Ikaros and RAG-2-Mediated Antisense Transcription Are Responsible for Lymphocyte-Specific Inactivation of NWC Promoter

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Abstract

Recombination activating gene-2 (RAG-2) and NWC are strongly evolutionarily conserved overlapping genes which are convergently transcribed. In non-lymphoid cells the NWC promoter is active whereas in lymphocytes it is inactive due to the DNA methylation. Analysing the mechanism responsible for lymphocyte-specific methylation and inactivation of NWC promoter we found that Ikaros, a lymphocyte-specific transcription factor, acts as a repressor of NWC promoter - thus identifying a new Ikaros target - but is insufficient for inducing its methylation which depends on the antisense transcription driven by RAG-2 promoter. Possible implications of these observations for understanding evolutionary mechanisms leading to lymphocyte specific expression of RAG genes are discussed.

Introduction

NWC (“Nad Wyraz Ciekawy”, which translates from Polish to “extremely interesting”) is the third evolutionarily conserved gene within the recombination-activating genes RAG-1 and RAG-2 locus [1] encoding a protein complex indispensable for the recombination of immunoglobulin and T-cell receptor minigenes [2–5]. The popular hypothesis on the origin of RAGs proposes that the transposon containing one of or both RAG genes infected a germ cell of an ancestor of jawed vertebrates (or deuterostomes), ultimately allowing for the development of lymphocytes [6,7]. The first exon of NWC gene and its promoter are located in the intron preceding the coding exon RAG-2 gene and these two genes are convergently transcribed [1] (Fig. 1A). The promoter of NWC gene is active in non-lymphoid cells and exhibits bidirectional activity, which can drive the transcription of both NWC and RAG-2 transcripts in some non-lymphoid cells [8]. Based on this observation, we have recently proposed that the bidirectional activity of NWC promoter could facilitate the integration and survival of RAG transposon in the ancestral genome [8]. Previously, we suggested that NWC transcription may negatively control RAG-1 and RAG-2 promoter activities in non-lymphoid cells owing to transcriptional interference caused by NWC transcription proceeding through RAG-2 promoter and RAG-1/ RAG-2 cis-regulatory elements localized upstream RAG-2 gene [9]. This hypothesis has not been verified so far, since due to the remaining activity of a secondary promoter [10], we have been unable to abrogate completely the transcription of NWC in mice, in which primary NWC promoter was deleted.

The primary NWC promoter is associated with a CpG island which is unmethylated in non-lymphoid cells and becomes methylated in immature T- and B-lymphocytes, which coincides with the promoter’s inactivation [11]. In lymphocytes the function of NWC promoter is taken over by RAG-1 promoter, which results in the expression of RAG-1/NWC hybrid transcripts [1]. The methylation of NWC promoter is not accompanied by other changes in chromatin organization, i.e. changes in posttranslational modifications of histone H3 [11] that are commonly associated with transitions between transcription permissive and repressive chromatin configuration and usually precede DNA methylation. Blocking DNA methylation with 5-azacytidine partially restores the activity of NWC promoter in lymphocytes [11], proving the primary role of DNA methylation in controlling its activity. The activation of NWC promoter is mediated by ZFP-143 transcription factor which binds to its two conserved elements, also possessing consensus binding sites for Ikaros transcription factor [8].

Ikaros is an essential transcription factor required for lymphocyte development. It is expressed in lymphoid cells, haematopoietic stem cells and some myeloid cells. Ikaros deficiency impairs the development of lymphoid and myeloid cell lineages [12]. Ikaros can be involved both in gene activation and repression and its activity occurs at different levels: by direct competition with the activator proteins for common binding sites at the target promoter.
by restructuring chromatin through targeting different types of chromatin remodelling factors [14–16] such as SWI/SNF (activator) or NuRD deacetylase (repressor) as well as by bridging the target genes destined for inactivation with centromeric foci, thereby facilitating their assembly into pericentromeric heterochromatin [17]. Ikaros target genes include RAG-1 and RAG-2 genes, which are tightly controlled throughout lymphocyte development. High and coordinated expression of RAG genes is regulated by the activity of several cis-elements localized mainly upstream RAG-2 gene [18–21]. Investigating the role of Ikaros in regulating V(D)J recombination in B-cell lineage Reynaud and colleagues [22] showed that Ikaros binds directly to regulatory elements of RAG locus in pro-B cells, namely with the Ep, D3, E- rag enhancers and RAG-1 promoter but not to RAG-2 promoter. These authors also compared the histone modification status of regulatory elements in RAG2+/− vs Ikfz−/− pro-B cells and concluded that activation of RAG transcription by Ikaros is accompanied by histone-H3 acetylation. Here we demonstrate that NWC represents a new target of Ikaros activity within RAG locus. We show that binding of Ikaros to NWC promoter downregulates NWC expression, but is unable to cause promoter methylation which is established by antisense transcription driven by the activity of RAG-2 promoter. We discuss how these two mechanisms: binding of Ikaros and antisense transcription may act in concert to inactivate NWC promoter.

Results

Ikaros binds to NWC promoter

We have recently shown that the promoter of NWC gene is activated by ZFP-143 transcription factor, which binds two inverted evolutionarily conserved sites of the promoter [8] spanning −10/−37 and −74/−100 nucleotides relative to transcriptional start site. We noticed that these regions also contain consensus binding sites for Ikaros transcription factor (TGGGAA) [12], which overlap with the ZFP-143 binding

Figure 1. Ikaros-binding properties of NWC promoter. (A) Structure of RAG/NWC locus and NWC gene promoter. Open gray and black boxes represent NWC, RAG-2 and RAG-1 exons, respectively. Horizontal arrows indicate the directions of the transcription and the numbers indicate the position of the sequences relative to NWC transcriptional start site. Red line indicates the localization of the EMSA probe (−119/+12) lying within the NWC promoter (−119/+125). Aligned sequences of murine (Mus musculus) and human (Homo sapiens) NWC promoters are shown. The sequence logos represent ZFP-143 binding sites and putative Ikaros binding sites. Vertical arrows indicate the location and nature of the mutations introduced in probes and reporter constructs used throughout this study. (B) EMSA experiment showing Ikaros binding to NWC promoter: first lane from the left – free probe, second – probe and Ikaros, next – probe, Ikaros and increasing molar excess (10, 50, 100x) of unlabelled specific (lanes 3–5) or non-specific competitors (Oct-2, for sequence see Materials and Methods) (lanes 6–9). (C) EMSA experiment showing Ikaros binding to non-mutated (NM), single mutated (mutA, mutT) or double mutated (mutAT) NWC promoter. Probes were tested in the absence (−) and presence (+) of Ikaros. (D) Competition of ZFP-143 and Ikaros for NWC promoter binding. Constant amount (150 ng) of ZFP-143 protein and increasing amount of Ikaros protein (Ikaros/ZFP-143 molar ratio: 0, 0.3, 0.5, 1, 1.5, 2, 2.5, 3, lanes 2–8) were used to bind to the probe corresponding to non-mutated NWC promoter.

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sequences (Fig. 1A). This observation raised the possibility that Ikaros could inactivate NWC promoter by competing with ZFP-143. As a first step towards verifying this possibility, we checked whether Ikaros is able to bind to the promoter using electromobility shift assay (EMSA). The recombinant HisTag-Ikaros protein yielded two complexes with the probe corresponding to −119/+12 portion of NWC promoter (Fig. 1B). The slow-mobility complex was disrupted when the probe, mutated at one of the two binding sites (TGAGAA) were used, while mutation of both binding sites resulted in complete disappearance of slow- and fast-migrating complexes (Fig. 1C). These results suggested that Ikaros is able to bind simultaneously and independently to both predicted sites in the promoter, which indicates that ZFP-143 and Ikaros share the same binding sites containing four strict consensus nucleotides (TCCG) indispensable for binding both proteins. In order to confirm this conclusion we compared their binding to the promoter in an EMSA competition experiment. Addition of increasing concentrations of HisTag–Ikaros to binding reactions containing the NWC promoter probe and a constant amount of ZFP-143 led to a gradual reduction in the abundance of the ZFP-143-containing complex. Addition of Ikaros in a ratio ranging from 0.3:1 to 2:1 produced an intermediate migrating complex containing presumably one molecule of Ikaros and ZFP-143, whereas addition of Ikaros in a ratio of 2:5:1 resulted in the disappearance of fast (ZFP-143) and intermediate (ZFP-143/Ikaros) complexes, producing ones containing only Ikaros (Fig 1D).

Ikaros downregulates the expression of NWC

In order to determine whether Ikaros influences the expression of NWC, HEK293T cells were transfected with pLVX Ikaros-IRES-GFP expression vector. Transfected cells were sorted based on the high GFP expression and assayed for the expression of NWC in Real-Time RT-PCR assay. As shown in Fig. 2A, overexpression of Ikaros led to a significant (~7 times) downregulation of NWC expression. In order to find out if the downregulation is due to a direct interaction of Ikaros with NWC promoter we tested the effect of Ikaros overexpression in cells co-transfected with reporter vectors containing NWC promoter constructs. Figure 2B shows that the overexpression of Ikaros resulted in a significant reduction of the promoter activity of constructs containing an NWC promoter fragment (−119/+125 relative to the NWC transcription start site) but had no effect on the control SV40 promoter. In order to confirm the specificity of this effect we used NWC promoter fragments containing point mutations affecting the consensus binding sequences for both Ikaros and ZFP-143 (mAT) or for Ikaros alone (mGC). The influence of introduced mutations on the ability of Ikaros and ZFP-143 to bind the constructs was verified by EMSA (Fig. 2C), confirming Ikaros-specific nature of mGC mutation. As expected, promoter containing mutations in binding sequences for both ZFP-143 and Ikaros showed significantly reduced activity when compared to a non-mutated promoter and the effect was not further enhanced with the overexpression of Ikaros. The promoter fragment containing the mutations in Ikaros binding sequences had similar activity as the non-mutated promoter, but, in contrast to the latter, its activity was not reduced after the overexpression of Ikaros (Fig. 2B). In order to determine whether the reduction in NWC promoter activity caused by Ikaros overexpression is accompanied by the methylation of NWC promoter we performed bisulfite sequencing of NWC promoter using HEK293T cells transfected with pLVX Ikaros-IRES-GFP vector. Ikaros overexpression did not result in any changes in the methylation status of NWC promoter as compared to non-transfected cells: NWC promoter was unmethylated in both cell types (0%–1%) (not shown). Lymphoid cell line (Jurkat) was used as a control in this experiment and was shown to have completely methylated NWC promoter. Altogether these results indicate that Ikaros is able to bind to NWC promoter and reduce its activity owing to the competition with ZFP-143 transcriptional activator for common binding sites but alone is not sufficient for the methylation of NWC promoter.

RAG-2 antisense transcription is responsible for NWC promoter methylation

Searching for a mechanism that could be responsible for lymphoid-specific methylation of NWC promoter we focused our attention on RAG-2 transcription, which, because of a convergent transcription of these two genes, represents an antisense transcription in relation to NWC. In order to test if RAG-2 transcription influences the methylation status of NWC promoter we have generated a transgenic mouse strain using BAC-based transgene containing complete murine RAG/NWC locus modified to express GFP under the control of RAG-2 promoter [18]. We have modified this transgene by inserting YFP gene in frame with NWC gene and a transcriptional termination cassette consisting of two SV40-polyA sequences linked with twelve loxP operators [23] immediately downstream of RAG-2 first exon (BAC-RG/Germ). A transgene without the termination cassette was used to generate the control mouse strain (BAC-RG/NY) (Fig 3A). The offspring of the transgenic founder animals was analyzed by flow cytometry. As expected, in the BAC-RG/NY control mouse, the highest level of the expression of RAG-2/GFP was detected in developing thymocytes and B lymphocytes while the highest expression of NWC/YFP was detected in testis (Fig 3B) consistent with our previous findings [1,10] that these cells express the highest level of NWC transcript. Figure 3B also shows that the BAC-RG/Germ/NY mice showed a strong reduction of expression of RAG-2/GFP providing evidence that the transcriptional termination cassette was functional. Since the cassette was also found to be bidirectional, it strongly reduced the expression of NWC-YFP reporter (Fig 3B) and made it impossible to monitor the potential effect of RAG-2 transcription termination on NWC promoter activity by flow cytometry. Therefore, we analyzed the methylation level of NWC promoter by bisulfite sequencing. Using transgene-specific primers (i.e. one of the primers in each pair in the nested PCR was complementary to GFP) we were able to distinguish transgenic from endogenous loci and thus determine the influence of the termination of RAG-2 transcription on NWC promoter methylation. As shown in Figure 3C, the termination of RAG-2 transcription significantly reduced the methylation level of NWC promoter as compared to the control mice, indicating that the methylation is due to a cis-mechanism. In double positive (CD4+/8+) thymocytes, single positive thymocytes, bone marrow preBII small cells and splenic B cells the methylation level was reduced from nearly 100% to ~40%. Importantly, in non-lymphoid tissues (liver, brain and testis) of both mouse strains, the methylation level of NWC promoter was similarly low (12%~17%), indicating that the differences in promoter methylation observed in lymphocytes were not due to the position effect of the integrated transgene.

Discussion

In the present study we have shown that two mechanisms contribute to the lymphocyte specific inactivation of NWC promoter: Ikaros induced repression and DNA methylation, gained through cis-antisense transcription driven by RAG-2
transcription. We have shown that Ikaros can bind to NWC promoter and outcompete its activator, ZFP-143 transcription factor, from the common binding sites. Within RAG/NWC locus Ikaros has two opposing roles in transcription regulation. It activates RAG transcription by binding to cis-regulatory elements and to RAG-1 promoter [22] and downregulates the expression of NWC gene.

Cis-antisense transcripts spanning the CpG islands are known to have a causative role in establishing the DNA methylation but the exact mechanism of their action is not clear. One possibility is that the observed cis effects are due to the ability of the antisense RNA to co-transcriptionally interact with the target DNA to form different kinds of RNA/DNA hybrids which may attract DNA methyltransferases guiding them to the target sequences [24]. Another possibility emerging from the analysis of the mechanism silencing KIR3DL1 promoter is that antisense transcription driven by a proximal promoter results in the formation of dsRNA which is processed into 28 base PIWI-like RNA [25]. The transcriptional shutdown of a single gene in a cluster of other closely related KIR genes would suggest that the antisense RNA is processed rapidly and mediates silencing without diffusion or transport away from the promoter. The chromatin modification accompanying the promoter methylation does not spread across the entire locus thus allowing for unaffected expression of other KIR genes [25]. Similar mechanism could be responsible for NWC promoter silencing as DNA methylation at the NWC promoter changes the chromatin structure only locally and does not affect the transcriptional activity of RAG genes [1,11]. Although because of the bidirectional activity of the transcriptional termination cassette we were not able to directly quantify the effect of RAG-2

Figure 2. Ikaros downregulates the activity of NWC promoter. (A) Real-Time RT-PCR analysis of endogenous NWC expression in non-lymphoid cells (HEK293T) transfected with Ikaros-expressing construct (pLVX-Ikaros-IRES-ZsGreen) or with empty vector (pLVX-IRES-ZsGreen) (100%). Transfection efficiency varied and approximated 80%. The MFI after sorting was ~1300. Expression values were normalized to GAPDH. The results shown are the means of three experiments with error bars representing ± 1 SD. The asterisk indicates a significant difference (p < 0.05) between both sample groups. (B) Activity of NWC promoter fragments containing point mutations in the Ikaros/ZFP-143 binding sites tested by luciferase assay in HEK293T cells ectopically expressing Ikaros gene (white bars) or transfected with empty vector (grey bars). pNWC-NM: non-mutated binding sites, pNWC-mGC: mutation specifically affecting the Ikaros binding site (see C), pNWC-mAT: mutation affecting both Ikaros and ZFP-143 binding sites. The relative promoter activities were normalized to the activity of promoter-less vector (pGL3-Basic). Vector containing SV40 promoter was used as a control. The results shown are the means of three experiments with error bars representing ± 1 SD. The asterisk indicates a significant difference (p < 0.05) between the activity of the given promoter in the presence or absence of Ikaros expression. (C) EMSA experiments verifying the Ikaros-specific nature of mutations (mGC) introduced to NWC promoter constructs. Non-mutated probes (NM) and probes with mutations affecting both Ikaros and ZFP-143 binding (mAT) were used as controls to monitor the efficiency of Ikaros and ZFP-143 binding.

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transcription termination on the activity of NWC promoter, we expect, based on previous experiments with blocking DNA methylation by 5-azacitidine, that the promoter activity is at least partially restored [11]. The incomplete re-expression of NWC in 5-azacitidine-treated lymphoid cells can be explained by the repression of NWC promoter by Ikaros and/or transcriptional interference caused by collision RAG-2 and NWC transcription processes.

Taking into consideration the widely accepted view that RAG genes “infected” the ancestral genome in the form of a transposon[5], the structure of NWC and modes of its regulation described in the present paper, we would like to add new elements to our recent proposal [8]. We propose that NWC locus was the original site of integration of RAG transponson rather than being part of it, which is suggested by the conserved structure of RAG/NWC locus (Fig. S1), multi-exon structure of NWC and by the fact that its homologues can be found in various invertebrate species (Fig. S2). Although the conserved NWC protein domains do not share homology to any known protein and the function of the protein is still obscure, the presence of NWC gene throughout animal kingdom and its mode of regulation suggest that it is a housekeeping gene. We earlier proposed that constitutive expression of NWC as well as the bidirectional activity of its promoter, a feature which characterizes many promoters controlling transposon-derived genes, could initially facilitate the integration and survival of RAG transposon, while cis-regulatory elements controlling lymphocyte specific expression of RAG genes were acquired later in evolution. The localization of the RAG-2 promoter at the 5′ end of the first exon of NWC resulted in convergent and overlapping transcription of NWC and RAG-2 genes, which, if occurred simultaneously, could result in down-regulation of both genes owing to transcriptional interference. Such mechanism could inhibit RAG expression in lymphocytes below the level required for efficient V(D)J recombination. We think that the expression of Ikaros protein at the initial stage of lymphocyte development in hematopoietic stem cells and its binding to NWC promoter causes the displacement of the activator protein ZFP-143 and shifts the equilibrium of opposing transcriptional processes in favor of RAG-2 transcription, which is additionally activated by Ikaros binding to RAG regulatory elements. RAG-2 transcription proceeding through NWC promoter causes its methylation and inactivation without engagement of other factors influencing chromatin accessibility thus preventing the spreading of the changes across the entire locus and enabling undisturbed RAG expression.

Figure 3. RAG-2 transcription is responsible for NWC promoter methylation. (A) The RAG/NWC locus of BAC-RG/NY and BAC-RGterm/NY transgenic mice. Relative positions of the exons encoding RAG-1 (black boxes), RAG-2 (open boxes), and NWC (gray boxes) are shown. Horizontal arrows indicate transcription start sites and orientations. Modifications of the locus are presented as green, yellow and red boxes representing GFP, YFP and the transcriptional termination cassette, respectively. (B) Cytofluorimetric evaluation of RAG-2/GFP protein expression in double positive thymocytes (upper histogram) and bone marrow B lymphocytes (middle histogram) and NWC-YFP in testis (lower histogram) of BAC-RGterm/NY (bold line), BAC-RG/NY (normal line) and non-transgenic control (dashed line). (C) Methylation status of transgenic NWC promoter of control (grey bars) and RAG-2 terminator-containing (white bars) loci.

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The evolutionarily conserved association of NWC with RAG-1 and RAG-2 genes raises additional questions concerning the nature of primordial RAG transposon. As a result of the discovery of Transib transposons closely resembling the core fragment of RAG-1 gene, a modified version of the hypothesis has been put forward, suggesting that only RAG-1 gene was a part of a mobile element which integrated within RAG-2 locus [26]. The discussion on the origin of RAG-1/RAG-2 genes has ignored, however, the existence of NWC gene, which most probably was the host gene for RAG transposon integration. Given that invertebrate species NWC gene is not associated with any gene resembling RAG-2 and the bidirectional activity of NWC promoter is typical for many host genes controlling transposon-related genes, it is reasonable to assume that the genetic element which integrated within NWC locus contained both RAG-1 and RAG-2 genes. Interestingly, in Strongylocentrotus purpuratus, the only organism outside jawed vertebrates phylum in which both RAG-1 and RAG-2 genes were found [27], the RAG and NWC loci are separate. Since it is highly unlikely that RAG-1-containing element integrated twice with RAG-2 locus this again argues for the “RAG-1 and RAG-2” hypothesis and suggests that the transposon independently infected the ancestors of jawed vertebrates and echinoderms. However, this point of view needs to be verified by extensive analysis of NWC transcriptional regulation both in vertebrates and invertebrates.

**Materials and Methods**

**Expression and purification of recombinant proteins**

PCR-amplified Ikaros cDNA was cloned into Neo/Xhol sites of the pET32a vector to obtain expression vector encoding N-terminal His-tagged recombinant Ikaros protein (pET32a-Ik). In order to express and purify the protein, overnight culture of E. coli BL-21 cells transformed with pET32a-Ik was diluted 500 times, cultured at 37°C until OD600 reached 0.5 and then incubated for the next 2 hours at 4°C. Then the cells were induced with IPTG (0.25 mM) and cultured overnight at room temperature in the presence of ZnCl₂ (100 μM). The bacterial pellet was resuspended in buffer A, sonicated and applied on HisPur Ni-NTA resin equilibrated with buffer A. Binding the recombinant protein was performed for 1 hour at 4°C and then the column was washed several times with buffer B and eluted with buffer C. After elution, the recombinant protein was dialyzed against buffer D.

Buffer A: 10 mM imidazole, 50 mM Tris-HCl, 300 mM NaCl, 10% glycerol, 0.05% Tween 20, pH 8.0. Buffer B: 20 mM imidazole, 50 mM Tris-HCl, 300 mM NaCl, 10% glycerol, 0.05% Tween 20, pH 8.0, Buffer C: 250 mM imidazole, 50 mM Tris-HCl, 300 mM NaCl, 10% glycerol, 0.05% Tween 20, pH 8.0, Buffer D: 20 mM HEPES pH 7.9, 0.2 mM EDTA, 20% glycerol, 100 mM KCl, 1 mM DTT. All buffers were supplemented with freshly added ZnCl₂ (20 mM), MgCl₂ (30 mM), CaCl₂ (50 mM) and PMSF (1 mM).

**ZFP-143 recombinant protein was obtained as previously described [8].**

**Electromobility shift assays**

The probe containing NWC promoter was obtained by amplifying DNA fragment encompassing −119/+12 nucleotides relative to NWC transcriptional start site using digoxigenin-labelled primer. PCR ligation method was used to obtain probes with mutations in ZFP-143 and/or Ikaros binding sites. The binding reaction was performed for 20 minutes on ice in 20 μl of reaction mix containing: binding buffer (20 mM HEPES pH 7.9, 0.2 mM EDTA, 20% glycerol, 100 mM KCl, 1 mM DTT, 10 μM ZnCl₂), 150 ng of purified recombinant protein, 1 μg poly(dI·dC) and 0.035 pmol of the probe. Where mentioned, 10, 50 or 100 μM molar excess of unlabelled specific (TCAGCTTTGGGAAATGTATCCCCTGCTA) or non-specific (Oct2) (GGCGTAAATCCTAAATTCACCGCC) competitor oligonucleotides was added. For ZFP-143/Ikaros competition, 150 ng ZFP-143 was used in the presence of increasing concentration of Ikaros protein (Ikaros/ZFP-143 molar ratio: 0.3, 0.5, 1, 1.5, 2, 2.5, 3, 4). The binding reaction was subjected to electrophoresis in 5% polyacrylamide at 4°C. The products were transferred on nylon membrane and detected with anti-digoxigenin antibodies according to manufacturer’s recommendations.

**Cell culture and sorting**

HEK-293T cells were cultured in DMEM medium (Sigma-Aldrich) supplemented with 10% of FBS (Invitrogen). Jurkat cells were cultured in RPMI medium supplemented with 10% FBS. For Ikaros expression, 0.5×10⁶ HEK-293T cells were plated on 10 cm dish and transfected with 2 μg of Ikaros expressing vector using MetafectenePro reagent (Biontex). The vector was constructed by cloning PCR-amplified Ikaros cDNA into pGEMT-Easy vector (Promega) and then subcloning the EcoRI fragment into pLVX-IRES-ZsGreen1 vector (Clontech). After 48 hours ZsGreen₁⁺⁺ cells were sorted using FACS-Aria instrument and used for downstream experiments (Real-Time RT-PCR and bisulfitesequencing).

**Real-Time RT-PCR**

RNA was isolated from cultured cell lines using TRIZol Reagent (Invitrogen) according to the manufacturer’s recommendations. Three micrograms of total RNA was digested with RNase-free DNase I (Thermo Scientific) and reversed transcribed with SuperScript III Reverse Transcriptase (Invitrogen) and random hexamer oligonucleotides at 50°C. Real-Time RT-PCR was performed on a DNA Engine Opticon 2 apparatus (Biorad) using Maxima SYBR Green qPCR Master Mix (Thermo Scientific). The thermal-cycling conditions comprised an initial denaturation step at 95°C for 10 min and 40 cycles of three-step PCR, including 15 s of denaturation at 95°C, 30 s of annealing at 55°C, and 30 s of elongation at 72°C. Expression values were normalized to HPRT. Standard curves were prepared for each primer pair by serial 5-fold dilutions of the template cDNA allowing determination of reactions efficiencies. One-way ANOVA followed by Tukey-Kramer post-hoc test was used for statistical analysis.

**Primers sequences:**

- **NWC:** GTCTGGCCCATATGTGCAGGATTG (forward)
- **NWC:** CTCTTCTACATCGTCCCAAATTC (reverse)
- **HPRT:** TGACCTTGATTTTATTTGCAACC (forward)
- **HPRT:** CGAGCAAGACGTTCAGTCCCT (reverse)

**Dual Luciferase Reporter assay**

pGL3-Basic-based reporter vectors containing NWC promoter were obtained as previously described [8]. PCR ligation method was used to obtain constructs with mutations in ZFP-143 and/or Ikaros binding sites. A total of 2×10⁵ HEK293T cells were transfected with 500 ng of firefly luciferase containing reporter plasmids and 50 ng of renilla luciferase containing plasmid (pRL-TK) using MetafectenePro reagent (Biontex) according to the manufacturer’s recommendations. The DLR assay was performed 24 h after the transfection using Dual-Luciferase Reporter Assay System reagents (Promega). In the experiments using Ikaros expression vector, the cells were transfected with 500 ng of Ikaros expression vector (pCDNA3-Ikaros, kind gift from S. Smale), 100 ng of firefly luciferase containing reporter plasmids and 50 ng of pRL-TK plasmid, and cultured for 48 hours. The cells were
lysed with 100 μl of passive lysis buffer and 15 μl of cell lysates were taken for each analysis. The data is presented as a ratio of firefly (FLU) to renilla (RLU) luciferase activity. One-way ANOVA followed by Tukey-Kramer post-hoc test was used for statistical analysis.

Bisulfite sequencing

Two micrograms of genomic DNA were treated with HCl (0.1 N) for 2 minutes at room temperature and denatured with NaOH (0.3 M) for 20 minutes at 37°C. The DNA was then treated with sodium bisulfite (35.5%, pH 5.0) in the presence of hydroquinone (0.5 mM) for 5 h at 55°C. The converted DNA was then bound and washed on the Genomic DNA Extraction column (Genoplast) and desulfonated by adding 0.15 NaOH in 90% EtOH on the column and incubating for 10 minutes at room temperature. DNA was washed and eluted from the column. Nested PCR amplification (2×30 cycles) was performed using primer pairs corresponding to the upper strand of the transgenic locus: outer 5′-TTTAAGGAGTGGAATGTGGTTAGTTA (forward) 5′-ACTCCACACAAACAAATCTATCTCC (reverse) and inner 5′-GGGATATGTTTTTTAGGATTTTTGGG (forward), 5′-AATCACCTATTTCAAAACTCAGAAA (reverse) or human locus: outer 5′-TCTCTCAATCTCTTACCTCCCAA (forward) 5′-GGTGCGTATGATGATTTTAGGGTTA (reverse) and inner 5′-TCCTAATCTCTTACCTCCAA-CACC (forward), 5′-GATTAGGATGGTTATTT (reverse) PCR product was directly cloned into pGEMT-Easy vector (Promega) and at least 20 individual clones were sequenced.

In order to test if PCR amplification of bisulfite-treated DNA did not produce artifacts by selective enrichment of unmethylated or methylated DNA fragments, cubic polynomial regression correction method was used [28]. A calibration experiment was performed based on amplified transgenic locus. A fully methylated PCR fragment was obtained using SssI methylase. Unmethylated or methylated DNA fragments were mixed to obtain DNA of known (0%, 25%, 50%, 75%, 100%) level of methylation and PCR fragment was obtained using SssI methylase. Unmethylated correction method was used [28]. A calibration experiment was performed exactly according to the published protocol [29]. Briefly, BAC-RG/NY transformed into E. coli SW105 strain was first modified by recombination by inserting the galK cassette flanked by 50 nt homology arms corresponding to the targeted region of RAG-2 gene and the cells were selected on minimal medium containing galactose as the sole source of carbon. Then the termination cassette flanked by the same homology arms was transformed to the cells and the cells were grown on medium containing glycerol and 2-deoxy-galactose (DOG) in order to select clones in which the termination cassette had replaced the galk cassette. Both constructs (BAC-RG/NY and BAC-RGterm/NY) were linearized with BstHII, purified by field inversion gel electrophoresis and used to generate transgenic C57BL/6 mice strains (Karolinska Center for Transgene Technologies, Karolinska Institutet, Stockholm). The transgenic founder mice were then bred in our animal facility and transgenic offspring was used for further experiments. Thymocytes (CD4+CD8+), splenic T-cells (CD4+CD8+), splenic B-cells (CD19+), bone marrow preB small cells (small CD25+CD19+) were isolated from 6-week-old mice by staining with anti CD4-PE/anti CD8-APC, CD19-PE, CD25-APC/CD19 PE antibodies (Becton Dickinson), respectively. The cells were sorted with FACS-Aria instrument. Non-lymphoid tissues (brain and liver) were used directly to isolate DNA and its subsequent methylation analysis.

Supporting Information

Figure S1 Structure of the RAG/NWC locus in vertebrate species. (TIF)

Figure S2 Multiple sequence alignment of vertebrate full-length NWC proteins and C-terminal portions of vertebrate and invertebrate NWC proteins. (TIF)

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Author Contributions

Conceived and designed the experiments: AL LB MK LS PK MC. Performed the experiments: AL LB MK LS SJ MC. Analyzed the data: AL LB MK LS SJ PK MC. Contributed to the writing of the manuscript: AL LB PK MC.
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