Sphingosine-1-phosphate lyase expression in embryonic and adult murine tissues

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Running title: SPL expression in embryonic and adult murine tissues
Abstract

Sphingosine-1-phosphate (S1P) is a bioactive sphingolipid involved in immunity, inflammation, angiogenesis and cancer. S1P lyase (SPL) is the essential enzyme responsible for S1P degradation. SPL augments apoptosis and is downregulated in cancer. SPL generates a S1P chemical gradient that promotes lymphocyte trafficking and as such is being targeted to treat autoimmune diseases. Despite growing interest in SPL as a disease marker, anti-oncogene and pharmacological target, no comprehensive characterization of SPL expression in mammalian tissues has been reported. We investigated SPL expression in developing and adult mouse tissues by generating and characterizing a β-galactosidase-SPL reporter mouse combined with immunohistochemistry, immunoblotting and enzyme assays. SPL was expressed in thymic and splenic stromal cells, splenocytes, Peyer’s Patches, colonic lymphoid aggregates, circulating T and B lymphocytes, granulocytes and monocytes, with lowest expression in thymocytes. SPL was highly expressed within the CNS, including arachnoid lining cells, spinal cord, choroid plexus, trigeminal nerve ganglion and specific neurons of the olfactory bulb, cerebral cortex, midbrain, hindbrain and cerebellum. Expression was detected in brown adipose tissue, female gonads, adrenal cortex, bladder epithelium, Harderian and preputial glands and hair follicles. This unique expression pattern suggests SPL has many undiscovered physiological functions apart from its role in immunity.

Supplementary key words  sphingosine phosphate lyase • sphingosine-1-phosphate • colon cancer • development • sphingolipid • signal transduction
INTRODUCTION

S1P is the final metabolic product of sphingolipid degradation. It circulates in the blood and lymph bound to lipoproteins and functions as a ligand for a family of five differentially expressed G protein coupled receptors, the S1PRs (1, 2). Acting through these receptors and downstream signaling events mediated by them, S1P influences cell migration, survival and stress responses (3). S1PR signaling is critical for development, as demonstrated by the embryonic lethal phenotype of S1PR1 knockout mice, due to severe hemorrhage associated with impaired vascular maturation (4). S1PR2 knockout mice are deaf, due to defects in the stria vascularis (5). Additional studies have demonstrated the importance of S1P transport and signaling in cardiac and brain development, reproduction and embryonic stem cell survival (6-17). S1P acts through its cognate receptors to control T, B, natural killer and hematopoietic stem cell trafficking as these cells emerge from bone marrow and thymus and transit peripheral lymphoid organs (18, 19). S1P signaling contributes to the regulation and constitutive activation of key transcription factors that play a central role in inflammation and cancer, including NFκB and STAT3 (20, 21). S1P is also a mediator of cardioprotection in response to ischemia and can reduce developmental cell death as well as apoptosis, tissue injury and infertility in response to ionizing radiation (22-24). Although many effects of S1P signaling are mediated through activation of its receptors, some of its actions, including activation of NFκB signaling and regulation of histone deacetylases appear to be mediated through receptor-independent, intracellular functions of S1P (21, 25).

S1P is generated via phosphorylation of the sphingoid base sphingosine by sphingosine kinases, SphK1 and SphK2 (26). These two enzymes exhibit different biochemical properties,
are expressed in different tissues, are located in different subcellular compartments, and manifest
distinct biological functions. S1P generated through the actions of sphingosine kinases can be
dephosphorylated by S1P phosphatases, SPP1 and SPP2 or through reactions catalyzed by
nonspecific lipid phosphatases, the LPPs (27-29). However, each of these reactions results in
regeneration of sphingo sine, which is then immediately available to be rephosphorylated. In
contrast to the phosphatases, SPL is responsible for catalyzing the irreversible degradation of
S1P to hexadecenal and ethanolamine phosphate by cleaving the C2-3 carbon bond in S1P in
what constitutes the final step of sphingolipid metabolism (30). As such, SPL functions as the
major regulator of S1P levels, and loss of its activity or expression results in significant
accumulation of S1P and other sphingolipid intermediates within cells and tissues (31). The
enzyme is conserved from yeast to man, and mutants in various model organisms have revealed
phenotypes including abnormalities of growth regulation and carbon metabolism in
Saccharomyces cerevisiae, developmental and migration defects in Dictyostelium discoideum,
reproductive defects in Caenorhabditis elegans, and degeneration of reproductive organs and
muscles in Drosophila melanogaster (32-36).

Mammalian SPL is encoded by the Sgpl1 gene, found on chromosome 10 of mouse and
human genomes (37, 38). Mice homozygous for a Sgpl1 null allele demonstrate negligible SPL
expression and enzyme activity in tissues, as well as high circulating and tissue levels of S1P,
indicating that Sgpl1 is the major or only gene responsible for SPL activity (31, 39). SPL null
mutants do not live past one month of age postnatally and exhibit a range of phenotypes
including skeletal and hematological abnormalities, elevated cytokines and pro-inflammatory
responses, impaired neutrophil and lymphocyte trafficking, elevated serum lipids, increased lipid
storage in the liver, and deficient adipose stores (39, 40). These findings collectively implicate SPL in development, lipid homeostasis and the regulation of innate and adaptive immunity.

Recent studies have implicated SPL as a potential immunomodulatory target in the context of autoimmune disease (41-43). LX2931, now in clinical trials for the treatment of rheumatoid arthritis, acts by inhibiting SPL, disrupting the S1P chemical gradient required for mature T cell egress from the thymus and peripheral lymphoid organs and thereby preventing immune-mediated tissue damage (43). Studies have also implicated SPL as a potential target for cardioprotection, radioprotection and recruitment of skeletal muscle stem cells for regeneration of injured muscle (44-46). SPL is downregulated in colon cancer and other malignancies (47-49). Coupled with the findings that overexpression of SPL promotes apoptosis in response to DNA damage, whereas knockdown of SPL promotes cell transformation, suggest that SPL silencing contributes to carcinogenesis and that SPL may serve an anti-oncogenic function by promoting the turnover of DNA-damaged cells (45, 50, 51). Although most of the functions and phenotypes associated with SPL and its disruption are attributed to its regulation of S1P, there is growing evidence that the products of the reaction catalyzed by SPL are important regulators of apoptosis and may have additional biological functions (52, 53).

Despite growing interest in SPL as a disease marker, potential anti-oncogene and pharmacological target, no comprehensive characterization of its expression pattern in mammalian tissues has been reported. In the current study, we explored SPL expression in embryonic and adult murine tissues by generating and characterizing a β-galactosidase (β-Gal) SPL reporter mouse, combined with immunohistochemistry (IHC), immunoblotting, and enzyme activity assays. In addition, we have explored SPL expression in the developing mouse embryo.
These studies provide a more complete characterization of SPL expression than has been previously reported.
MATERIALS AND METHODS

Generation of SPL β-Gal reporter mice and other animals used

The SPL reporter mouse line was generated during the production of a conditional SPL knockout line. To produce both these lines, we generated a sequence-replacement targeting vector designed to insert a gene-trapping cassette containing a lacZ reporter within intron 9-10 of Sgpl1. The gene-trapping vector, which was flanked by Frt sites, consisted of a splice acceptor site (flanked by a loxP and a lox71 site) followed by a promoter-less βgeo cassette. This is the same gene-trapping cassette used by the KOMP: Knock-Out Mouse Project (54). We utilized the gene-trapping cassette designed to yield an in-frame fusion with exon 9 of Sgpl1. The gene-targeting vector was designed to insert a single loxP site into intron 12-13. The integrity of the completed gene-targeting vector was verified by restriction endonuclease digestion and by DNA sequencing.

The gene-targeting vector was electroporated into strain 129/OlaHsd embryonic stem (ES) cells, which were then subjected to selection in G418. G418-resistant ES cell clones that produced a fusion transcript extending from exon 9 of Sgpl1 to βgeo (indicating the expected recombination event had occurred) were identified by 5′ rapid amplification of 5′ complementary DNA ends (RACE). PCR was used to identify clones that also retained the distant loxP site. One such clone was then injected into C57BL/6 blastocysts to generate chimeric mice. A total of 10 high-percentage male chimeras were obtained, and those mice transmitted the targeted mutation to their offspring. The reporter mice were backcrossed six times and subsequently maintained on the C57BL/6 background.
Wild type C57BL/6 mice were obtained from Taconic (Oxnard, CA). C57BL/6 mice carrying the Sgpl1 null allele described previously (39) were obtained from Philippe Soriano (Mt. Sinai School of Medicine, NYC). Animals were maintained in the Children’s Hospital Oakland Research Institute (CHORI) AAALAC Accredited Animal Facility. All experiments were conducted in accordance with CHORI Institutional Animal Care and Use Committee approved protocols.

Whole mount survey of SPL β-Gal reporter expression in adult murine tissues

Organ dissection, fixation and staining for β-Gal activity in adult male and female reporter mice was conducted essentially as described (55).

Frozen section tissue staining for β-Gal activity

Tissue sections were fixed and stained for β-Gal activity essentially as described (56). Briefly, tissue samples from SPL reporter and wild type mice were collected and fixed in LacZ fixative solution containing 0.2% glutaraldehyde, 5mM EGTA (pH 7.3), 100 mM MgCl₂, in 0.1 M NaPO₄ (pH 7.3) for 4 hours at room temperature with a solution change at 2 hours. The samples were transferred into 15% sucrose in PBS for 4 hours at room temperature and then into 30% sucrose in PBS overnight at 4°C. Tissues were embedded in OCT and 10 micron frozen sections were cut. Air dried sections were washed 3 times in PBS, transferred to X-Gal staining solution: 1mg/ml of 5-bromo-4-chloro-3-indolyl β-D-galactosidase, 2 mM MgCl₂, 5 mM potassium ferrocyanide, 5 mM potassium ferricyanide, 0.01% sodium deoxycholate, and 0.02% Nonidet-P40 overnight at 37°C. The following day samples were washed in distilled water, counterstained with Nuclear Fast Red, and coverslipped with aqueous mounting media. For each
tissue examined, corresponding tissue from a wild type mouse lacking the SPL reporter was used as a negative control.

**IHC for SPL expression**

Formalin-fixed and paraffin embedded murine tissues were deparaffinized and incubated for 30 min in 3% hydrogen peroxide/methanol to quench endogenous peroxidases. Sections were rinsed in PBS and immunostained with anti-(murine) SPL antisera at 1:200 dilution in 0.5% PBS/Ova Albumin at 37°C for 1 h after antigen retrieval with citrate buffer, pH 6.0 in a small autoclave set for 125°C for 2 min; slides were cooled for 1 h at RT before adding secondary antibody. Secondary antibody was biotinylated anti-rabbit (Vector laboratories) diluted 1:1000 in 0.5% PBS/Ova Albumin and incubated for 30 min at RT. Sections were incubated with Elite ABC kit (Vector Laboratories) for 30 min, rinsed in PBS, and detection was performed with DAB (Vector Laboratories) for 2 min, and counterstained in Hematoxylin. Tissues from a SPL knockout mouse were used as a negative control for IHC studies.

**Flow cytometry**

Blood cells and splenocytes were pooled and labeled with fluorescent antibodies including the B cell marker B220-APC, T cell marker CD3-FITC, polymorphonuclear cell marker Gr-1 PerCP Cy5.5 and monocyte marker Cd11b-PE, all obtained from eBiosciences, Inc. (San Diego, CA). Cell separation was performed on a Becton Dickinson FACSArria I cell sorter using FACSDiva software. Granulocytes were identified as CD11b+ and Gr-1+, whereas monocytes were identified as Cd11b+ and Gr1-. Cells were homogenized and used for immunoblotting.
**Immunoblotting**

Immunoblotting was performed using a rabbit polyclonal antibody generated against the C-terminal peptide of murine SPL of the sequence: C-VTQGNQMNGSPKPR. Murine SPL antibody was used at 1:1000 dilution. Actin antibody (Sigma) was used as a loading control at 1:20,000 dilution. Immunoblotting was conducted essentially as described previously (47).

**SPL enzyme activity assays**

SPL enzyme activity was performed using tritium radiolabeled dihydrosphingosine-1-phosphate substrate as described previously (57).

**Relative levels of tissue SPL expression by IHC and frozen section β-Gal staining**

In order to provide a rough estimate of total organ SPL expression levels that would allow for comparison with SPL activity levels in different organs, we utilized a scoring system for β-Gal or SPL IHC. This scoring system reflected both the staining intensity level as well as the distribution of stain (percentage of cross-sectional area that expresses SPL at any given level) within an organ. A full cross-section of the organ was assessed, thereby providing information incorporating the signal from many substructures. This represents a modified combined intensity and distribution score (58). In other words, the score for an organ reflects total expression levels that can be affected by high, intense expression in specific substructures as well as low but detectible expression (as compared to the appropriate negative control tissue) in large areas of tissue. The tissue with the most robust cell staining, small intestinal villous tips, was both uniformly and strongly positive, and thus assigned a value of 1.0 (i.e., 100% value). All other tissues were then scored relative to this reference tissue. For example, the thymus in total would
give a low score due to the low to undetectable staining in immature lymphocytes. However, because the thymic epithelium was uniformly stained with moderate to high intensity, the score for thymic tissue was very high (0.95). In contrast, a minority of heart muscle cells exhibited consistently weak staining intensity, and other heart muscle cells showed no stain, resulting in a score of 0.05.

**Embryo isolation**

C57BL/6 mice heterozygous for the *Sgpl1* null allele were mated in order to generate litters containing some SPL knockout embryos and some embryos wild type for SPL. Embryos were removed from the uterus and yolk sac of pregnant mice at various days of gestation as described (59). Genotyping was performed to identify knockout embryos that were employed as negative controls for IHC staining. Representative parasagittal sections of E17 embryos were stained for SPL IHC as described below.
RESULTS

Generation of β-Gal SPL reporter mice

Global disruption of murine SPL expression results in 100% lethality in the weaning period (31, 39). Therefore, in order to investigate tissue-specific SPL functions in the adult mouse, we employed a combination of recombineering and Cre-lox technologies to generate an SPL conditional knockout mouse model. To facilitate the analysis of SPL expression in different murine tissues, the targeting vector was designed to introduce a bacterial LacZ gene 5’ to exon 9, thereby creating a β-Gal reporter under control of the Sgpl1 gene regulatory region and resulting in SPL truncation (SPL reporter mice). Mice heterozygous for the SPL allele containing the reporter are born at expected mendelian frequency, exhibit no developmental or other phenotypes, and have normal reproductive function and lifespan, similar to mice heterozygous for a Sgpl1 null allele. These reporter mice were utilized to characterize SPL expression as described below. The targeting cassette that harbored the reporter gene was subsequently deleted in order to generate a Sgpl1 allele in which exons 9-11 containing the enzyme cofactor-binding site are “floxed” for the establishment of tissue-specific SPL knockout mice lines described elsewhere (Ashok Pandurangan, Padmavathi Bandhuvula, Ashok Kumar, Meng Zhang, Yuko Yoshinaga, Mikhail Nefedov, Alexander D. Borowsky, Pieter de Young, Loren G. Fong, Stephen G. Young and Julie D. Saba., manuscript in preparation).

Whole mount survey of β-Gal stained adult SPL reporter mouse organs

In order to undertake a comprehensive survey of SPL expression in the adult mouse, we performed whole mount analysis of β-Gal expression in organs dissected from male and female
SPL reporter mice. SPL exhibits a broad range of expression levels in adult tissues. Little or no expression is detected in the whole mount analysis of tongue, spleen, pancreas, quadriceps muscle, sciatic nerve, heart atrium and ventricle, lung, aorta, diaphragm, mammary gland, pituitary gland, lacrimal gland, mesenteric adipose tissue, penis, prostate, vesicular gland, trachea, esophagus and duodenum (data not shown). Staining in some tissues such as the salivary gland appears to be non-specific, and faint expression is found in the thyroid gland (data not shown). In contrast, significant expression is observed in brain, spinal cord, trigeminal nerve ganglion, thymus, kidney, bladder, skin, preputial gland, Harderian gland, ribcage, liver, stomach, jejunum, ileum, cecum, colon, brown adipose tissue, adrenal cortex, and male and female reproductive organs (Figure 1A-X). In some tissues, including thymus, preputial gland, Harderian gland, jejunum and ileum, β-Gal expression is strong and uniform. In other cases, staining reveals specific patterns of expression, such as in the cortex, medulla and papilla of the kidney, discrete glomeruli of the olfactory bulb, specific regions of the hippocampus, midbrain, hindbrain and granular layers of cerebellum, spinal cord gray matter, costal cartilage of the ribcage, fundus of the stomach, medulla of the adrenal gland, hair follicles of the skin and epithelium of the bladder.

**Histological analysis of SPL reporter mouse tissues by β-Gal staining**

To gain more detailed analysis of some organs that stained strongly in whole mount survey, selected tissues were fixed and stained for β-Gal. As shown in Fig. 2A, brown fat exhibits strong SPL expression. In the skin, the sebaceous glands are strongly positive (Fig. 2B), and weak staining is detected in the squamous epithelium (Fig. 2C). The Harderian gland of the
eye stains strongly. The tracheal epithelium, spleen and glandular stomach all show some SPL expression (Fig. 2D).

**Survey of specific SPL adult tissues of interest by IHC**

We next evaluated SPL expression by IHC in a range of tissues, including tissues both exhibiting and lacking SPL expression as determined in the whole mount survey. As shown in Fig. 3A, bladder transitional epithelium shows strong but patchy expression. Skeletal muscle is weakly positive, and in the panel shown in Fig. 3B, there is a neurovascular bundle with similar weak positive staining in the nerve fibers. The bone marrow compartment shows specific immunoreactivity in the osteoclasts as well as rare cells with relatively abundant cytoplasm and immature nuclei located in the marrow itself, possibly representing marrow stromal cells (Fig. 3C and F). However, the majority of the hematopoietic cells are negative. The ovarian primordial oocytes adjacent to the primary follicle are positive for SPL, whereas the oocyte in the primary follicle is negative (Fig. 3D). Colon tissues show faint luminal membrane positivity in the epithelial cells, as well as occasional cells in the gastrointestinal associated lymphoid tissue, possibly lymphoid stromal or dendritic cells (Fig. 3E). The adrenal x-zone cells show moderate expression levels (Fig. 3G). The kidney is weakly positive in some tubules (Fig. 3H). The spleen shows positive cells throughout the red pulp predominantly, with elongated cytoplasm likely representing splenic stromal supporting cells (Fig. 3I).

**SPL expression in gut-associated lymphoid tissues, lymph nodes and blood cells**

Mice homozygous for the *Sgpl1* null allele exhibit lymphopenia, retention of mature T cells in the thymus, and neutrophil trafficking defects. These findings suggest that SPL has
multiple roles in innate and adaptive immunity. In order to provide a more in-depth analysis of SPL expression in immune cells and tissues, we evaluated SPL expression by IHC in gut-associated lymphoid tissues including Peyer’s Patches of the jejunum (Fig. 4A) and colorectal lymphoid aggregates of the colon (Fig. 4B), as well as in peripheral lymph nodes (Fig. 4C). In addition, we used immunoblotting to compare the levels of SPL expression in peripheral blood leukocytes and the stromal and non-stromal fractions of thymus and spleen, as shown in Figs. 4D-E. As shown in Fig. 4A-C, SPL is expressed in each of the adult peripheral lymphoid organs. In the spleen, SPL is readily detected and expressed in both splenocyte and stromal cell fractions with relatively equal intensities (Fig. 4D). In contrast, the stromal cells of the thymus exhibit a high level of SPL expression, whereas expression in the thymocyte fraction is barely detectible (Fig. 4D). These findings are consistent with our results from histological analysis of the thymus using both IHC and β-Gal methodologies. SPL is also expressed in circulating blood cells including B and T lymphocytes and granulocytes, with lower but detectible expression in monocytes (Fig. 4E).

Survey of SPL IHC positive brain tissues

By histochemical analysis, the expression of SPL within each anatomical section of the brain appears highly specific, consistent with the distinct expression patterns observed in each brain compartment in the whole mount β-Gal survey. The Purkinje layer neurons of the cerebellum are moderate in intensity but are uniformly immunoreactive (Fig. 5A). Cells lining the choroid plexus are positive (Fig. 5B). Olfactory neurons at the periphery of the olfactory bulb are one of the strongest SPL expressing tissues (Fig. 5C). The expression in the olfactory epithelium is found in the sensory neurons. SPL is expressed in the cell body proper and also in
the axonal processes to the olfactory bulb. Brain stem neurons also demonstrate relatively strong SPL expression (Fig. 5D), comparable to the cerebellum and stronger than the cerebral cortical neurons, which are weakly to moderately positive. Meninges, particularly the arachnoid lining cells, are positive (Fig. 5E). Interestingly, some of the thalamic nuclei neurons are positive, while others are negative for immunoreactivity (Fig. 5F). Glial astrocytes (5F, most cells), oligodendroglia (5A top, and 5F smallest nuclei), granular cell layers of the cerebellum (5A, bottom), and the majority of the brain monocytes/microglia are negative for immunoreactivity (not shown). Cell types in the brain were identified by morphologic features on standard stains including the hematoxylin counter staining in the IHC studies shown.

Validation and comparison of SPL expression detected by β-Gal reporter and IHC

In order to establish if the SPL reporter is a reliable indicator of SPL expression, we compared histological results of β-Gal staining of (SPL reporter mouse) tissues known to express high levels of SPL with IHC staining of the corresponding tissues from a WT control mouse. For SPL IHC, we employed a murine SPL antibody generated against a peptide corresponding to the C-terminus of murine SPL. This antibody demonstrates a high degree of specificity, as shown by absence of staining in tissues of mice homozygous for the Sgpl1 null allele. It should be noted that SPL knockout mouse tissues are the appropriate control for IHC studies, whereas tissues from a wild type mouse lacking the SPL reporter insertion are the appropriate controls for β-Gal staining. As shown in Fig. 6A, thymic epithelium of a WT mouse stains strongly using the SPL IHC detection system. In contrast, no staining is observed in the thymus of a SPL null mouse (Fig. 6A, inset). As shown in Fig. 6B, SPL protein expression in the thymus determined by β-Gal staining shows corresponding patterns, with expression restricted to thymic epithelial cells of the
stroma and little or no staining in thymocytes, and no staining in wild type mouse tissues (Fig. 6B, inset). As shown in Fig. 6C, IHC of WT mouse jejunum shows strong staining in the differentiated enterocytes at the tip of the villi with less staining toward the base of the crypts, with no staining in SPL null mouse intestines (Fig. 6C inset). Similarly, Fig. 6D shows robust β-Gal staining in the SPL reporter mouse jejunum, whereas no staining is detected in the WT mouse jejunum (Fig. 6D, inset). Brain cortical neurons exhibit much lower SPL expression levels. However, positive cells are similarly identified by both SPL detection systems, but not in the corresponding negative control tissues (Fig. 6E and F). Overall, these results establish the specificity and sensitivity of both IHC and β-Gal reporter systems to detect SPL expression in tissues expressing relatively high and low levels of SPL.

Comparison of SPL expression among different tissues

To provide a comparison of SPL activity among different tissues of the adult mouse, we performed SPL enzyme activity assays in a range of organs from three individual mice (Fig. 7A). SPL activity was highest in the small intestine (jejunum), with lowest levels in heart and brain, and intermediate levels in colon and thymus. We observed consistent levels of SPL enzyme activity within the same tissues of different individual mice. In order to provide comparison between SPL activity and SPL expression levels as determined by our different histological detection systems, we employed a method that allows a rough estimate of total organ expression levels using each staining technique, as described in Materials and Methods and as published previously (58). This method reflects a combined intensity and distribution score. The score for each organ was normalized to the most intense and consistent area of expression, the intestinal villi (Fig. 7B-C), which was arbitrarily set at 1. Strong IHC staining was observed in the
intestines of adult SPL reporter mice (100%). Based on this reference tissue, uniform staining was also observed in the thymus (95%). In the stomach, 50% positive cells occurred. Much lower levels of SPL expression were found in the liver (30%), colon (20%), brain (15%), lung (10%), kidney (10%), and heart (5%). Relative expression of SPL, as indicated by β-Gal staining of tissues harvested from adult SPL reporter mice, is summarized in Fig. 7C. Based on strong positive staining in the intestines (considered 100% reference tissue), SPL was also robustly expressed in the thymus (95%) and spleen (90%). Staining in the stomach was 40%. Much lower levels of SPL expression were found in the liver (20%), brain (15%), colon (15%) lung (10%), kidney (10%), and heart (5%). There was good correlation between the enzyme activity, β-Gal and IHC results in all of the above tissues. In contrast, IHC staining in the spleen was low (25%) compared to β-Gal activity (90%) and restricted to scattered splenic dendritic cells. This suggests either a short lived/rapid turnover of the SPL protein compared to the stability of β-Gal, or alternatively a source of falsely positive β-Gal activity in the spleen, although this was not seen in controls supporting the former conclusion.

Expression of SPL during murine development

The expression of SPL was investigated in day 17 murine embryos (Fig. 8.I). As in the adult, the intestinal epithelium and thymus exhibit the strongest expression of all the tissues. Positive cells are also noted in the developing sensory ganglia, olfactory lobe, nasal epithelium, salivary gland, bladder epithelium, pituitary gland, brown fat, skin and hair.

Histological analysis of the tissues exhibiting the strongest and most specific staining are shown in Fig. 8.IIA-I. The mid-small intestine (jejenum) shows diffuse strong epithelial staining with additional SPL positive cells scattered in the basal layers. The proximal small intestine and
duodenum show weaker luminal epithelial staining, but still have scattered strong basal cells. The embryonic thymus is strongly positive, with a high density of the positive signal in thymic epithelial cells and a low signal in the lymphocytes. The upper respiratory mucosa and olfactory epithelium express high levels of SPL. The bladder transitional epithelium is strongly positive with much weaker expression in the smooth muscle. The trigeminal ganglion cells show scattered cells with strong expression. The developing bone at the epiphysis shows extensive staining in the osteoblasts. The developing exocrine pancreas is positive predominantly in ductal cells and more weakly in acinar cells. The pituitary gland shows scattered strongly positive cells in the adenohypophysis.
DISCUSSION

SPL is required for mammalian developmental and physiological functions, as shown by the congenital abnormalities, immunological and metabolic dysregulation, runting and short lifespan observed in SPL null mice (39). To gain more insight into possible functions of SPL in the developing and adult mouse, we have examined the spatial expression pattern of murine SPL using a combination of β-Gal reporter whole mount and histological stains, immunohistochemistry and enzyme activity assays. A high degree of correlation was found between the three detection systems, demonstrating that the results are robust and validating the SPL reporter as a reliable system for SPL expression. A limited number of tissues, including the pituitary gland and spleen, gave discrepant results between different detection methods. This may be explained by poor penetration of detection reagents in the whole mount assay or the expression of SPL in specific cell types deep within the tissues that are not evident in whole mount assay. Some differences were also observed in intensity of SPL expression as determined by IHC versus SPL reporter. While this could reflect technical conditions, such as the efficiency of antigen retrieval in different tissues with IHC detection, positional effects on the lacZ reporter or endogenous β-Gal activity, the evaluation of multiple animals (both control and reporter/knockout), suggest that the differences are not based on artifact. Instead, it seems likely that some tissues have higher levels of transcription as detected by the lacZ reporter, even without higher levels of protein as detected by IHC. This suggests that there is post-transcriptional regulation or more rapid protein degradation in some tissues. Overall, a combination of detection systems, biochemical assays and appropriate negative controls provides
the most reliable information regarding SPL expression and function in individual tissues of interest.

Our results in small intestines, which express the highest levels of SPL among all the organs evaluated, are consistent with previous reports (47, 48). SPL expression in intestinal and colonic epithelia likely represents its role in the metabolism of dietary sphingolipids (60). Downregulation of SPL in neoplastic intestinal tissues, combined with the established role of SPL in regulation of apoptosis, suggest that SPL may also facilitate the turnover of damaged intestinal epithelial cells.

Similarly, SPL is highly expressed in the thymus, corroborating previous reports (61, 62). SPL expression is restricted to the epithelial cells of the thymic stroma. It has been suggested that SPL expression in this location serves to maintain low thymic S1P levels, thereby generating the S1P gradient that facilitates mature lymphocyte egress, a concept that has not yet been established experimentally (42). SPL expression was not detected in thymic lymphocytes using β-Gal, IHC or immunoblotting methods, nor was it detected by IHC in blood cells of the bone marrow. These findings, in combination with the notable SPL expression in peripheral lymph nodes, gut-associated lymphoid tissues and circulating leukocytes, suggest that SPL expression may be induced in lymphocytes and other blood cells as they reach maturity. More precise studies of SPL expression during the stages of lymphocyte maturation and in other blood cell populations will be required to confirm this possibility.

A lower than expected frequency of SPL homozygous null mice are observed in crosses of mice heterozygous for the null allele, and congenital anomalies including runting, vascular and kidney defects and thoracic malformations of the sternum, ribs and vertebrae were all observed (39). These findings are consistent with our observation that SPL is strongly expressed
in specific regions of the adult and developing kidney and ribcage cartilage. The function of SPL in these locations is not known but may relate to the role of S1P in vascular maturation and/or cell migration, which has been shown to be defective in SPL null mouse embryonic fibroblasts. The pattern of SPL expression observed in the day 17 embryo indicates that the adult SPL expression pattern is established well before birth. Analysis of SPL expression in early stage embryos will provide more insight regarding the contribution of SPL function to the development and/or prenatal functions of specific organs.

Some of the SPL expression patterns we observed in developing and adult mice have not been previously reported or examined histologically. For example, SPL appears to be highly expressed in sebaceous glands of the skin, with lower but detectible expression in skin epithelial cells. This finding is interesting in light of previous reports that Sgpl1 expression is altered in atopic skin disease in canines and humans (63, 64). Additionally, the fatty acid dehydrogenase whose deficiency is responsible for the skin disorder Sjogren-Larsson syndrome was recently shown to catabolize the SPL product hexadecenal (65). Finally, a recent study implicated SPL and its impact on S1P metabolism and calcium homeostasis as potentially playing a role in Darier’s disease, which is caused by a defect in keratinocyte adhesion and differentiation (66). Our finding of SPL expression in discrete glomeruli of the olfactory bulb and embryonic olfactory mucosa are consistent with reports demonstrating Sgpl1 expression and activity in this location by other methods (67, 68). Combined with the finding of high SPL expression in the preputial gland, which is responsible for synthesis of pheromones, these findings suggest that SPL may play a role in olfactory organ function and the biosynthesis, metabolism and/or detection of pheromones. We recently showed that one of the products of the SPL reaction, hexadecenal, has biological activity and activates JNK signaling (52). Interestingly, some long
chain aldehydes function as pheromones, which suggests the possibility that SPL may be required for production of hexadecenal in olfactory organs (69-71). Our finding of SPL expression in Purkinje cell layer of the cerebellum is consistent with a recent study by Hagen and colleagues (72). SPL expression in other specific neurons of the forebrain, midbrain, hindbrain, cerebellum, spinal cord and trigeminal ganglion have not been previously reported and suggest that SPL may have important functions in these locations. This notion is supported by the observation that SPL null mice demonstrate neurodegeneration of the brain and that neurons which robustly express SPL are the first to degenerate in SPL null mice (72). SPL expression is critical for preventing S1P-induced apoptosis in hippocampal neurons, and sphingosine kinases are required for normal brain development (17, 72, 73). These observations suggest the need for tight regulation of S1P levels in the developing and adult brain. SPL expression in the dorsal root ganglia of the spinal cord, which contains the cell bodies of afferent spinal nerves, is interesting in light of the finding that S1P and other lysophospholipid mediators are important in nocioception (74, 75). The observation that SPL is highly expressed in brown adipose tissue is interesting in light of the previous finding that SPL null mice are lean, despite exhibiting high tissue and circulating lipid levels (31). SPL expression in the liver is consistent with the finding of metabolic and immune disregulation in liver tissues of SPL null mice (31). The finding of SPL expression in the developing and adult bladder epithelium and bronchial epithelium has not been previously reported. SPL expression in these locations and in the epithelium of the gut could indicate tissue-specific functions involved in maintenance of barrier integrity, immune defenses, and/or promotion of injured cell turnover.

The levels of S1P among different rodent tissues have been quantitated previously by several groups using different biochemical methods. In 1997, Yatomi showed that S1P levels in
rat tissues varied from lowest in liver, heart and skeletal muscle to highest in intestines, with intermediate levels in brain, kidney and spleen (76). S1P levels were not measured in the thymus (which we measured) but were high in testis (which we did not measure). In a study by Jiang and Han, S1P was determined in brain compartments and was found to be highest in spinal cord and brainstem, with lowest levels in cortex and intermediate levels in cerebellum (77). These findings show that SPL expression roughly correlates positively with S1P levels. This could be interpreted in two ways. SPL could have little bearing on S1P levels, since its high expression does not result in low S1P levels. Alternatively and most likely, high SPL expression is required in tissues subject to high S1P biosynthesis or uptake. Mice with reduced global SPL activity exhibit high levels of S1P in most tissues in comparison to control mice, as shown by others and ourselves (31, 40, 42, 44, 78). Even in tissues such as heart and skeletal muscle, wherein baseline SPL levels are very low to undetectable, SPL inhibition has significant effects on S1P levels. More detailed analysis of the impact of SPL expression in different tissues will require the conditional disruption of SPL in specific organs and tissues. It should be kept in mind that gross changes in tissue S1P levels may not accurately reflect the importance of highly localized gradients that influence the viability and migration of discrete cell populations. In addition, S1P levels may be controlled by S1P phosphatases, such as has been demonstrated in the thymus (29).

Lastly, SPL expression was not detectible in some tissues, notably the heart and skeletal muscles. However, we have shown that SPL is induced in the murine heart and skeletal striated muscle in response to ischemia and chemical injury, respectively (44, 46). SPL expression is regulated by both transcriptional and posttranscriptional mechanisms during development and in response to radiation, ischemia and other insults. Thus, it is important to recognize that the
baseline SPL expression patterns we observe likely underestimate the functions of SPL, especially those that may come into play under stressful conditions in which apoptosis is exacerbated and tissues are subject to injury or degeneration.
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FIGURE LEGENDS

**Fig. 1.** Whole mount survey of β-Gal stained adult SPL reporter mouse organs. (A) Whole brain. (B) Olfactory bulb, showing staining of specific olfactory ganglia. (C) Forebrain. (D) Midbrain. (E) Cerebellum and brainstem. (F) Hindbrain. (G) Basicranium, showing positively stained trigeminal nerve ganglia and unstained pituitary. (H) Spinal cord, showing grey matter staining. (I) Thymus and heart. Thymus is intensely stained. The pin is within the ventricular wall of the bisected heart, and the view shows the interior of the ventricle. (J) Kidney, showing strong staining in the renal pelvis. (K) Bladder, showing epithelial staining. (L) Female gonads including ovaries, fallopian tubes and uterus. (M) Foot. (N) Skin. (O) Preputial gland. (P) Ribcage. (Q) Liver. (R) Stomach. (S) Jejunum, with a Peyer’s Patch in the center and showing less staining than surrounding jejunal tissue. (T) Ileum. (U) Brown adipose tissue. (V) Harderian gland and eye. (W) Adrenal gland. (X) Tail. These data are representative of results obtained from four animals (two male and two female).

**Fig. 2.** β-Gal staining of SPL reporter mouse tissues. Adult mouse tissues were fixed and stained for β-Gal as described in Materials and Methods. (A) Brown fat. (B) Sebaceous glands and squamous epithelium of the skin. (C) Harderian gland. (D) tracheal epithelium. (E) spleen. (F) stomach. These data are representative of results from three animals.

**Fig. 3.** SPL IHC of wild type adult mouse tissues. (A) Bladder. (B) Skeletal muscle. (C) Bone marrow. (D) Ovary. (E) Colon. (F) Bone marrow. (G) Adrenal gland. (H) Kidney. (I) Spleen. These data are representative of results from three animals.
Fig. 4. SPL expression in gut-associated lymphoid tissues, lymph nodes and blood cells. (A) Peyer’s Patch of the jejunum. (B) Colorectal lymphoid aggregate. (C) Lymph node. These data from A-C are representative of results from three animals. (D) Thymocytes and splenocytes were isolated by mechanical dissociation of thymus and spleen, respectively. Remaining, thymic and splenic stromal cells were washed twice with PBS. Cell homogenate was subjected to immunoblotting and probed with an SPL antibody and an actin antibody. Shown is a representative immunoblot with the relative expression of SPL in each cell population given below, as determined by ImageJ quantification of autoradiogram signal and shown as SPL signal normalized to actin signal. (E) Leukocytes from spleen and whole blood were isolated, labeled with markers specific for each cell type indicated and separated using a cell sorter as described in Materials and Methods. Cell homogenates from each fraction were subjected to immunoblotting and probed with SPL and actin antibodies. Shown is a representative immunoblot of SPL expression in T lymphocytes, B lymphocytes, granulocytes and monocytes. The relative SPL expression of each cell population is given below, as determined by ImageJ quantification of autoradiogram signal and shown as SPL signal normalized to actin signal.

Fig. 5. Survey of SPL IHC positive brain tissues. (A) Cerebellum showing positive Purkinje cells and processes, as well as arachnoid meninges cells (top). Granular cell layers are non-immunoreactive (asterisks) and cerebellar astrocytes are also negative (arrows). (B) Choroid plexus lining cells are positive. (C) Olfactory neurons and their processes (left) at the edge of the olfactory bulb are one of the strongest staining tissues. (D) Brain stem neurons are relatively strong, comparable to the cerebellum and stronger than neurons of the cortex and thalamus. (E)
Meninges (left) and cerebral cortical neurons are moderately positive, while astrocytes are negative (arrows). (F) Some individual thalamic nuclei cells are positive (arrowheads), while others are negative for immunoreactivity.

**Fig. 6.** Validation of SPL detection systems and comparative IHC and β-Gal staining in adult mouse tissues. (A) SPL IHC of thymus from a WT mouse. Inset shows lack of staining in tissues from a SPL knockout mouse. (B) SPL expression by β-Gal staining of thymus from a SPL reporter mouse. Inset shows lack of staining in tissues from a wild type mouse lacking the SPL reporter insertion. (C) SPL expression by IHC of small intestinal villi from a WT mouse. Inset shows lack of staining from a SPL knockout mouse. (D) β-Gal staining of small intestinal villi from a SPL reporter mouse. Inset shows lack of staining in tissues from a WT mouse. (E) SPL expression by IHC of brain cortical neurons from a WT mouse. Inset shows lack of staining from a SPL knockout mouse. (F) SPL expression by β-Gal staining of brain cortical neurons from a SPL reporter mouse. Inset shows lack of staining in tissues from a wild type mouse. Bar = 100 µm.

**Fig. 7.** Relative SPL expression and activity in different adult mouse tissues. (A) SPL enzyme activity levels. (B) Relative SPL protein expression determined by quantitative analysis of IHC using polyclonal anti-murine SPL antibody. (C) Relative β-Gal staining intensity determined by quantitative analysis of β-Gal staining. AU = arbitrary units.

**Fig. 8.** Late stage embryo tissues. (I) Midsection E18 with highest areas of expression indicated by labels. (II) Histological sections showing the strongest and most specific expression. (A)
Mid-small intestine, jejunum. (B) Proximal small intestine, duodenum. (C) Thymus. (D) Upper respiratory mucosa and olfactory epithelium. (E) Bladder transitional epithelium. (F) Trigeminal ganglion. (G) Bone epiphysis. (H) Pancreas. (I) Pituitary gland. These data are representative of results from three late stage embryos.
FIGURES

Fig. 1
Fig. 8

I

Sensory Ganglia
Nasal Epithelium
Pituitary
Salivary Gland
Thymus
Brown Fat
Skin/Hair Follicles

1 mm

II

A
B
C

D
E
F

G
H
I

100 µm