten the fate of the species.

The demonstration of FasL expression in the placenta is of special interest, as the mechanisms preventing rejection of the embryo by the maternal immune system are subject to speculation. Preliminary immunohistochemical analysis of FasL expression seemed to localise the staining in the fetal trophoblast covering the placental villi (Xerri, unpublished observation). The trophoblast is situated at a strategic interface between maternal and fetal circulations; the apoptotic signals produced by trophoblastic FasL could regulate the immune traffic and maintain maternal tolerance by preventing the activated lymphocytes from entering the fetal cellular compartment. Another possibility may be that FasL regulates the growth of placenta itself via an autocrine-paracrine loop involving the Fas–FasL apoptotic pathway. This possibility, however, appears unlikely as Fas expression was not detected in the FasL positive placenta specimen. This negative result does not exclude a transient upregulation of Fas during critical stages of placental maturation.

It has long been thought that immune privilege in specific human organs was passively maintained by particular anatomical barriers which prevent inflammatory cells or antigens from entering the privileged sites. Although this hypothesis appears unlikely regarding tissues expressing FasL, it seems relevant for the blood–brain barrier, because our results showed a lack of FasL expression in the brain, as previously demonstrated in mice brain.
Fas ligand in human tissues