NF-κB–inducing kinase plays an essential T cell–intrinsic role in graft-versus-host disease and lethal autoimmunity in mice

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NF-κB–inducing kinase (NIK) is an essential upstream kinase in noncanonical NF-κB signaling. NIK-dependent NF-κB activation downstream of several TNF receptor family members mediates lymphoid organ development and B cell homeostasis. Peripheral T cell populations are normal in the absence of NIK, but the role of NIK during in vivo T cell responses to antigen has been obscured by other developmental defects in NIK-deficient mice. Here, we have identified a T cell–intrinsic requirement for NIK in graft-versus-host disease (GVHD), wherein NIK-deficient mouse T cells transferred into MHC class II mismatched recipients failed to cause GVHD. Although NIK was not necessary for antigen receptor signaling, it was absolutely required for costimulation through the TNF receptor family member OX40 (also known as CD134). When we conditionally overexpressed NIK in T cells, mice suffered rapid and fatal autoimmunity characterized by hyperactive effector T cells and poorly suppressive Foxp3+ Tregs. Together, these data illuminate a critical T cell–intrinsic role for NIK during immune responses and suggest that its tight regulation is critical for avoiding autoimmunity.

Introduction

Naive T cells require several signals to become activated in vivo. TCR stimulation and costimulation via CD28 are required for cell division and IL-2 production, but in the absence of infection or damage, these 2 signals are insufficient for an effective T cell response, and T cells die or become anergic after initial proliferation (1). Additional costimulation (signal 3) is required for continued clonal expansion, survival, and differentiation into effector subsets, and TNF receptor family members (TNFRs) constitute an important group of costimulatory molecules that relay signal 3 to antigen activated T cells (2). In particular, OX40 (also known as CD134 and TNFRSF4) is critical for optimal CD4+ T cell expansion, survival, differentiation, and memory responses to a variety of infectious and noninfectious immune challenges, including autoimmunity and allogeneic responses in the context of graft-versus-host disease (GVHD) (3).

OX40, like other TNFRs as well as the TCR complex itself, activates NF-κB (3). Activation of NF-κB can be described in terms of 2 interrelated pathways (4). The canonical NF-κB pathway depends on inhibitor of NF-κB kinase (IKK) subunit β, which rapidly phosphorylates inhibitory IκB proteins, leading to their degradation. This releases active dimers composed primarily of p50:RelA or p50:p52 c-rel subunits to translocate to the nucleus and induce proinflammatory gene transcription. In contrast, the noncanonical (also known as alternative) NF-κB pathway depends on accumulation of NF-κB–inducing kinase (NIK) and subsequent phosphorylation of IKK subunit α, which then targets the inhibitory subunit p100 (also known as NF-κB2) for partial degradation, producing the active p52 subunit. In this pathway, gene transcription is mediated primarily by active p52:RelB dimers. Kinetics also distinguish the canonical from the noncanonical NF-κB pathways, in that activation of the canonical pathway is immediate, but its duration short as a result of rapid negative feedback, whereas activation of the noncanonical pathway is more delayed but sustained (4).

While lesions in the canonical NF-κB pathway have devastating effects on the immune system (4), genetic manipulation of the noncanonical NF-κB pathway has more limited effects because of the limited number of receptors that activate this pathway. NIK knockout mice and alymphoplasia (aly) mice, which harbor a naturally occurring loss-of-function mutation in NIK (5), have disorganized lymphoid structure in the spleen and thymus, no LNs, and few peripheral B cells (6). These defects have been attributed to defective signaling by lymphotoxin β receptor (LTβR) and other TNFRs on lymphoid organ stromal cells and lack of survival signals transmitted by B cell activating factor receptor (BAFFR) on peripheral B cells (7–9). In addition, some defects in APC function have been attributed to defective signaling though CD40 (10, 11).

In contrast, steady-state peripheral T cell populations are fairly normal in NIK-deficient mice, and the role of NIK in T cell function remains unclear. Autoimmunity has been reported in aly/aly mice (12), but this appears to be due to a lack of TNFR signaling in thymic epithelial cells that mediate negative selection and development of Tregs (13). Similarly, although aly/aly mice do not reject skin grafts, this was attributed to their lack of LNs, rather than a T cell–intrinsic defect (6, 14). Findings of in vivo investigations of NIK-deficient T cells have been equivocal. Unfractionated aly/aly T cells showed a diminished response to anti-CD3 stimulation, but allogeneic proliferative responses were normal (15), and sorted naive aly/aly T cells were subsequently shown to be hyper-responsive to TCR stimulation in vitro and in vivo (16). In contrast, 3 in vivo studies showed that NIK-deficient T cells fail to induce autoimmune disease in mouse models of arthritis and EAE (17–19).

To explain these contradictory data, we propose that NIK is required for naive T cell responses in vivo by signaling downstream of costimulatory TNFRs that promote acquisition of

Conflict of interest: The authors have declared that no conflict of interest exists.

Citation for this article: J Clin Invest. 2011;121(12):4775–4786. doi:10.1172/JCI44943.
NIK-deficient CD4+ T cells do not cause lethal GVHD. To test whether NIK is required in a T cell–intrinsic manner for allogeneic responses, we sorted naive CD4+ T cells from aly/aly mutant mice or littermate controls and transferred them into sublethally irradiated MHC class II mismatched congenic (H-2bm12×CD45.1)F1 recipients. Transfer of T cells with 2 normal copies of NIK caused lethal GVHD in all recipients, which was preceded by rapid weight loss starting 2 weeks after transfer (Figure 1). In stark contrast, aly/aly T cells caused no GVHD lethality or weight loss. Interestingly, recipients of heterozygous aly/+ T cells showed an intermediate phenotype, experiencing delayed disease and a 70% mortality rate. The importance of NIK levels seen here mirrors recent reports showing that loss of a single copy of NIK can rescue TNF receptor–associated factor 2 and TNF receptor–associated factor 3 knockout mice from perinatal lethality (20, 21). The failure of NIK-deficient T cells to induce lethal GVHD is likely a result of poor accumulation of allospecific T cells in the recipients, as illustrated in a separate experiment in which we transferred a smaller, nonlethal number (1×10^6) of naive CD4+ T cells into sublethally irradiated (H-2bm12×CD45.1)F1 recipients. In this case, despite lack of overt disease, control CD4+ T cells accumulated in mismatched recipients, and 10-fold more control CD4+ T cells than aly/aly T cells were recovered from recipient spleens 1 month after transfer (Table 1).

**Results**

NIK-deficient CD4+ T cells do not cause lethal GVHD. To test whether NIK was essential for lethal GVHD in a CD4+ T cell–intrinsic manner, Naive NIK-deficient T cells responded normally to TCR stimulation in vitro and to alloantigen in vivo under conditions of low costimulation in which T cells divide but do not differentiate or cause disease. In contrast, NIK was absolutely essential for OX40-mediated costimulation of allospecific T cells in vivo, and OX40 activated the NIK-dependent noncanonical NF-kB pathway in primary T cells. Finally, overexpression of NIK in T cells led to an aggressive lethal autoimmunity associated with decreased function of Tregs and effector T cells that are resistant to suppression. We therefore concluded that NIK is essential for T cell function in vivo and that its tight regulation in T cells is crucial to prevent autoimmunity.

**Figure 1**

NIK-deficient CD4+ T cells do not cause lethal GVHD. 1×10^6 naive +/+ , aly/+ , or aly/aly CD4+ T cells were injected intravenously into sublethally irradiated (H-2bm12×CD45.1)F1 recipients. (A) Survival is combined data from 4 independent experiments; each genotype was represented in 2–4 experiments. P < 0.05, +/+ versus aly/+ and aly/+ versus aly/aly; log-rank test. (1 mouse in the aly/aly group died only 4 days after transfer, which is much earlier than GVHD onset in this model). (B) Body weight data (± SEM) are representative of 2 independent experiments.

**Table 1**

NIK-deficient T cell accumulation after transfer into (H-2bm12×CD45.1)F1 MHC mismatched recipients

<table>
<thead>
<tr>
<th>Recipient manipulation</th>
<th>aly/+ donor</th>
<th>aly/aly donor</th>
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<tbody>
<tr>
<td>Low costimulation</td>
<td></td>
<td></td>
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<tr>
<td>Unirradiated, control Ig</td>
<td>5.3 ± 0.36</td>
<td>6.5 ± 0.99</td>
</tr>
<tr>
<td>High costimulation</td>
<td></td>
<td></td>
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<tr>
<td>Unirradiated, anti-OX40</td>
<td>17.4 ± 1.96</td>
<td>8.8 ± 1.83*</td>
</tr>
<tr>
<td>Irradiated</td>
<td>2.4 ± 1.7</td>
<td>0.2 ± 0.02*</td>
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In the unirradiated antibody-treated conditions, recipients received 50 μg antibody intravenously with 7×10^4 donor CD4+ T cells. Accumulation of donor T cells in spleen was assessed 5 days later (n = 3 per group); data are representative of 3 independent experiments. In the irradiated condition, recipients received 6 Gy γ irradiation approximately 3 hours prior to transfer of 1×10^6 donor CD4+ T cells, and accumulation of donor T cells in spleen was assessed 4 weeks later (n = 6 per group). All values (mean ± SD) represent the number of donor cells recovered × 10^6. *P < 0.05 versus aly/+ control.
transferred into irradiated H-2<sup>bm12</sup> recipients (23), and we confirmed that naive Ox40<sup>−/−</sup> CD4<sup>+</sup> T cells did not cause lethal GVHD in our similar model (Supplemental Figure 2). Therefore, we considered the possibility that the inability of NIK-deficient T cells to mediate lethal GVHD might be owing to a requirement for NIK downstream of OX40 and perhaps other TNFRs.

To test this hypothesis, we used a model of GVHD previously described by our lab, in which costimulation by agonist anti-OX40 antibody produces lethal GVHD in unirradiated (H-2<sup>bm12</sup>×CD45.1)F1 recipients of B6 CD4<sup>+</sup> T cells. Without irradiation of recipients and without anti-OX40, donor CD4<sup>+</sup> T cells proliferate and accumulate, but are rendered anergic and do not acquire effector function (24), because in the absence of irradiation damage and inflammation, costimulatory molecules are not upregulated (22). The OX40 stimulus causes alloreactive donor T cells to differentiate to IFN-γ producers. It acts directly on donor cells, as shown by lack of a response by Ox40<sup>−/−</sup> donor T cells transferred alone into WT (H-2<sup>bm12</sup>×CD45.1)F1 recipients (24). We tested naive aly/aly or littermate control donor CD4<sup>+</sup> T cells in this model to assess whether NIK is essential for OX40-mediated costimulation. When naive CD4<sup>+</sup> T cells were cotransferred with control antibody and assayed 5 days later, aly/aly and control CD4<sup>+</sup> T cells behaved identically: both proliferated more than 7 times, as assessed by complete CFSE dilution (data not shown); both expanded to similar numbers (Table 1); and neither upregulated CD25 or produced IFN-γ upon ex vivo stimulation (Figure 3). In contrast, coadministration of agonist anti-OX40 antibody had very disparate effects on control versus aly/aly donor T cells. In recipients of control CD4<sup>+</sup> T cells, anti-OX40 increased cell accumulation 3-fold (Table 1), induced massive CD25 upregulation (Figure 3A), and induced effector differentiation such that donor cells produced copious IFN-γ upon ex vivo stimulation with IL-12 and IL-18 or with PMA and ionomycin (Figure 3, B and C). Moreover, approximately 20% of control T cells made IFN-γ even without stimulation (Figure 3, B and C). In contrast, anti-OX40 administration had almost no effect on aly/aly T cells, as assessed by cell numbers, CD25 upregulation, and IFN-γ production (Table 1 and Figure 3). We conclude that NIK is essential for OX40 to costimulate and impart a lethal phenotype to responding alloreactive CD4<sup>+</sup> T cells.

Because NIK is required for normal thymic epithelial cell development, we more stringently tested the T cell–intrinsic role of NIK using aly/aly or control T cells obtained from BM chimeras in which the BM recipients (and thus thymic stroma) were WT. aly/aly T cells that developed in a NIK sufficient thymus also failed to respond to costimulation through OX40 by upregulating CD25 or acquiring the capacity to secrete IFN-γ (Supplemental Figure 3).

In addition to costimulating effector T cells, OX40 may also augment immune responses in some situations by inhibiting Treg differentiation (25–27). We tested whether this function of OX40 also relies on signaling through NIK by inducing Treg differentiation in vitro in the presence or absence of agonist anti-OX40 antibody. OX40 inhibited induced Treg (iTreg) differentiation by approximately 40% in NIK-sufficient CD4<sup>+</sup> T cells, whereas it had no effect on iTreg induction in aly/aly CD4<sup>+</sup> T cells (Supplemental Figure 4).

OX40 activates noncanonical NF-κB in a NIK-dependent manner. NIK is necessary to activate the noncanonical NF-κB pathway downstream of a limited number of TNFRs, including BAFFR, LTBR, and CD40. To assess whether OX40 also depends on NIK to activate the noncanonical NF-κB pathway, we examined canonical and noncanonical NF-κB signaling downstream of OX40 in aly/aly and littermate control T cells. We stimulated purified naive aly/aly or control CD4<sup>+</sup> T cells with anti-CD3 and anti-CD28, and after 48 hours we added control antibody or agonist anti-OX40 for 5 minutes or 20 hours to assess NF-κB activation. Stimulation with anti-OX40 for 5 minutes induced equivalent IkBα phosphorylation in T cells of both genotypes (Figure 4A), which indicates that OX40
initiates robust canonical NF-κB signaling, as previously reported (28), and that this pathway is NIK independent.

We examined the effect of OX40 ligation on noncanonical NF-κB activation by assessing p100 processing in whole-cell extracts. Naive CD4+ T cells contained low levels of p100 (Figure 4B). As previously reported (16), naive aly/aly T cells expressed less p100 than control T cells, but by 48 hours after TCR stimulation, p100 levels were the same between genotypes (Figure 4B and Supplemental Figure S5). Although p100 expression increased upon TCR stimulation in both control and aly/aly T cells, processing to p52 was minimal at 24 and 48 hours; 20 hours later (i.e., 68 hours of TCR stimulation), the p52/p100 ratio had increased in both control and aly/aly cultures (Figure 4B), showing that after 2–3 days of in vitro stimulation, CD4+ T cells processed p100 in a partially NIK-independent manner. However, agonist anti-OX40 antibody augmented p100 processing in control T cells, but had no effect on aly/aly T cells (Figure 4B). Thus, although TCR stimulation alone induced both NIK-dependent and NIK-independent p100 processing, OX40 further increased p100 processing, and this effect was entirely NIK-dependent.
To assess downstream events in the NF-κB pathway, we performed EMSA on nuclear extracts from OX40-stimulated *aly*/*aly* and control cells. We found maximal DNA binding activity 48 hours after TCR stimulation, which then declined by 20 hours later with control antibody (Figure 4C, top). Anti-OX40 treatment maintained NF-κB binding, and this effect was similar between control and *aly*/*aly* T cells. This reflects the dominant contribution of canonical NF-κB activation downstream of the TCR and OX40. Supershift analysis using anti–c-Rel antibody or combined anti–c-Rel and anti-RelA antibodies allowed discrimination of discrete RelA and RelB bands, indicative of canonical and noncanonical NF-κB signaling, respectively. OX40 stimulation upregulated RelA DNA binding activity in both *aly*/*aly* and control T cells (Figure 4C, middle). In contrast, OX40 stimulation upregulated RelB DNA-binding activity only in control, but not in *aly*/*aly*, CD4⁺ T cells (Figure 4C, middle and bottom). Together, these data showed that OX40 activated both canonical and noncanonical NF-κB pathways in T cells, but only activation of the noncanonical NF-κB pathway required NIK. By extension, these data indicate that the seemingly minor contribution of RelB to total NF-κB activity may be essential to the function of OX40 in vivo.

NIK overexpression in T cells causes lethal T cell–mediated inflammation and iTreg-mediated suppression. Given the essential role for NIK in our model of OX40-induced GVHD, we asked whether overexpression of NIK in the absence of OX40 or other TNFR costimulation was sufficient to recapitulate this effect. To this end, we obtained conditional NIK transgenic mice in which a single copy NIK-IRES-GFP transgene preceded by a loxP-flanked stop cassette is inserted into the ubiquitously expressed ROSA-26 locus (referred to herein as NIKtg mice; ref. 29). We crossed NIKtg mice with CD4-Cre transgenic mice — generating mice referred to herein as CD4-NIKtg — in order to turn on expression of the NIK transgene specifically in T cells, as confirmed by flow cytometry (Supplemental Figure 6). Expression of the NIK transgene in T cells caused an upregulation of NF-κB2 (p100 plus p52) and an increase in processing of p100 to p52 that was similar to the level of p100 processing seen in WT B cells (Supplemental Figure 7). Unexpectedly, CD4-NIKtg mice died before weaning. They were
Overexpression of NIK in T cells causes lethal inflammation. (A and B) CD4-NIKtg and age-matched littermate WT control mice were weighed and euthanized when moribund. Data in A show average weights (± SD) from 2 combined litters. (C) Sections of colon and lungs from moribund CD4-NIKtg and age-matched littermate WT control mice were assessed for leukocyte infiltrates by hematoxylin-eosin stain. Scale bars: 100 μm. Data are from 2 combined litters.

Born at a normal Mendelian frequency of 50%, but began to lose weight rapidly around day 20 (Figure 5A), developed loose stools, and quickly became moribund, requiring euthanasia between days 23 and 30 (Figure 5B). The lungs of all moribund CD4-NIKtg mice had moderate to massive lymphocytic infiltration (7 of 7), as did the colons of 6 of 7 CD4-NIKtg mice (Figure 5C). Age-matched littermate control organs were all normal (4 of 4).

The rapid-onset mortality accompanied by lymphocytic infiltration of multiple organs that we observed in CD4-NIKtg mice was similar to the phenotype of Foxp3−/− and Foxp3+null mouse, which harbor an inactivating mutation in Foxp3 (30). These mice die of T cell–mediated autoimmunity resulting from lack of regulation by Foxp3+ Tregs. To determine whether NIK overexpression interferes with Treg and/or effector T cell development, we assessed lymphocyte populations from CD4-NIKtg mice at day 11, before onset of outward disease symptoms. Spleen and thymus cellularity were similar between CD4-NIKtg and control littersmates (Supplemental Figure 8), but peripheral and mesenteric LNs were noticeably larger (data not shown). The proportion of CD8+ T cells was slightly lower in LN and spleen of CD4-NIKtg mice, and the proportion of total CD4+ T cells was the same in spleen and LN and slightly lower in the thymus of CD4-NIKtg mice (Supplemental Figure 8). To our surprise, when we examined CD4+ T cells more closely, we found that a significantly greater percentage of CD4+ T cells was Foxp3+ in CD4-NIKtg lymphoid organs compared with lymphoid organs of control littersmates (Figure 6, A and C). Despite this large increase of Foxp3+ T cells, the conventional T cell subsets (CD8+ and Foxp3− CD4+) showed an activated phenotype, as assessed by a 2- to 3-fold increase in the percentage of CD44hi cells (Figure 6, B and D, and data not shown), which suggests they are poorly regulated by Tregs. In CD4-NIKtg mice, both Foxp3+ and Foxp3− T cell populations overexpress NIK, so it was unclear whether CD4-NIKtg Tregs are poor regulators or whether CD4-NIKtg conventional T cells (Tconvs) are resistant to regulation. CD4-NIKtg Foxp3−/CD4+ T cells had significantly lower CD44 and Foxp3 expression compared with control littersmates (Figure 6, E and F), consistent with poorer Treg function (31). To directly address this possibility, we performed in vitro Treg suppression assays using combinations of WT and CD4-NIKtg Tregs and Tconvs. In this assay, WT Tregs inhibited the ability of WT Tconvs to proliferate in response to polyclonal stimulation with anti-CD3 and APCs. While WT Tregs efficiently suppressed WT Tconvs, showing suppression at a 1:18 Treg/Tconv ratio, CD4-NIKtg Tregs were incapable of suppressing WT Tconvs, even at a 1:1 Treg/Tconv ratio (Figure 6, G and H). In addition, CD4-NIKtg Tconvs were difficult to suppress, showing only modest regulation by WT Tregs, even at the highest 1:2 Treg/Tconv ratio (Figure 6, G and I). This suggested that overexpression of NIK both increases Tconv resistance to Treg-mediated suppression and impairs Treg function.

Despite in vitro resistance to Treg-mediated suppression, CD4-NIKtg Tconvs were suppressible in vivo, as assessed by BM chimeras. Lethally irradiated CD45.1 recipients of a mixture of CD4-NIKtg and congenically marked WT BM remained healthy for at least 10 months and did not show signs of inflammation such as enlarged LNs. The same was true for recipients of CD4-NIKtg BM alone, due to radioreistant Tregs that reconstitute the host after irradiation and have previously been shown to prevent disease after transplantation of Foxp3+BM (32). The mixed BM chimeras demonstrated that the CD4-NIKtg Tconv and Treg phenotypes are intrinsic rather than secondary to inflammation, because although the mice remained healthy, (a) the CD4-NIKtg Tconvs had an expanded population of CD44hi cells; (b) WT Tregs were expanded relative to CD4-NIKtg Tregs; and (c) NIKtg Tregs showed decreased CD44 expression, a phenotype consistent with poor regulatory capacity (Supplemental Figure 9).

These chimeras also allowed us to obtain CD4-NIKtg Tregs from healthy mice to stringently test the Treg-intrinsic role of NIK. The decreased suppressive capacity of CD4-NIKtg Tregs from 11-day-old CD4-NIKtg mice could have been secondary to an inflammatory environment in these prediseased mice, rather than an intrinsic effect of NIK in Tregs. We sorted CD4-NIKtg and WT Tregs and Tconvs from healthy BM chimeras and performed in vitro Treg suppression assays using various combinations of Treg and Tconv cell populations. The results of this experiment were similar to those obtained using 11-day-old CD4-NIKtg mice: CD4-NIKtg Tregs from healthy BM chimeras were poorer suppressors...
of WT Tcons than were WT Tregs, and CD4-NIKtg Tcons were more resistant to suppression by WT Tregs than were WT Tcons (Figure 7). However, CD4-NIKtg Tregs from the healthy mice were only modestly impaired in suppressive capacity, which suggests that both Treg-intrinsic and inflammatory phenomena contribute to the demise of Treg function in CD4-NIKtg mice.

These in vitro data suggested that overexpression of NIK in T cells causes aggressive lethal autoimmunity via intrinsic effects on both Tcons and Tregs; combined, these effects lead to hyperactive effector T cells. Furthermore, the observation that lethally irradiated CD45.1 WT recipients of CD4-NIKtg BM (which harbor radioresistant WT host Tregs) remained healthy, but Rag1−/− recipients (which lack host Tregs) succumbed to disease (data not shown), suggested that WT Tregs can regulate CD4-NIKtg effector T cells and, by extension, that CD4-NIKtg Tregs cannot. To formally test this, we made mixed BM chimeras in which 1:1 mixtures of CD4-NIKtg plus Foxp3scurfy BM or CD4-NIKtg plus WT (FoxP3scurfy littermate) BM were used to reconstitute Rag1−/− recipients. In this experiment, the sole difference between the groups was the absence or presence of WT Foxp3+ Tregs. We found that 80% of the CD4-NIKtg plus WT BM recipients survived, but nearly all of the CD4-NIKtg plus FoxP3scurfy BM recipients succumbed to disease within 8 weeks (Supplemental Figure 10). We thus concluded that CD4-NIKtg Tregs have poor regulatory function in vivo.

In our genetic systems of NIK induction, T cells overexpressed NIK from the double-positive thymocyte stage. We wondered whether...
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Acute NIK induction in mature Tregs would have the same effect, so we developed a system to induce NIK expression in vitro using TAT-Cre fusion protein transduction, in which we attained approximately 50%–70% transduction efficiency. We treated purified NIKtg or control CD4+ T cells with TAT-Cre, cultured under Treg-inducing conditions, and then sorted iTregs from control cultures and GFP+ iTregs from NIKtg cultures. NIK overexpression did not interfere with iTreg differentiation in these conditions, as assessed by Foxp3 and CD25 expression (Figure 8A and data not shown). However, when we tested these sorted iTregs for suppressive function against WT Tconv, we found that iTregs in which NIK was acutely expressed were much poorer suppressors than TAT-Cre–treated control iTregs (Figure 8, B and C). This showed that NIK overexpression affected Treg function independently of developmental effects that could occur during selection in the thymus.

Discussion

NIK has been studied primarily in terms of its role in immune system organogenesis, lymphoid architecture, and naive B cell survival. Here, we identify a role for NIK as a critical mediator of T cell responses following antigen exposure by signaling downstream of OX40 and probably other costimulatory TNFRs. We found that NIK was absolutely required for lethal CD4–mediated GVHD, but not for initial naive T cell responses through the antigen receptor in vitro. Likewise, in unirradiated recipients without exogenous costimulation, NIK-deficient T cells expanded normally.

Recently, Hofmann et al. showed that NIK is important for DCs to prime effective T cell responses (18). They found that whereas single NIK−/− BM chimeras are resistant to EAE, single NIK+/− BM chimeras and mixed NIK−/− Rag−/− BM chimeras (which harbor WT DCs) are susceptible to EAE. However, it is unclear in this experiment whether there is also a role for NIK in the T cells, since the authors compared the mixed chimeras with single NIK+/− BM chimeras whose DCs are not WT, but heterozygous for NIK. We and others have shown that NIK heterozygosity confers a significant in vivo effect in various cell types, and it is likely that NIK+/− DCs are also at least partially impaired. Indeed, similar to Jin et al. (19), Hofmann et al. also showed that purified NIK KO T cells transferred into WT hosts (with WT DCs) could not mediate EAE (18). Although NIK is clearly important in DCs and other APCs to elicit optimal T cell responses, in our experiments, we carefully purified T cells to be able to identify cell–intrinsic effects of NIK in vivo and in vitro. Furthermore, we identified OX40 as a specific receptor expressed by T cells that relies absolutely on NIK to mediate its T cell costimulatory effects.

Several lines of evidence suggested to us that NIK might act downstream of costimulatory TNF receptors on T cells. First, NIK intrinsically inhibits Treg-suppressive capacity and increases Tconv resistance to suppression. CD4-NIKtg and littermate WT control Tregs and naive T cells were sorted from spleen and LN of healthy BM chimeras and cultured with CD45.1 Tregs or naive T cells plus WT APCs and soluble anti-CD3. (A) Naive T cells were labeled with CellTrace Violet to assess division over the course of 3 days. Gates indicate proportion of T cells that underwent at least 1 cell division. (B and C) Data (± SD) are representative of 2 independent experiments.
known receptors that rely on NIK to activate the noncanonical NF-κB pathway. Second, several of the TNF receptors expressed on T cells have previously been shown to induce constitutive p100 processing when overexpressed in cell lines (33), and OX40-induced NF-κB activation in HEK293 cells was inhibited by transduction with dominant-negative NIK (34). TNFRII on human primary T cells was recently shown to induce p100 processing (35). Finally, the T cell phenotypes of NIK-deficient mice are surprisingly similar to those of OX40−/− mice. T cell development and peripheral populations are near normal in OX40−/− mice, as are most in vitro responses and initial clonal expansion upon antigen recognition in vivo. However, maximum T cell expansion and differentiation to effector function depend on OX40/OX40L interactions in vivo (3). Particularly relevant to our studies, OX40−/− and Ox40−/− mice suffer much milder disease in models of autoimmunity and GVHD (3). When we tested the role of NIK in OX40-mediated GVHD, we found that the costimulatory effects of OX40 activation were almost completely abrogated in NIK-deficient T cells, demonstrating a direct requirement for NIK downstream of OX40. Likewise, the ability of OX40 to inhibit iTreg induction was abolished in NIK-deficient T cells. Although we cannot rule out other possible roles for NIK in the CD4+ T cell response to OX40 signaling, we believe sustained activation of NF-κB through the noncanonical pathway may be necessary. In support of this concept, we showed that OX40 activated the noncanonical NF-κB pathway in primary CD4+ T cells, and that this was NIK dependent.

It is likely that OX40 is not unique among T cell costimulatory TNF receptors in using NIK to activate the noncanonical NF-κB pathway. Jin et al. showed that costimulatory effects of LIGHT in vitro depend on part on NIK (19), and, as mentioned above, activation of TNFRII can induce p100 processing in T cells (35). Other likely candidates include CD30 and 4-1BB, both of which contribute to GVHD (22, 23), and both of which have been shown to induce p100 processing when overexpressed in cell lines (33).

It is somewhat surprising that NIK deficiency had such a profound effect on T cell responses to OX40, given that canonical NF-κB activation was preserved and that the noncanonical pathway seemed to make a minor contribution to overall NF-κB activity (Figure 4). However, it should be noted that in other cell types, small differences in noncanonical NF-κB activation have profound biological consequences. In conditional NIK transgenic B cells, for example, processing of p100 to p52 is increased only 30% above WT B cell levels, yet peripheral B cell numbers are increased 3-fold (29). It may be that specific OX40 target genes depend on p52: RelB dimers to induce their transcription, as has been shown for LTβR target genes (36). Another possibility, not mutually exclusive from the former, is that NIK-driven activation of the noncanonical NF-κB pathway may prolong NF-κB signaling via degradation of p100, which is induced by NF-κB activation and acts like an IκB to inhibit both canonical and noncanonical NF-κB activation (37, 38). This might explain how small differences in the level of NIK (i.e., a 2-fold decrease in aly/+ heterozygous mice) could mediate large physiologic effects: NIK-induced degradation of p100 could simultaneously activate the noncanonical NF-κB pathway and relieve a brake on both NF-κB pathways.

Likewise, we found that a modest increase in NIK expression in T cells exerted profound physiological effects. The conditional NIK transgene induced p100 processing in T cells similar to that in WT resting B cells, yet T cells became highly activated, and mice succumbed to rapid fatal autoimmunity. Surprisingly, this occurred despite increased numbers of Foxp3+ Tregs. NIK overexpression...
affected both Tconvs and Tregs, with the combined result being overactivated effector T cells that were not controlled by Tregs. WT Tregs could control the disease mediated by CD4-NIKtg T cells in vivo, whereas NIKtg Tregs could not. This suggests that NIK overexpression in Tregs is critical for disease induction, but does not rule out a role for NIK in Tconvs. In particular, even in healthy mixed BM chimeras, NIKtg Tconvs retained an activated phenotype and remained difficult to suppress in vitro. Thus, we believe that NIK overexpression in both Tconvs and Tregs contributes to the rapid-onset autoimmunity observed in CD4-NIKtg mice.

 Constitutive activation of the noncanonical NF-κB pathway via a ubiquitously expressed p52 transgene also predisposes mice to multiquor organ autoimmunity, and the p52 transgenic T cells are more activated and more resistant to apoptosis (39). Moreover, although p52 transgenic mice do not suffer early postnatal mortality, normal or transgenic expression of p52 in the absence of the inhibitory p100 subunit leads to a lethal inflammatory syndrome, although the role of T cells in mediating autoimmunity was not tested (40, 41). The exacerbated disease caused by the absence of p100 indicates that NIK overexpression may have provided a powerful T cell stimulus in our experiments by relieving p100-mediated NF-κB inhibition as well as by increasing p52-mediated gene transcription.

 In many ways, NIK overexpression in T cells mimicked constitutive OX40 activation, which induces an activated phenotype, enhances cell survival, and, importantly, confers resistance to Treg-mediated suppression in Tconvs (3). OX40 ligation not only enhances T cell responses to foreign antigens, tumor antigens, and alloantigens, but also can exacerbate autoimmune diseases when used as an adjuvant along with self-antigen. Interestingly, OX40L transgenic mice develop a T cell–mediated autoimmune syndrome characterized by a large population of CD44hi memory CD4+ T cells and colon and lung leukocyte infiltration reminiscent of our CD4-NIKtg mice, but with delayed kinetics (42). The phenotype of CD4-NIKtg Tregs also paralleled observed effects of OX40 on this cell type. Numerous studies have shown that activation of OX40 inhibits the suppressive function of Tregs (25–27). Although some studies have also shown that OX40 signaling increases Treg-mediated suppression, this appears to result from positive effects of OX40 on Treg homeostasis and expansion, such that lack of OX40 in vivo reduces Treg numbers, rather than affecting Treg function on a per-cell basis (43, 44). Additionally, 2 other TNFR ligands that inhibit Treg function may rely, at least in part, on NIK for signaling. Glucocorticoid-induced TNF receptor family related protein ligand (GITRL) and TNF-α both have been shown to activate NF-κB in Tregs and to inhibit Treg function (45, 46), and as mentioned above, activation of TNFRII induces modest p100 processing in human Tconvs (35). Curiously, NIK KO Tregs are hyperproliferative upon GITR ligation, which may be caused by a disproportionate increase of cells in the Treg subset that possess an activated/memory phenotype (47). This is suggestive of complex effects of NIK in Tregs that may depend on activation of particular receptors and/or integration with other signals.

 How does upregulation of NIK interfere with the Treg program? Comparing the essentially nonfunctional CD4+NIKtg Tregs from predeceased mice (Figure 6) with the moderately impaired CD4-NIKtg Tregs from healthy chimeras (Figure 7) suggests that NIK overexpression impairs Treg function sufficiently to allow Tconvs to escape suppression, and the ensuing inflammatory environment further degrades Treg function in a positive feedback loop. A similar phenomenon occurs when Dicer is conditionally deleted in Foxp3+ Tregs to block microRNA production. Those mice develop a severe fatal autoimmune syndrome similar to that observed in our CD4-NIKtg mice (48). Foxp3+ Tregs are expanded, but nonfunctional. Like CD4-NIKtg Tregs, the Dicer−/− Tregs are functionless in the inflammatory environment of the sick males, but retain some function in healthy heterozygous females. Together, these data show that diverse mechanisms are required to maintain optimal Treg function, and modest Treg destabilization can easily tip the balance to frank autoimmunity.

 Our data suggest that manipulating levels of NIK in T cells may have therapeutic value. While a large number of NF-κB blocking compounds are in clinical development and trials, we are aware of none shown to be specific for NIK or the noncanonical NF-κB pathway. Decreasing NIK activity in T cells might significantly ameliorate autoimmune and alloresponses, like GVHD and transplant rejection, without crippling the immune system as severely as do inhibitors of canonical NF-κB activation. The fact that even heterozygous aly+/− mice were partially protected from lethal GVHD is promising, as it suggests complete ablation of NIK is not necessary for drug efficacy. Moreover, GVHD is ameliorated when the noncanonical NF-κB subunit, RelB, is eliminated in APCs (49). Therefore, combined inhibition of the noncanonical NF-κB pathway in T cells and APCs might be a particularly effective therapy. Conversely, upregulating NIK in DCs has been suggested as a potential therapeutic to increase vaccine efficacy (50). We showed here that upregulation of NIK provided a strong activation stimulus in T cells as well. As above, this suggests that combined modulation of NIK in T cells and DCs could have additive or synergistic effects, particularly in the setting of cancer immunotherapy, in which a major obstacle is overcoming Treg-mediated suppression of immune responses. However, caution clearly needs to be employed in developing therapies aimed at increasing NIK, since the potential for devastating autoimmunity and lymphomagenesis is present when the noncanonical NF-κB pathway is not tightly regulated (29, 39, 51, 52).

Methods

Mice. Mice were housed under specific pathogen–free conditions at the Oregon Health and Science University animal facility. B6.Thy.1, B6.CD45.1, FoxP3tm1ef, and Rag2−/− mice were from The Jackson Laboratory. (H-2b×H-2d)F1 mice were made by crossing B6.C-H2b×d/Ktheg mice (The Jackson Laboratory) to B6.CD45.1 mice. Aly/NscJcl mice (CLEA Japan) were maintained by breeding female aly/+ to male aly/aly mice. Offspring were genotyped by assessing serum IgA levels as described previously (6). The aly+/− mice, with 2 normal copies of NIK on the aly background, were made by aly/+ to aly+/+ crosses and were genotyped by PCR. ROSA26StopNik transgenic mice were provided by K. Rajewsky (Harvard Medical School, Boston, Massachusetts, USA) and M. Schilde-Supprian (Max Plank Institute of Biochemistry, Munich, Germany), and mice heterozygous for the NIK transgene were bred to Cd4-Cre homozygous transgenic mice (53) (Taconic). Female FoxP3tm1ef mice were from The Jackson Laboratory. Mice were housed under specific pathogen–free conditions at The Jackson Laboratory. Rag2−/− mice were from Taconic. Male FoxP3tm1ef mice were bred to male C57BL/6 mice. Male FoxP3tm1ef offspring were identified phenotypically and by FACs analysis of T cells. Primers used for genotyping aly/aly mice were 5′-GACATCCCGAGCTACTTCAACA-3′ and 5′-GGTGTCTTTACCTTTCTCATCCACCA-3′. Primers used for genotyping CD4-NIKtg mice (EGFP amplification) were 5′-ATGAAAGCGAGCACTTCTCAA-G-3′ and 5′-GGCGGATTTGAGGTCTTGATGCGG-3′.

Antibodies. Rat anti-OX40 (clone OX86), anti–IL-4, and anti–IFN-γ blocking antibodies were from BioXCell. Control rat IgG was from Capell, ICN Pharmaceuticals. Anti-CD3 was from eBiosciences, and anti-CD28 was from Biolegend. Rabbit anti–phospho-IκBε was from Cell Signaling Technology.
rabbit anti-actin and rabbit anti-Erk2 were from Santa Cruz Biotechnology, rabbit anti-mouse-p100/p52 was a gift from S. Ley (National Institute for Medical Research, London, United Kingdom), and HRP-conjugated goat anti-rabbit was from BD Biosciences. Fluorescently conjugated antibodies used for FACS were anti-CD44-FITC, anti-CD4-PerCP-Cy5.5 or −PE-Cy7, anti-CD8-PE-Cy7, anti-CD45.1-PE or −PE-Cy7, anti-CD25-PE or −biotin, anti-IFN-γ-allophycocyanin, anti-CD69-FITC, anti-Foxp3-Alexa Fluor 647, anti-CD3-biotin, polyclonal chicken anti-OKT40-biotin, and streptavi- din-allophycocyanin. All staining antibodies were from eBioscences except anti-CD25-biotin (BD Biosciences) and chicken anti-OKT40 (gift from A. Weinberg,Earle A. Chiles Research Institute, Portland, Oregon, USA).

Flow cytometry. Spleen cell suspensions were stained directly ex vivo or stimulated for 5 hours and then stained for surface markers and intracellular cytokines as previously described (54). Foxp3 staining was performed per the manufacturer’s instructions (Biolegend Foxp3 Fix/Perm intracellular cytokines as previously described (54). Foxp3 staining was anti-CD25–biotin (BD Biosciences) and chicken anti-OX40 (gift from S. Ley) and analyzed using FlowJo (Tree Star).

Adoptive transfers. Donor lymphocyte single-cell suspensions were prepared from spleen, and total CD4+ T cells were purified by magnetic negative selection (EasySep; Stem Cell Technologies). Naive CD4+ T cells (CD4+CD25+CD44lo/mid) were then sorted on a FACSVantage (BD Biosciences) to greater than 98% purity. In some experiments, cells were CFSE labeled prior to transfer. 1 × 10^6 cells were injected intravenously into lethally irradiated (12 Gy, split dose) CD45.1 recipients or sublethally irradiated (6 Gy, single dose) Rag1−/− recipients. For CD4−NIKtg plus Foxp3−/− and CD4−NIKtg plus WT BM chimeras, CD4−NIKtg BM was picked in Rag1−/− mice for 3 weeks prior to making the mixed BM chimeras. This was due to difficulty in precisely timing B6Foxp3−/− congenic and CD4−Cre+NIKtg pregnancies.

Histology. Mouse organs were fixed with 10% (v/v) buffered formalin and embedded in paraffin. Sections were stained with hematoxylin and eosin.

Statistics. The significance of differences between groups was calculated using 2-tailed Student’s t test. Log-rank test was used to assess differences in survival between aly genotypes. Differences with a P value less than 0.05 were considered statistically significant.

Study approval. All protocols in the present studies were reviewed and approved by the Institutional Animal Care and Use Committee of Oregon Health and Science University.

Acknowledgments

We thank A. Weinberg and S. Ley for anti-OX40 and anti-p100 antibodies, respectively; K. Rajewsky and M. Schmickl-Supprian for NIKtg mice; M. Boyd for cell sorting; P. Stork and M. Takaguchi for scurfy mice, or WT control litters of Foxp3−/− mice or from femurs and tibias of adult aly/a ly mice or +/+ mice. Single-cell suspensions of BM were depleted of mature T cells via magnetic separation using anti-CD3–biotin. 0.25–1.0 × 10^6 BM cells were injected intravenously into lethally irradiated (12 Gy, split dose) CD45.1 recipients or sublethally irradiated (6 Gy, single dose) Rag1−/− recipients. For CD4−NIKtg plus Foxp3−/− and CD4−NIKtg plus WT BM chimeras, CD4−NIKtg BM was picked in Rag1−/− mice for 3 weeks prior to making the mixed BM chimeras. This was due to difficulty in precisely timing B6Foxp3−/− and CD4−Cre+NIKtg pregnancies.

Received for publication August 20, 2010, and accepted in revised form September 21, 2011.

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