Decreased signal transducers and activators of transcription (STAT) protein expression in lymphatic organs during EAE development in mice

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Abstract

Experimental autoimmune encephalomyelitis (EAE) is mediated by myelin-specific CD4+ T cells secreting Th1 and/or Th17 cytokines. Signal transducer and activator of transcription (STAT) family proteins have essential roles in transmitting Th1 and/or Th17 cytokine-mediated signals. However, most studies demonstrating the importance of the STAT signaling system in EAE have focused on distinct members of this family, often looking at their role specifically in the central nervous system, or in vitro. There is limited information available regarding the temporal and spatial expression patterns of each STAT protein and interplay between STAT proteins over the course of EAE development in critical lymphatic organs in vivo. In the present study, we demonstrate dramatic and progressive decrease of all six STAT family members (STAT1, STAT2, STAT3, STAT4, STAT5, STAT6) in the spleen and lymph nodes through the course of EAE development in SJL/J mice, in contrast with almost steady expression of thymic STAT proteins. Decreased splenic and lymphatic STAT expression was accompanied by significant enlargement of the spleen and lymph nodes, and histological proliferation of T cell areas with remodeling of the splenic microstructure in EAE mice. All STAT family members except STAT2 were mainly confined in T cell areas in spleen, whereas they were distributed in a protein specific manner in thymus. We present here a comprehensive analysis of all six members of the STAT family in spleen, lymph nodes and thymus through the development phase of EAE. Results suggest that EAE induced inflammatory T cells may develop distinct biological features different from normal splenic T cells due to altered STAT signaling.

Keywords: Signal transducer and activator of transcription, signalling, experimental autoimmune encephalomyelitis, multiple sclerosis, mice

Introduction

Experimental autoimmune encephalomyelitis (EAE) in mice is a commonly used animal model of multiple sclerosis (MS) that has provided valuable insight into the pathobiology of the human disease [1,2]. EAE is induced in mice by active priming with myelin proteins or specific myelin peptide epitopes in adjuvant. Demyelination and paralytic episodes are associated with infiltration of myelin-specific inflammatory Th1 CD4+ T cells into the central nervous system (CNS) [3]. Although initially described as an IL-12 driven Th1 cell mediated disease, it was recently shown that EAE can also be an IL-23 driven Th17 cell mediated disease [4]. Moreover, EAE can be induced by adoptive transfer of myelin-specific CD4+ T cells [5]. All the above data demonstrate the significance of T cells in EAE induction [6]. However, the mechanisms which underlie T cell proliferation and differentiation associated with EAE induction and development are not fully understood, and details of the signaling pathways that induce Th1 or Th17 cells in EAE are complex and not completely known.

It has been shown that the cytokine milieu is important for CD4+ T cell proliferation and differentiation [4]. Signal transducer and activator of transcription (STAT) family proteins have essential roles in transmitting many cytokine-mediated signals and thereby have similarly crucial roles in T helper cell proliferation and differentiation [7]. The binding of cytokines to their cognate receptors differentially activates 6 STAT proteins (STAT1–6) [8], which in turn regulate the expression of genes involved in T cell homeostasis, growth, differentiation, apoptosis and immune response [7]. Previous studies demonstrated that blocking of IL-12 in Th1 cells [9] or IL-17A in Th17 cells [10,11], two cytokines that signal through the Janus Kinases (JAK)-STAT pathway, suppressed EAE development [12,13]. Therefore, the STAT signaling pathway has been targeted as a site for potential intervention for EAE and MS treatment [9-11,13-15].

While a role of STAT signaling in EAE has been suggested, studies are limited. However, studies in other models have demonstrated the general importance of this pathway in maintenance of the cellular integrity and healthy growth of T cells. For example, dysregulation of the JAK-STAT pathway plays a role in many primary human T cell lymphomas [16], and STAT knockout mice exhibit severely impaired lymphoid development and differentiation [17]. Specifically, STAT1 [18] and STAT4 [19,20] function predominantly in promoting Th1 cell differentiation [21], whereas STAT3 is a key signaling molecule essential for Th17 lineage commitment [22]. STAT6 triggers Th2 cell devolution [23], and IL-2-activated STAT5 facilitates Treg cell differentiation [24].

Thus, the importance of the STAT signaling system in T cell development, differentiation, and survival is well established. However, studies of the importance of STAT signaling in EAE
development have largely focused on just one or two specific STATs, often examining the role in the CNS [25,26] or in vitro [10]. In contrast, less information is available from lymphatic organs in vivo in EAE. This is important to examine because EAE is a systemic autoimmune disease, with myelin-reactive T cells first generated and expanded in the peripheral lymphatic system. Prior limited studies examining effects of STAT deletion on CNS inflammation in EAE are likely downstream effects, influenced by T cell differentiation and proliferation in the periphery prior to cells gaining entry into the CNS. The temporal and spatial expression patterns of all members of the STAT family over the course of EAE development and the interplay between STAT proteins in mediating T cell proliferation and differentiation in critical lymphatic organs in vivo are not known. In the current studies, we defined the time course of changes in STAT expression in relation to the development of EAE in primary (thymus) and secondary (spleen and lymph nodes) lymphatic organs. We characterize the cell types and histological microstructure in those lymphatic organs in order to identify altered distribution of STAT proteins.

Materials and methods

Mice

Eight-week-old female SJL/J mice (n=34) were purchased from the Jackson Laboratory (Bar Harbor, ME) for induction of EAE. Treatment of animals was reviewed and approved by the Institutional Animal Care and Use Committee at the University of Pennsylvania.

Induction of EAE

EAE was induced as previously described [27]. Briefly, eight-week-old female SJL/J mice, each weighing 20g (20±1g) were anesthetized with ketamine/xylazine and were injected subcutaneously at two sites on the back with 0.1 mL solution containing 0.5 mg/mL proteolipid protein peptide 139–151 (PLP; GenScript, Piscataway, NJ) emulsified in complete Freund’s adjuvant (CFA; Difco, Detroit, MI) containing 0.5 mg/mL mycobacterium tuberculosis (Difco). Control mice were injected with equal volumes of PBS and CFA. All mice were injected with 200 ng pertussis toxin (List Biological, Campbell, CA) in 0.1 mL PBS intraperitoneally on day 0 (day of immunization) and again on day 2.

Thymus, lymph nodes and spleen were harvested from mice on day 5 (n=12, six animals for each group, Control and EAE, respectively) and day 10 (n=12, six animals/group) post-immunization. On day 12 (n=10, five animals/group), only thymus and spleen were harvested. Harvested thymus and spleen were weighed first before being frozen in liquid nitrogen and then stored at -80°C until being used for protein extraction for Western blot analysis. A portion of thymus and spleen from two mice of each group was embedded in OCT compound (Sakura Finetek, Torrance, CA), slowly frozen on dry ice, and stored at -80°C for sectioning for immunohistochemical analysis.

Antibodies

All rabbit anti-mouse STAT antibodies (STAT1, STAT2, STAT3, STAT4, STAT5 and STAT6) were purchased from Santa Cruz (Dallas, TX) and used at a dilution of 1:200 for both Western blot and immunohistochemical analysis, except for STAT1, for which a dilution of 1:10,000 was used. The STAT5 antibody detects both STAT5 A and B isoforms. The mouse anti-human β-actin antibody was purchased from Sigma (Sigma-Aldrich, St. Louis, MO).

Protein extraction

Protein was extracted from cytosolic and nuclear cellular fractions. Thymus, lymph nodes and spleen were ground into small pieces, homogenized in TED buffer (50 mM Tris [pH 7.4], 10 mM EDTA, and 1 mM diethyldithiocarbamic acid (DEDTC)) containing 2 mM octyl glucoside, and centrifuged at 16,000 x g for 30 min at 4°C. The supernatant (cytosol) was retained and stored at -80°C until used for electrophoretic analyses. The crude pellets containing cell nuclei from the spleens, lymph nodes and thymus were sonicated (8-sec cycle, three cycles; Branson Sonifier, Danbury, CT) in 500 µl TED sonication buffer (20 mM Tris [pH 7.4], 50 mM EDTA, and 1 mM DEDTC containing 45 mM octyl glucoside). The sonicates were centrifuged at 16,000 x g for 25 min at 4°C. The recovered supernatants were stored at -80°C until electrophoretic analyses were performed. In lymph nodes, STAT expression was only analyzed in the nuclear fraction since STAT expression in spleen was almost identical in both cytosolic and nuclear compartments.

Western blotting

The protein concentration was determined in both cytosolic and nuclear fractions using the BCA protein assay according to the manufacturer’s instructions (ThermoFisher Scientific, Rockford, IL). Proteins (30µg/lane) were then separated by 7.5% SDS-PAGE and electrophoretically transferred to a nitrocellulose membrane with use of a Bio-Rad transfer blot cell. The membranes for immunoblotting were blocked with 2% BSA in 10 mM Tris-Cl buffer containing 0.1% Tween 20. After blocking, blots were incubated with rabbit anti-mouse STAT1, STAT2, STAT3, STAT4, STAT5 or STAT6 antibodies at 4°C overnight, then washed and incubated with peroxidase-conjugated goat anti-rabbit IgG (Amersham Life Sciences, Arlington Heights, IL) at room temperature for 1 h. After each antibody incubation, blots were washed four times (10 min each) in wash buffer. Protein bands were visualized by means of an enhanced chemiluminescence Western blotting detection kit (Amersham). To determine the molecular size of the proteins, standard molecular weight marker protein (Bio-Rad) was run in an adjacent lane. For normalization of signals, blotted membranes were stained for β-actin (Sigma). The intensity of each band was determined using Image J software (NIH.gov). Briefly, the first band is framed with the rectangle selection tool in Image J, then the “Analyze-Gels-Select Next Lane”
command is used to select an equal area in adjacent lanes. Average pixel intensity is plotted across the selection box in each lane, with the protein band seen as a peak above the background pixel intensity. The area under the peak is calculated. For each protein, the band intensity (area under the peak) is normalized to the band intensity of the beta-actin band in each lane calculated by the same method.

**Immunocytochemistry**

Frozen thymus and spleen were sectioned at 5 µm on a cryostat (Microm HM 550, ThermoFisher Scientific, Kalamazoo, MI). There were not enough lymph nodes to proceed with immunohistochemistry in the same animals after the tissue was used for protein extraction for Western blot analysis. Sections were stained for detection of STAT proteins using the avidin-biotin complex method (Elite kits, containing avidin [solution A] and biotin [solution B]; Vector Laboratories, Burlingame, CA). Sections were first fixed in 3.7% formaldehyde (Fisher Scientific) at room temperature followed by incubation in cold methanol for 4 min and acetone for 1 min at -20°C. The sections were then rinsed with 0.05 M Tris-Cl (TBS), pH 7.4 two times (5 min each). All sections were incubated with the primary antibodies (STAT1, STAT2, STAT3, STAT4, STAT5, or STAT6) overnight at 4°C. Sections were then rinsed 2 times (5 min each) in TBS before incubation with biotinylated goat anti-rabbit IgG in TBS for 1 h at room temperature, then rinsed 2 times (5 min each) in TBS. Finally, the sections were transferred to AB solution (22.5 µl solution A and 22.5 µl solution B in 10 ml TBS) for 30 min. The AB solution was made at least 30 min prior to use. Immunostaining for all STATs was visualized with 3,3-diaminobenzidine tetrahydrochloride (DAB; Sigma), followed by hematoxylin counterstaining. Specificity of immunostaining was verified by including the following controls: 1) no addition of primary antibody, 2) replacement of primary antibodies with normal rabbit or normal mouse serum, 3) Western blot analysis.

**Statistical analysis**

Each STAT protein expression level was calculated as the ratio of the Western blot band intensity of STAT:β-actin. Comparison of the mean protein expression level between groups (Control: n=5-6 vs EAE: n=5-6 per group) was made by unpaired Student’s t-test. Statistical significance was assumed at the 5% level. Data were presented throughout as mean ± SEM.

**Results**

**Enlarged spleen in EAE mice**

EAE was induced in 8 week old female SJL/J mice by immunization with PLP. At the time of sacrifice, the spleen and thymus were removed and weighed. There was a significant increase of the average wet weight of the spleens from EAE mice on days 5, 10 and 12 post-immunization (*P<0.01), with the weight of spleens from EAE mice found to be double the weight of spleens from control mice (Figure 1). There was no significant change in the average thymus weight between EAE and control mice (data not shown). The visualized size of lymph nodes dissected at the time of sacrifice was also noted to be larger in EAE mice than in control mice (data not shown).

**Stat expression is significantly decreased in the spleen during EAE development**

STAT proteins move between cytoplasm and nuclei [28], therefore, each STAT family member was quantified in both cytoplasmic and nuclear extracts isolated from thymus and spleen. We first examined expression of all six STAT proteins in the primary (thymus) and the secondary (spleen) lymphatic organs of EAE mice 12 days post-immunization, the peak time of onset of EAE [27]. Protein was extracted from 10 mice (five controls and five EAE). Western blot analysis demonstrated that expression of all STAT family members, except STAT2, decreased significantly (***P<0.01, **P<0.05, Figures 2A and 2B) in EAE mice in both cytosolic and nuclear fractions of protein extracts isolated from spleen. No significant change in STAT2 level was found (Figures 2A and 2B). In contrast, in the thymus only STAT5 levels decreased significantly in EAE mice (Figure 3).
A trend of decline in the level of expression of STAT6 in thymus from either cytosolic or nuclear extracts of EAE mice did not reach statistical significance (Figure 3).

Time course of changes in splenic, lymphamatic and thymic stat expression during development of EAE
Because significant changes in expression levels of almost all members of the STAT family were noted in splenic protein extracts at peak onset of EAE, STAT expression was also measured at earlier time points to assess temporal changes in protein levels. Spleen, lymph nodes and thymus were isolated from SJL/J mice on day 5 (prior to onset of any clinical EAE symptoms) and day 10 (earliest onset of EAE symptoms in some mice) post-immunization. 12 mice (six controls and six EAE) were sacrificed at each time point. On day 5, there was a significant decrease (**P<0.01, Figures 4A and 4B) and day 10 (**P<0.01, Figures 4A and 4B) proteins in both cytosolic and nuclear fractions from EAE mice as compared with extracts from control mice. There was a trend toward decreased expression of STAT6 protein in both cytosolic and nuclear fractions from EAE mice, however this did not reach significance (P>0.05). STAT1, STAT2, STAT3 and STAT4 levels were equivalent between EAE and control mice. Values are presented as mean ± SEM.
Figures 4A and 4B. The level of STAT1 protein in the spleen did not change in either cytosolic or nuclear extracts of the EAE mice compared with control mice on day 5 (Figures 4A and 4B). There was no change in expression levels of any STAT family members in the thymus in either cytosolic or nuclear extracts of EAE mice as compared to control mice on day 5 (Figure 5), and similarly the level of all six STAT proteins in extracts from lymph nodes of EAE mice was equal to lymph node expression levels in control mice, in both nuclear (Figure 6) and cytosolic fractions (data not shown), on day 5.

There was a more prominent and consistent decrease (Figures 7A and 7B) in expression of all STAT proteins in the spleen of EAE mice on day 10 post-immunization. Analysis of both cytosolic and nuclear fractions of protein extracts from spleen showed significantly lower levels of STAT1, STAT4, STAT5 and STAT6 in EAE mice (**P<0.01, Figures 7A and 7B) as compared with control mice. STAT3 was only reduced in the nuclear (**P<0.01, Figures 7A and 7B), but not cytosolic (P>0.05, Figures 7A and 7B) fractions of protein extracts from EAE mouse spleens. STAT2 protein was not detectable in splenic extracts from any mice on day 5 or day 10 (data not shown) post-immunization. Similar to splenic STAT proteins, protein levels of all STAT family members, except STAT1, were lower in lymph node extracts from EAE animals as compared with extracts from control mice.
Figure 6. Protein was extracted from lymph nodes isolated from EAE mice 5 days post-immunization. (A) Western blots of the nuclear fractions of protein extracts from lymph nodes of control (lanes 1–6, n=6) and EAE (lanes 7–12, n=6) mice are shown. (B) There was no difference in the level of expression of any of the STAT proteins in lymph nodes from EAE mice as compared with levels in control mice (P>0.05). Values are presented as mean ± SEM.

control mice, although the decrease in STAT5 did not reach statistical significance (Figure 8).

STAT1, STAT3, STAT4, STAT5 and STAT6 protein levels in both cytosolic and nuclear fractions of extracts from thymus of EAE mice isolated 10 days post-immunization were no different than levels found in control mice (Figure 9). STAT2 was not detectable in protein extracts from the thymus of any mice on day 5 or day 10 (data not shown).

Immunolocalization of stat proteins in the spleen and thymus

Because profound changes in protein expression levels of most STAT family members were observed in EAE, immunohistochemical staining of tissue sections was used to determine the pattern of STAT expression and to localize changes within spleen and thymus. Spleen and thymus were dissected from control and EAE mice following sacrifice on day 12 post-immunization. Tissues were frozen, cut into 5μm sections, and stained with antibodies to specific STAT family members for immunohistochemical visualization of STAT protein expression. In sections of spleen from control mice (Figure 10A-10F, the top row), STAT1 (A), STAT2 (B), STAT3 (C), STAT4 (D), STAT5 (E) and STAT6 (F) proteins were mainly localized in the T cell area of the white pulp, with predominantly nuclear staining (Figure 10A-10F). In sections of thymus from control mice (Figure 10G-10L, the bottom row), immunostaining for STAT proteins (G: STAT1; H: STAT2; I: STAT3; J: STAT4; K: STAT5; L: STAT6) was also localized within nuclei, except STAT2 (H), which was present in both cytoplasm and nuclei.

There was enormous histological reconstruction of the immunostaining pattern in spleens isolated from EAE mice (Figure 11), as compared to the pattern in control mouse
spleens. Multiple circular secondary lymphatic follicles were observed, with excessive proliferation of T cell areas. Immunostaining for all STAT proteins in spleens from EAE mice was mainly confined within these follicles, except STAT2, which was localized in all other areas of the spleen, but excluded from the T cell areas of the follicles. In sections of spleens from control mice, immunostaining for all STAT proteins was observed in both red as well as white pulp (Figure 11, the top row), whereas in spleens from EAE mice (Figure 11, the bottom row) all STAT proteins except STAT2 were primarily localized in the T cell areas of the newly formed follicles. The wider distribution of STAT protein expression shows that the number of positively stained cells was larger in spleens from control mice than in spleens from EAE mice (Figure 11). The intensity of the immunostaining for all splenic STAT proteins was also stronger in spleen sections from control mice than in spleen sections from EAE mice (Figure 11).

**Discussion**

EAE is a T cell mediated CNS disease [29,30]. Although the relative roles of Th1 [31,32] or Th17 [4] or both T cell lineages are still being examined, induction of EAE by adoptive transfer of either activated Th1 or Th17 cells [6,32] further reinforces the idea that EAE, and possibly human MS, are T cell mediated autoimmune CNS diseases. Enlargement of the spleen and lymph nodes observed in EAE mice in the current study is consistent with proliferation of lymphatic cells. The detectable physical expansion was restricted to secondary
lymphatic organs, with no visible change in thymus after EAE induction. While these findings may be expected due to the recognized role of T cells in EAE pathogenesis, interestingly a prior study reported a noted decrease in the overall weight of the spleen after EAE induction [33], in which homogenized brain tissue was used to induce EAE in Lewis rats and spleen weight was recorded at the time of death around 25 days post-immunization. In the current study, spleen weight was measured earlier, through the course of EAE development, on days 5, 10 and 12 post-immunization. Different experimental factors may account for the contrary observations, such as different species (mice vs rats), the antigens used to produce EAE (PLP vs whole brain homogenate), and the timing of recorded spleen weight (EAE development vs chronic phase of EAE).

To explore potential pathological mechanisms driving T
of all family members in lymph nodes of EAE mice equal to
similarity, but was more delayed in onset, with expression
family members in spleen began to decline by day 5 post-
STAT family members (all except STAT2) were downregulated in
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lymphatic organs during EAE development.

In addition it is not known whether STAT proteins play a more
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mice are highly susceptible to EAE and develop more severe
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severe EAE than wild-type mice [43]. Similarly, STAT1 knockout
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been suggested by analysis of individual gene knockout
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family kinases [36,37]. Upon cytokine binding, activated Jaks
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activate gene transcription.
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family kinases [36,37]. Upon cytokine binding, activated Jaks
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phosphorylate tyrosine residues on the cytokine receptor
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promotes phosphorylation of STAT, leading to its release
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from the cytokine receptor, dimerization, and translocation
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into the nucleus. STAT dimers then bind promoter regions
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of cytokine-inducible genes and activate gene transcription
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[36,37], facilitating T cell proliferation and differentiation.

Distinct functions of individual STAT family members have
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been suggested by analysis of individual gene knockout
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mice [38,39]. STAT4 is activated mainly by IL-12 and type I
IFNs, and it functions predominantly in promotion of Th1
cell differentiation [40]. STAT6 is activated in response to IL-4
and functions as a molecular switch for initiation of the Th2
cell differentiation program [40,41]. After activation by IL-6,
STAT3 triggers Th17 commitment [22]. On the other hand, IL-
2-activated STAT5 facilitates Treg cell differentiation [10,28].
In EAE, studies using mice containing deletions of individual
STAT family members suggest variable effects of this family
on demyelinating disease. STAT6-deficient mice develop more
severe EAE than wild-type mice [23]. Similarly, STAT1 knockout
mice are highly susceptible to EAE and develop more severe
and accelerated disease with atypical neuropathologic features
[18]. However, mice deficient in STAT4 are resistant to EAE,
with minimal inflammatory infiltrates observed in the CNS
[23]. STAT3 is essential for early embryonic development [42],
therefore isolated function of STAT3 in EAE can’t be assessed
by conventional gene knockout. Thus, important functions of
specific STATs are recognized in EAE, but overall changes and
interplay of members of this family are not well understood.
In addition it is not known whether STAT proteins play a more
critical role in development of myelin reactive T cells in the
peripheral immune system, or modulate entry of pathogenic
T cells into the CNS and mediate localized tissue responses. It
is for this reason that all six members of the STAT family were
analyzed in the primary (thymus) and secondary (spleen and
lymph nodes) lymphatic organs during EAE development.

Results show that on day 12 post-immunization, a time when
clinical EAE nears its peak [27], protein levels of five of the six
STAT family members (all except STAT2) were downregulated in
spleen, whereas in thymus only STAT5 decreased significantly.
Downregulation of STAT expression began even earlier, and
progressed during EAE induction, as expression of most STAT
family members in spleen began to decline by day 5 post-
immunization, with further decreases found on day 10. In
lymph nodes, expression of STAT family members followed a
similar pattern, but was more delayed in onset, with expression
of all family members in lymph nodes of EAE mice equal to
levels in control mice 5 days post-immunization, but expression
of all STAT proteins, except STAT1, were decreased on day 10.
This study provides the first comprehensive analysis of STAT
protein expression during EAE development in critical primary
and secondary lymphatic organs in vivo. A previous study [43]
reported expression of STATs 1-6 was significantly enhanced
in brain and spinal cord of EAE rats. Interestingly, the cells
expressing STATs in the CNS [43] were astrocytes and microglia,
not T cells, and STAT expression in spleen, lymph nodes and
thymus were not reported. Another recent study [44] examined
Jak-STAT expression in vitro, in splenocytes isolated from EAE
animals. The authors demonstrate that amelioration of EAE by
Plumbagin, a natural bicyclic naphthoquinone, is associated
with downregulation of the JAK-STAT pathway, as opposed to
our results suggesting downregulation is associated with
induction of the EAE disease itself. Differences may be seen
because our results represent effects on splenic and lymphatic
STAT expression in vivo, while they measured expression in vitro
after isolation of splenocytes from EAE animals.

While prior analysis of STAT knockout mice demonstrated
distinct, sometimes opposite, functions of individual STAT
family members in EAE, our results show almost all members
decrease over a similar time course in spleen and lymph
nodes of EAE mice. Germine knockout studies analyze a
zero (knockout) vs all (wild type) effect of a given gene, while
the downregulation of the STAT pathway in EAE spleen and
lymph nodes is a gradual and progressive process, and most
STATs were not turned off completely even by day 12 post-
immunization. Also, the complex interplay between all STAT
proteins together, along with other pathways functioning in
primary and secondary lymphatic organs, may be orchestrated
in an orderly fashion through the development of EAE. This
may not be replaced by analysis of the absence of one
individual gene. Therefore, it is also important to consider
the overall pattern of changes in this family in conjunction
with individual gene knockout data in order to extend and
expand our understanding of the function of the STAT family
in mediating EAE.

The significance and mechanism underlying the decrease of
almost all STAT family members in spleen and lymph nodes of
EAE mice is not yet understood, although prior studies and the
current data suggest possible important mechanisms. Previous
studies described a number of negative regulators of the Jak-
STAT signaling pathway, including tyrosine phosphatases [45],
suppressors of cytokine signaling (SOCSs, 46), and protein
inhibitors of activated STAT (PIASs, 45). STAT signaling is also
regulated by STAT acetylation through SirT1 [47]. Future
studies of these regulators should be considered to determine
the cause of the massive downregulation of splenic and
lymphatic STAT proteins associated with EAE development.
Nonehedless, the suppression of STAT expression suggests
the biological importance of this signaling pathway in EAE
induced inflammatory T cell proliferation and differentiation.
Whether an alternative cellular signaling pathway replaces the
STAT system in EAE T cells, or inhibition of the splenic STAT system is a prerequisite for normal T cells to be transformed into myelin reactive T cells merits further study.

Observed changes in expression of specific STAT family members in the current study may suggest potential mechanisms by which STAT signaling modulates EAE development. Interestingly, STAT5 was the only protein which decreased significantly in thymus upon EAE induction. A human STAT5 mutation results in severe immunodeficiency [48], and STAT5 facilitates regulatory T cells, not Th1, Th2 or Th17 differentiation [24], thus it is possible STAT5 downregulation reduces Treg populations that could inhibit EAE. Lack of thymic enlargement and absence of any change in expression of other STAT proteins in thymus suggests this pathway is more important for T cell proliferation in the secondary lymphatic organs.

Another distinct and isolated change occurred in lymph nodes, where expression of all STAT family members except STAT1 decreased on day 10 post-immunization. STAT1 knockout mice are more prone to EAE and develop more severe and accelerated disease [18]. Thus, sustained STAT1 expression in EAE lymph nodes may act as one mechanism to fight against the occurrence and progression of the disease after EAE induction.

In addition to enlargement in the size of the spleen, histological analysis revealed tissue reorganization and redistribution of the localization of STAT proteins. There are many newly formed secondary follicles, which explain the physical enlargement of EAE spleen. All STAT proteins except STAT2 were localized mainly in T cell areas of the white pulp of control mice and in T cell areas of spleen follicles of EAE mice, consistent with prior studies suggesting the STAT pathway may regulate T cell proliferation and differentiation [22,28,34,35]. The distribution of STAT 2 differed from other STAT proteins. Notably, STAT2 was the only protein not restricted to nuclei, but found also in cytoplasm of the thymus. Moreover, in spleen STAT2 was localized to other cell types, but not the T cell area. The physiological implication of the difference in cellular distribution of STAT2 is not clear, although it suggests that this member of the STAT family, unlike other STATs, may not play a prominent role in EAE development.

Conclusion
We present here detailed expression analysis of all six members of the STAT family in thymus, lymph nodes and spleen through the development phase of EAE in mice. Downregulation of splenic and lymphatic STAT protein expression started on day 5 post-immunization, and progressed to nearly undetectable levels by day 12, which was in contrast with almost steady expression of STAT proteins in thymus of EAE mice. Decreased STAT expression was accompanied by physical enlargement of the spleen, histological remodeling of the microstructure and proliferation of T cell areas in EAE spleen. All STATs except STAT2 were mainly confined to T cell areas in spleen, whereas thymic STATs were distributed in a protein specific manner. Our data suggest that downregulation of almost all STAT signaling is important for pathologic transformation of inflammatory T cells during EAE development.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions

| Authors’ contributions | WXX | LZ | KED | KSS |
|------------------------|-----|----|-----|-----|
| Research concept and design | ✓ | - | - | ✓ |
| Collection and/or assembly of data | ✓ | ✓ | ✓ | - |
| Data analysis and interpretation | ✓ | - | - | ✓ |
| Writing the article | ✓ | - | - | - |
| Critical revision of the article | ✓ | - | - | ✓ |
| Final approval of article | ✓ | - | - | - |
| Statistical analysis | ✓ | - | - | ✓ |

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