Role of Nr13 in regulation of programmed cell death in the bursa of Fabricius

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Apoptotic cell death is developmentally regulated in the chicken bursa of Fabricius. Although apoptosis is low in the embryonic bursa, cell death increases markedly after hatching. The expression of Bcl2 family cell death antagonists was examined to identify the genes that regulate bursal cell apoptosis. The expression of Bcl-xL, A1, and Mcl1 was detected in both embryos and hatched birds, whereas Nr13 was expressed at high levels in embryonic bursa, and decreased significantly after hatching, correlating inversely with apoptosis. The oncogene v-rel and phorbol myristate acetate, two known inhibitors of bursal cell apoptosis, induced Nr13 expression. Overexpression of Nr13 in DT40 bursal lymphoma cells protected them from low serum-induced apoptosis. The mechanism of inhibition of apoptosis by Nr13 is likely to involve a critical BH4 domain and interaction with death agonist Bax. Deletion of the BH4 domain converted Nr13 into a death agonist. Bax coimmunoprecipitated with Nr13 and Bax was induced, whereas Nr13 levels diminished when bursal lymphoblasts were induced to apoptosis by dispersion. Bursal transplantation studies demonstrated that Nr13 could prevent the in vivo programmed elimination of bursal stem cells after hatching, suggesting that Nr13 plays a role in maintaining bursal stem cells.

[Key Words: Nr13; apoptosis; bursa; transplant; stem cells]

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Results

Expression of antagonists of apoptosis in bursal development

The expression of Bcl2 members of death antagonists was first investigated to determine which member of the family correlates with bursal cell survival. We analyzed expression of other death antagonists in the Bcl2 family, including Bcl-xL, Nr13, A1, and Mcl1. Both A1 and mcl1 are hematopoiesis-related genes able to protect hematopoietic cells from interleukin-3 withdrawal-induced apoptosis [Lin et al. 1996; Zhou et al. 1997]. Nr13 is a relatively newly identified member of the Bcl2 family, initially identified as a v-src-activated gene in developing neural tissues, with a pattern of membrane localization similar to that of Bcl2 [Gillett et al. 1995]. Ectopic overexpression of quail Nr13 protects cultured mammalian hematopoietic cells against factor withdrawal-induced apoptosis [Mangenev et al. 1996].

The expression of these four genes was analyzed by Northern analysis. Nr13 RNA was present in bursal follicles at days 15, 17, and 21 of embryogenesis, but was no longer present 28 days after hatching [Fig. 1]. Mcl1 RNA was present in bursal follicles at days 15, 17, and 21 of embryogenesis, but was no longer present 28 days after hatching. Northern analysis. Nr13 RNA was present in bursal follicles after hatching [Mangeney et al. 1996]. Apoptosis is particularly prominent in bursal follicles after hatching [Neiman et al. 1994b], and embryonic and post-hatching bursal lymphocytes can be induced to undergo apoptosis by loss of cell contact [Neiman et al. 1991].

Embryonic bursal follicles contain a transplantable stem cell compartment, comprised of 1% or less of the bursal population [Toivanen and Toivanen 1973; Pink et al. 1985], which is also subject to programmed elimination after hatching. It has been demonstrated that there are no transplantable bursal stem cells left in the bursa by 4 weeks after hatching [Thompson and Neiman 1987].

We investigated the expression of bcl2 family genes during B-cell development in the bursa to identify regulators of this extensive cell death activity in the bursa. Previous studies [Eguchi et al. 1992] and our own unpublished results indicated that bcl2 itself was expressed at a very low level, if at all, in the bursa. Therefore, we examined the expression of other family members. Our results suggest that Nr13 is an important protein for bursal B-cell viability and programmed survival of bursal stem cells.

Induction of apoptosis in bursal cells by dispersion causes an increase of Bax and a decrease of Nr13

Bursal cells require cell–cell interaction to maintain their viability. When embryonic bursal cells are dispersed, they rapidly undergo apoptosis, and can become terminal deoxynucleotide end-labeling (TUNEL) positive within an hour [Neiman et al. 1991]. We dispersed day 21 embryonic bursal cells by passing through nylon mesh as described [Neiman et al. 1991] to trigger apoptosis. Proteins were extracted at different time points after dispersion for Western blot analysis. The levels of Nr13 decreased by 2 hr after dispersion (Fig. 2a). The levels of Bax, a death agonist, were analyzed by Western blot, and the Bax levels increased after 1 hr of dispersion [not shown]. As shown in Figure 2b, we repeated this analysis to study the kinetics of Bax increase. Bax was not detectable in undispersed bursa, but appeared by 30 min after bursal cell dispersion and continued to increase for at least 2 hr. Although Bax was reported to be a transcriptional activation target of p53 [Miyashita and Reed 1995], this induction of Bax with dispersion of bursal cells is probably independent of p53, because p53 pro-
tein and mRNA levels are not altered by the dispersion process (Neiman et al. 1994a). We conclude that the ratio of Nr13 to Bax decreases dramatically after dispersion of bursal lymphocytes.

**Induction of Nr13 by v-rel and phorbol ester, two known inhibitors of bursal apoptosis**

Phorbol myristate acetate (PMA) is known transiently to inhibit dispersion-induced bursal apoptosis (Asakawa et al. 1993; Compton and Waldrip 1998). We tested a chicken bursal lymphoma-derived cell line, DT40 (Baba et al. 1985), to determine whether PMA could induce Nr13 expression. Nr13 RNA increased within 1 hr of PMA treatment and continued to increase at least until 6 hr (Fig. 3a). PMA induction of Nr13 was also observed in dispersed embryonic bursal cells (Fig. 3b). These results suggest that Nr13 could be an intermediate in PMA-induced protection from cell death.

Another known inhibitor of bursal cell apoptosis is proto-oncogene v-rel, a member of the NF-κB transcription activators. v-rel can transform bursal cells in vivo and block apoptosis induced by bursal dispersion [Neiman et al. 1991; White et al. 1995]. We constructed retroviral vectors by cloning v-rel [White and Gilmore 1993] into the LXSN vector [Fig. 4a] and infected DT40 cells with v-rel vectors to test whether v-rel would affect Nr13 expression. After the cells were selected with G418, the control DT40 cells and the DT40 cells infected with v-rel were incubated at various temperatures to examine their growth and effects on Nr13 expression. The growth rate of DT40 cells was not affected by v-rel at 37°C, 40°C, and 42°C [not shown]. Northern blot analysis demonstrated that Nr13 mRNA increased...
threefold in DT40 cells when the temperature was shifted from 37°C to 42°C (near the physiological body temperature of chicken) (Fig. 4b, lanes 1, 5). Nr13 RNA was only slightly increased at 37°C by v-rel [Fig. 4b, lanes 1, 2] but was significantly enhanced by v-rel at 42°C [Fig. 4b, lanes 5, 6]. At 44°C the growth rate of the DT40 cells was impaired (not shown) and no effect of v-rel on Nr13 RNA was observed. These results suggest that rel-related Nr13 activation may contribute to v-rel-induced inhibition of apoptosis.

Overexpression of Nr13 in DT40 cells protects DT40 cells from serum withdrawal-induced apoptosis

We then tested the DT40 cell line to determine whether increased levels of Nr13 could inhibit apoptosis in these cells. DT40 cells express constitutively high levels of c-myc as a result of retroviral promoter insertion ( Hayward et al. 1981). Like previously reported myc-overexpressing Rat1 cells [Evan et al. 1992], serum deprivation of DT40 cells suppresses net cell growth in the culture as a result of increased apoptosis. We studied whether Nr13 could protect DT40 cells from serum withdrawal-induced apoptosis. As shown in Figure 5a, Nr13 retroviral vectors LNRSN and LNRCG were constructed in pLXSN [neomycin resistance expressing] and pLXCG [green fluorescence protein (GFP) expressing] plasmids, and PG13 packaging lines expressing these viruses were prepared [Miller and Rosman 1989; Miller et al. 1991]. The DT40 cells have a low but detectable level of Nr13 expression. After infection with the LNRSN virus and selection with G418, DT40 cells expressed at least a 5- to 10-fold increase of Nr13 protein (Fig. 5b). A similar increase of Nr13 was observed in DT40 cells infected with LNRCG virus and sorted for GFP-positive cells by flow cytometry (not shown). In standard media [10% bovine calf serum (BCS) and 2% chicken serum (CS)] parental wild-type DT40 cells grew for 3 days and leveled off. Under reduced serum condition (1% CS and no BCS), both wild-type DT40 cells and DT40-overexpressing Nr13 cells were able to grow at similar rate for 2 days. Net growth of wild-type DT40 cells ceased after 2 days, and there was net cell death attributable to apoptosis, whereas the growth of Nr13-overexpressing cells was closer to that of DT40 in normal serum (Fig. 5c). This experiment indicated that Nr13 can protect bursa-derived cells against serum withdrawal-enhanced apoptosis in the presence of elevated Myc.

Potential mechanisms of Nr13 function

Nr13 interacts with Bax  Bax is a death-promoting agonist that interacts physically with Bcl2 and Bcl-xL [Oltvai et al. 1993]. Bax-deficient mice have the phenotype of lymphoid hyperplasia in both T and B lineages [Knudson et al. 1995]. Quantitative analysis in IL-3-dependent FL5.12 cells showed that, when half or more of endogenous Bax was heterodimerized with Bcl2 or Bcl-xL, apoptosis induced by IL-3 withdrawal was repressed (Yang et al. 1995). On the other hand, when Bax was displaced from Bcl2 or Bcl-xL as a result of binding of Bcl2 and Bcl-xL by Bad, another Bcl2 family death agonist, cell death was promoted [Yang et al. 1995]. Therefore, cell survival or death is thought to be dependent on the interaction between Bcl2/Bcl-xL and Bax. Both Bax and Nr13 can be detected in DT40 cells, thus, we investigated whether Bax interacts with Nr13 by coimmunoprecipitation. Cell lysates from DT40 were immunoprecipitated with Nr13 antisera and the immunoprecipitates were analyzed by Western immunoblotting using Bax antisera. Bax was detected in the whole cell lysate, the supernatant of the immunoprecipitate, and the Nr13 immunoprecipitate, but not in control immunoprecipitate with normal rabbit serum (see Fig. 2c). This experiment suggests that a portion of Bax is complexed with Nr13 in DT40 cells, and that Nr13 may control apoptosis in bursal cells by interaction with Bax. When the level of Bax increases relative to the level of Nr13, for example 1 hr after bursal dispersion, bursal cells undergo apoptosis.

BH4 domain of Nr13 is required for protection of cell death and deletion of BH4 domain converts Nr13 into a death agonist  Four conserved homology domains of
the Bcl2 family have been defined: BH1, BH2, BH3 and BH4 (Hanada et al. 1995; Reed 1997). The BH1 and BH2 domains are known to be essential for homodimer or heterodimer formation. BH3 plays the role of ‘ligand’ during heterozygous dimerization of Bcl2 family members. The BH4 domain is an amino-terminal α-helix that appears to play a regulatory role by docking of Raf1 [Wang et al. 1996a], Bag1 [Wang et al. 1996b], calcineurin [Shibasaki et al. 1997], or Ced4/Apaf1 [Chinnaiyan et al. 1997]. It was recently reported that the BH4 domain of Bcl2 has a caspase cleavage site at its carboxy-terminal end that can be cleaved by a caspase, converting Bcl2 into a Bax-like death effector (Cheng et al. 1997). We analyzed the predicted secondary structure of Nr13 and compared it with Bcl2 by NNPREDICT, a protein secondary structure prediction program (Kneller et al. 1990).

We found that Nr13 also has seven predicted α-helices similar to Bcl2 [Fig. 6a]. However, unlike Bcl2 or Bcl-xL, Nr13 has a relatively short loop between the amino-terminal helices. Moreover, no clear sequence homology to the amino-terminal BH4 domain of Bcl2 could be identified in Nr13, and a caspase cleavage site (aspartic acid) was not found. There is, however, an apparent α-helix at the Nr13 amino-terminus that could be the potential counterpart of the BH4 domain. This portion of Nr13 sequence was deleted and tested to determine the function of this BH4-like helix. The Nr13 deletion mutant was cloned into pLXSN to generate a retroviral vector, and DT40 cells were infected and selected for G418 resistance as was previously done with full-length Nr13. Because available antibodies to full-length Nr13 did not react with the deletion mutant protein, the expression of

Figure 6. BH4 domain of Nr13. [a] Predicted secondary structures of chicken Bcl2 and Nr13. The primary sequences and the domain structures are shown at top. The predicted secondary structures are shown on the bottom. [H] helix; [-] no prediction; [E] β strand. [b] Confirmation of expression of the truncated mutant by RNase protection assay. Deletion between ATG initiation codon and NarI site was generated as described in Materials and Methods. The probe for RNase protection assay is shown at left. The first lane of the gel is the marker and the size of each marker is indicated. The second and third lanes are DT40 cells and DT40 cells infected with truncated mutant vectors. [c] Growth curves of DT40 cells and DT40 cells expressing amino-terminal truncated mutant in 1% CS media. (●) DT40 1% CS; (○) DT40-del-Nr 1% CS. Cell count was performed with trypan blue exclusion.
truncated Nr13 was confirmed by a RNase protection assay (Fig. 6b). In this assay, endogenous Nr13 mRNA produced a 190-base RNA fragment, and the retroviral mRNA resulted in a 120-base RNA fragment. The signal from retroviral mRNA was also at least three fold more intense than that from the endogenous mRNA, indicating overexpression from the retroviral promoter. Phenotypically, the DT40 cells expressing the deletion Nr13 mutant have a normal growth curve under standard growth conditions in 10% BCS and 2% CS. However, when treated with 1% CS media, the amino-terminal deletion mutant-expressing cells were more susceptible than wild-type DT40 cells to serum withdrawal-induced cell death (Fig. 6c). Thus, the BH4 domain of Nr13 is essential for the inhibition of apoptosis under these conditions, and loss of the BH4 domain appeared to convert Nr13 into a death agonist.

**Nr13 protects bursal stem cells from programmed elimination**

The in vivo function of Nr13 was investigated using a bursal transplant model established in this laboratory. Donor bursal cells from unchallenged embryonic bursa can reconstitute recipient birds treated with cyclophosphamide, which ablates the lymphoid population of bursal follicles. Nr13 was introduced into donor cells by ex vivo cocultivation with LNRSN vectors for expression in reconstituted bursal follicles and their progeny. Bursal transplantation was performed as previously described (Neiman et al. 1985; Thompson et al. 1987) except that day 15 syngenic embryonic bursal cells were used as donors and cyclophosphamide-treated day 18 embryos were used as recipients. The transplanted birds were sacrificed at 4 weeks after hatching. Reconstituted bursal follicles appeared normal on routine histological examination. Western blot analysis detected Nr13 expression in reconstituted follicles compared with normal 4-week bursa, which lacks detectable Nr13 (Fig. 7a). The primary transplanted bursal cells 4 weeks after hatching were used as donors for a secondary transplantation experiment into cyclophosphamide-ablated recipient birds. Histological examination of the secondary transplant bursas revealed that 10–55% of the follicles were reconstituted and appeared normal (Fig. 7b, middle, and c). Bursas from either cyclophosphamide-ablated controls or five control transplants using donor cells from 4-week normal bursa resulted in empty follicles (Fig. 7b, left, and c), consistent with previous reports demonstrating absence of stem cells in bursa after hatching (Toivanen and Toivanen 1973; Pink et al. 1985). Western blot analysis confirmed that Nr13 was detected in secondary reconstituted follicles transduced with LNRSN (Fig. 7a). The reconstitution of follicles and detection of Nr13 in secondary transplant assays suggested that Nr13 induced persistence of stem cells. We concluded that Nr13 overexpression in bursal stem cells maintained the viability of the transplantable stem cells after hatching. To confirm this observation we repeated this transplantation experiment using the LNRCG vector [see Fig. 5a] to in-
of the cell death pathway. Nevertheless, as suggested by responses to phorbol ester and v-rel, Nr13 may play a role in the essential survival machinery in many types of bursal cells. Moreover, Nr13 may be central and essential to survival in the stem cell compartment, and perhaps in tumor cells derived from bursal stem cells. DT40 is derived from a bursal lymphoma induced by myc oncoprotein overexpression in bursal stem cells (Baba et al. 1985). We did obtain evidence that survival of these cells, at least in culture, was markedly influenced by Nr13, being enhanced by overexpression and diminished by a BH4 deletion mutation of Nr13.

Nr13 and Bax

Bax is a death agonist thought to function in part by interacting with and preventing Bcl2 or its homologs from binding with the CED4 homolog, Apaf1 (Oltvai et al. 1993; Sedlak et al. 1995). This interaction allows Apaf1 to activate a caspase cascade and induce cell death. Bax is also thought to trigger apoptosis by its pore forming activity (Schlesinger et al. 1997), which is also blocked by Bcl2. We used dispersion as a model to induce bursal cell death, and found that levels of Bax increase (and Nr13:Bax ratio decreases) with dispersion-induced cell death. However, Nr13 does not by itself appear to protect normal bursal cells from dispersion-induced apoptosis, although Nr13 interacts with Bax in DT40 cells based on coimmunoprecipitation. We have not obtained direct experimental evidence that Nr13 is able to attenuate the death effects of Bax, and we have not determined whether Bax has any more direct killing mechanism in bursa independent of Bcl2 family members. Currently we are characterizing the chicken bax gene to address these issues.

PMA induction of Nr13

Inhibition of bursal apoptosis by phorbol esters has been documented (Asakawa et al. 1993; Compton and Waldrip 1998). Phorbol esters activate the protein kinase C (PKC) family, which currently has at least 12 member isoenzymes. The classic PKC-α, PKC-βI, PKC-βII, and PKC-γ isoforms are activated by phorbol esters and are calcium dependent. The novel PKC-δ, PKC-ε, PKCη, and PKC-θ isoforms are calcium independent but activated by phorbol esters. All these isoforms have been linked to apoptosis in different cell lines, but results are conflicting (Deacon et al. 1997). In some systems, PMA treatment induces apoptosis, but in other systems such as the bursa, PMA inhibits apoptosis. We demonstrated by Northern blot analysis that PMA induced Nr13 at the transcriptional level. This induction could contribute to the mechanisms by which PMA acts to block cell death. However, simple overexpression of Nr13 does not by itself block dispersion-induced bursal cell death, indicating that induction of Nr13 is not sufficient to fully explain this effect of PMA.
Inhibiting bursal apoptosis by v-rel or other members of the NF-κB family

v-rel is one of the members of the NF-κB complex and is able to transform avian B cells [Neiman et al. 1991; Gilmore et al. 1996]. v-rel contains multiple internal mutations and a 118 amino-acid carboxy-terminal deletion compared to c-rel [Sarkar and Gilmore 1993]. This group of transcription activators plays an important role in signal transduction and proliferation, and also can either block or enhance apoptosis. Although multiple targets of the NF-κB factors have been identified [Gilmore et al. 1996], none of them are clearly linked to apoptosis except a recently identified chicken inhibitor-of-apoptosis (IAP) gene [You et al. 1997]. We observed here that Nr13 is induced by v-rel, and that both Nr13 gene transcription and v-rel activation of Nr13 expression is most efficient near physiological temperatures for chicken. Sequence analysis of the Nr13 promoter suggests a possible NF-κB binding site [G. Gillet, unpubl.] which could explain why Nr13 is induced by v-rel. These results suggest that v-rel and other factors could inhibit apoptosis through activation of Nr13. However, because expression of Nr13 does not, by itself, block apoptosis induced by dispersion in bursa, Nr13 is not the only determinant of v-rel-mediated inhibition of apoptosis. For example, the previously mentioned chicken IAP is induced by v-rel and may by itself, or in combination with Nr13, explain the effect of v-rel on bursal cell apoptosis.

The Nr13 BH4 domain

Although the functions of the conserved BH1, BH2, and BH3 domains are important to homodimer or heterodimer formation, the BH4 domain, which binds at least Raf1 [Wang et al. 1996a], CED4, calcineurin [Shibasaka et al. 1997], and BAG1 [Wang et al. 1996b], may function as a regulatory domain. Deletion of the BH4 domain of Bcl2 and the apparent BH4 helix of Nr13 both result in loss of the death-protecting function. It was reported that caspase cleaves at an aspartic acid residue immediately carboxy-terminal to the BH4 domain and converts Bcl2 into a death agonist [Cheng et al. 1997]. Because Bcl2 is considered to function upstream of the caspases, this cleavage could serve as a feedback amplification signal for the death cascade. A comparison of the sequence of Nr13 does not reveal an analogous aspartic acid for caspase cleavage. Moreover, Bcl2 and Bcl-XL both have a long flexible loop structure between the BH4 and BH3 domains [Muchmore et al. 1996], which also appears to be much shorter in Nr13. A third difference between Nr13 and Bcl2 is the presence of a carboxy-terminal tyrosine residue in Bcl2, which is important in regulation of G0-G1 cell cycle progression [Huang et al. 1997]. The BH4 domain of Nr13 does not have a similar tyrosine residue. Currently we are investigating whether Nr13 can affect cell cycle progression. These structural dissimilarities suggest that Nr13 may have regulatory interactions different from those of Bcl2 and Bcl-XL. Nr13 might interact with different proteins through the BH4 domain, and thereby play unique roles that are different from those of other Bcl2 family members.

Other implications of the bursal transplantation experiments

Overexpression of Nr13 in the chicken bursa did not lead to the development of neoplastic lesions, but did lead to the persistence of transplantable bursal stem cells. The morphology of the primary and secondary reconstituted follicles were normal in appearance. Nr13 overexpression in the bursal follicles might be similar to events in the development of human low-grade follicular lymphomas, which overexpress Bcl2 through a t[14,18] translocation [Bakhshi et al. 1985]. We have not completed long-term observation of transplanted birds. It will be interesting to learn whether the Nr13 overexpression in the bursa will eventually lead to the development of low-grade lymphomas. Finally, this technique for induced persistence of transplantable stem cells may also provide a useful tool for further investigation and characterization of these stem cells.

Materials and methods

Cloning of chicken Mcl1 and A1

Previously unidentified chicken homologs of A1 and Mcl1 were found by random single pass sequencing of an oligo (dT)-primed cDNA library prepared from activated ConA-treated splenic lymphocytes. The complete sequence of these clones was obtained by primer walking. The Mcl1 clone was incomplete and contained sequence encoding the carboxy-terminal 209 amino acids. This region showed 60% sequence identity with human Mcl1 [Kozopas et al. 1993], including the highly conserved Bcl2 homology (BH) domains. The A1 cDNA contained the entire coding region and encodes a protein of 174 amino acids that is nearly identical in size [175 amino acids] to human A1 [Karsan et al. 1996]. The predicted amino acid sequence shows 46% overall sequence similarity to the human A1 cDNA with two highly conserved BH domains. The GenBank accession numbers for chicken Mcl1 and A1 are AS120210 and AS120211, respectively.

Cell culture

Previously described retroviral packaging lines PE501 and PG13 were grown in DME supplemented with 10% fetal bovine serum [Miller and Rosman 1989]. DT40 bursal lymphoma-derived cells have been described [Baba et al. 1985]. DT40 cells were grown in DME supplemented with 10% BCS, 10% tryptose phosphate [DIFCO] and 2% chicken serum [Sigma]. For low serum condition experiments, DT40 cells were grown without BCS in DME supplemented with 10% tryptose phosphate and 1% CS.

Bursal transplant experiments

The technique of primary and secondary bursal transplantation has been described [Neiman et al. 1985; Thompson et al. 1987]. Briefly, fertilized eggs from inbred SC strain White Leghorn chicken were obtained from Hylane Farms, Dallas Center, IA. Cyclophosphamide, 1.25 mg/day, was injected into chorioallanto-
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toxic veins on days 15, 16, and 17 to ablate the bursa. Donor bursal cells were obtained from normal 15-day-old embryos or transplanted birds 3–4 weeks after hatching. Donor cells were resuspended in Hams media and dispersed by passing through sterile mesh. The dispersed bursal cells were cocultivated with retroviral vector-producing packaging cells overnight before injected into cyclophosphamide-ablated recipient birds. For secondary transplants, the recipient birds were newly hatched chicks injected with 3 mg of cyclophosphamide for 3 days to ablate the bursal lymphocytes and the donor cells were bursal lymphocytes from 4-week-old primary transplant birds. Transplanted birds were sacrificed at 4 weeks for histological and biochemical analysis. Frozen sections of the bursa were used for GFP detection.

Construction of retroviral vectors and packaging cell lines

The plasmid pBS–NR with Nr13 cDNA (Gillett et al. 1995) was digested with EcoRI and the 1.6-kb Nr13 fragment was inserted into the poly cloning site of pLXSN (Miller and Rosman 1989) or pLXCG retroviral vectors to construct pLNRSN and pLNRCG. pLXCG was derived from pLNCG (Rasko 1997) by replacing the BamHI–XbaI (SV40 promoter–neomycin resistance gene) fragment of LXSN with an analogous [CMV promoter-GFP] fragment from pLNCG. pMH105 plasmid containing v-rel (White and Gilmore 1993) was generously provided by Dr. Thomas Gilmore (Boston University). v-rel was excised with XbaI from pMH105 and inserted into an HpaI site of pLXSN vector by blunt end cloning. To generate amino-terminal helix deletion of Nr13, pBS–NR was digested with SmaI and NarI, and religated with linker oligonucleotides to create a 60-nucleotide deletion. Junctional sequence was confirmed by automated sequence analysis. All retroviral constructs were transfected into PE501 cells with linker oligonucleotides to create a 60-nucleotide deletion. To generate amino-terminal helix deletion of pMH105 and inserted into an HpaI site of pLXSN vector by blunt end cloning. To generate amino-terminal helix deletion of Nr13, pBS–NR was digested with SmaI and NarI, and religated with linker oligonucleotides to create a 60-nucleotide deletion. Junctional sequence was confirmed by automated sequence analysis.

Immunoprecipitation and immunoblotting analysis

DT40 cells were harvested and washed with cold phosphate buffered saline twice and lysed with buffer containing 50 mM tris (pH 7.5), 100 mM NaCl, 5 mM EDTA, 1% Triton X-100, 1% NP40, 1 mM PMSF protease inhibitor. The lysate was sonicated and centrifuged for 15 min. The supernatant was mixed with preimmune serum for 30 min. Immunoglobulin was removed by mixing with Pansorbin for 30 min. Pansorbin was removed by centrifugation, and the supernatant was incubated with Nr13 antiserum at 1:50 dilution for 4 hr. Pansorbin was added for another 45 min. The immune complex was precipitated and washed three times with RIPA buffer. SDS loading buffer was added to the washed Pansorbin and boiled 5 min to solubilize the immune complex before SDS-PAGE analysis. After the electrophoresis, the protein was transferred to a nitrocellulose membrane for standard Western blot analysis using a commercially available anti-Bax antiserum at concentration of 1:1000 (Santa Cruz Biotechnology, CA).

Northern blot analysis and RNAse protection assay

RNAs from DT40 cells or bursal tissues were isolated with RNAzol reagent (Tel-Test, Inc., Friendswood, TX). Samples were analyzed on 1% agarose gel, and transferred to Hybond membranes. Probes were labeled with 32P by random priming to a specific activity ~2 × 106 cpm/µg. The RNAse protection probe was cloned in antisense orientation into a Bluescript-KS plasmid (Stratagene, La Jolla, CA). RNA was transcribed in vitro by T7 RNA polymerase using [α-32P]UTP as label. RNA samples were hybridized with labeled probe overnight, followed by 10 mg/mL RNase digestion for 45 min. The protected fragments were analyzed by 6% urea–polyacrylamide gel and autoradiography.

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