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# Basic helix-loop-helix proteins E2A and HEB induce immature T-cell receptor rearrangements in nonlymphoid cells

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T-cell receptor (TCR) gene rearrangements are mediated via V(D)J recombination, which is strictly regulated during lymphoid differentiation, most probably through the action of specific transcription factors. Investigated was whether cotransfection of RAG1 and RAG2 genes in combination with lymphoid transcription factors can induce TCR gene rearrangements in nonlymphoid human cells. Transfection experiments showed that basic helix-loop-helix transcription factors E2A and HEB induce rearrangements in the TCRD locus (Dô2-Dô3 and Vô2-Dô3) and TCRG locus (ψ Vγ7-Jγ2.3 and Vγ8- $J\gamma$ 2.3). Analysis of these rearrangements and their circular excision products re-

vealed some peculiar characteristics. The Vô2-Dô3 rearrangements were formed by direct coupling without intermediate  $D\delta 2$ gene segment usage, and most  $D\delta 2-D\delta 3$ recombinations occurred via direct coupling of the respective upstream and downstream recombination signal seguences (RSSs) with deletion of the  $D\delta 2$ and D<sub>03</sub> coding sequences. Subsequently, the E2A/HEB-induced TCR gene recombination patterns were compared with those in early thymocytes and acute lymphoblastic leukemias of T- and B-lineage origin, and it was found that the TCR rearrangements in the transfectants were early (immature) and not necessarily Tlineage specific. Apparently, some parts

of the *TCRD* (V $\delta$ 2-D $\delta$  region) and *TCRG* genes are accessible for recombination not only in T cells, but also in early B-cells and even in nonlymphoid cells if the appropriate transcription factors are present. The transfection system described here appeared to be useful for studying the accessibility of immuno-globulin and TCR genes for V(D)J recombination, but might also be applied to study the induction of RSS-mediated chromosome aberrations. (Blood. 2001;98: 2456-2465)

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#### Introduction

Antigen recognition by lymphocytes is dependent upon successful rearrangement of immunoglobulin and T-cell receptor (TCR) genes from variable (V), diversity (D), and joining (J) gene segments through the process of V(D)J recombination.<sup>1</sup> The rearrangement processes are mediated by the recombination-activating gene (RAG)-1 and RAG2 proteins, which specifically recognize the recombination signal sequences (RSSs) that flank the coding regions of the V, D, and J gene segments.<sup>2-4</sup> RSSs are consensus sequences consisting of a heptamer and nonamer separated by a 12-base pair (bp) or 23-bp spacer. Site-specific cleavage at the borders of RSSs and coding elements by the RAG proteins is followed by a process of rejoining of DNA ends in which the double-strand break-repair enzymes play a central role. Antigen receptor assembly is critically dependent upon expression of the lymphoid-specific RAG proteins; this is further illustrated by the fact that ectopic RAG expression results in site-specific recombination both in vitro and in vivo.2-4

The whole process of V(D)J recombination is ordered and tightly regulated during lymphoid differentiation.<sup>5</sup> The hierarchical order is apparent at different levels. First, cells committed to the B-cell lineage undergo immunoglobulin rearrangements, whereas TCR genes rearrange in T-cell precursors. Nevertheless, so-called cross-lineage rearrangements might occasionally occur in precursor B and T cells; this phenomenon is particularly evident from malignantly transformed lymphoid precursor cells, ie, B- and T-lineage acute lymphoblastic leukemias (ALLs).<sup>6,7</sup> Furthermore, immunoglobulin (Ig) heavy chain (*IGH*) rearrangements are known to precede Ig kappa (*IGK*) and Ig lambda light-chain recombination.<sup>8</sup> Similarly, data from human T-ALL and sorted human thymocyte subpopulations indicate that TCR $\delta$  (*TCRD*) recombination occurs prior to TCR $\gamma$  (*TCRG*) and TCR $\beta$  (*TCRB*) rearrangement, whereas the TCR $\alpha$  (*TCRA*) locus is rearranged at a late stage only.<sup>9-11</sup> The ordered process is also apparent from the finding that D<sub>H</sub>-J<sub>H</sub> or D $\beta$ -J $\beta$  rearrangements generally precede the coupling of their respective V gene segment usage, as is apparent from murine thymic ontogeny where V $\gamma$  and V $\delta$  gene rearrangements occur in waves with different V segments being used in fetal and adult thymocytes.<sup>14,15</sup>

It has been suggested that the tight and hierarchical regulation of the rearrangement processes can be explained by differential chromatin accessibility to the V(D)J recombinase,<sup>16</sup> which in turn is controlled by transcription factors binding to promoters and enhancers. E proteins are an important class of transcription factors in lymphoid differentiation. They consist of a helix-loop-helix (HLH) dimerization motif and a basic DNA binding domain that binds to conserved E-box motifs, as identified in immunoglobulin, *TCRB*, and CD4 enhancers. Members of the E-protein family

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include E2-2, HEB, and the E12 and E47 splice variant products of the E2A gene.<sup>17</sup> E12 and E47 are differentially expressed in a wide variety of tissues, but exist as homodimers only in B-lineage cells.18,19 In T cells, E-box binding complexes are heterodimers of E2A and HEB. E2A<sup>-/-</sup> mice have a block in B-cell differentiation prior to the start of the immunoglobulin recombination process.<sup>20,21</sup> Induction of D<sub>H</sub>-J<sub>H</sub> and IGK rearrangements upon ectopic E2A expression in a pre-T-cell line or even nonlymphoid human cells illustrates its involvement in regulation of V(D)J recombination.  $^{22,23}\ \text{E2A}^{-/-}$  mice also have a severe, though less complete, defect in T-cell differentiation at the double-negative stage. Interestingly, in E2A<sup>-/-</sup> mice, certain TCRγδ T-cell subsets are lacking, owing to an impaired ability to rearrange particular TCRG and *TCRD* gene segments.<sup>15</sup> Targeted disruption of the *HEB* gene gives rise to a partial block in early T-cell differentiation prior to the development of double-positive cells.24

It is now generally accepted that V(D)J recombination processes are also involved in the formation of particular chromosome aberrations in human leukemias.<sup>25-27</sup> This especially concerns aberrations in human T-ALL, in which *TCRB* or *TCRD* gene segments are translocated to oncogenes, resulting in activation of these oncogenes through TCR regulatory elements.<sup>28</sup> Although little is known about the exact molecular processes, it is tempting to speculate that transcription factor–induced accessibility of the involved loci is a critical step in the formation of these aberrations.

Here we show that 2 types of E proteins, E2A and HEB, have the ability to target the recombination machinery to TCR loci in nonlymphoid cells. Expression of E2A or HEB in the presence of RAG1/RAG2 appeared to induce immature types of *TCRD* rearrangements, several *TCRG* rearrangements, but no *TCRB* recombination.

#### Materials and methods

#### Cell culture

Nonlymphoid BOSC 23 cells were cultured in Dulbecco modified Eagle medium with 10% fetal bovine serum at 37°C with 5% CO<sub>2</sub>.<sup>29</sup>

#### **DNA constructs**

The E12 and E47 variants of the human E2A transcription factor gene were cloned into the pH $\beta$ APneo vector, which has been previously described.<sup>30</sup> The human HEB transcription factor gene was cloned in the pXS vector, which is derived from pcDL-Sra296.<sup>31</sup> The pEBB-Rag1 and pEBB-Rag2 expression vectors have been described before.<sup>32</sup>

#### Transfection protocol

Transfections were performed via calcium phosphate precipitation as described.<sup>23</sup> BOSC 23 cells were plated on the day prior to transfection at a density of  $4.5 \times 10^6$  cells per 10-cm dish. On the day of the transfection, 18 to 24 µg total DNA, including 6 µg each expression vector or carrier DNA, was used per transfection. The cells were harvested 3 days after transfection.

#### Polymerase chain reaction analysis of TCR rearrangements (coding and signal joints)

By polymerase chain reaction (PCR), 200 ng genomic DNA, isolated from the various BOSC 23–transfected cell cultures, was analyzed in a 50- $\mu$ L reaction volume containing 1 × Taq Gold buffer (Applied Biosystems, Foster City, CA), 1.5 mM MgCl<sub>2</sub>, 12.5 pmol each primer, 200  $\mu$ M deoxy–nucleoside 5' triphosphate, and 1 U AmpliTaq Gold (Applied Biosystems). PCR reactions were performed on an ABI480 machine as follows: 10 minutes preactivation at 94°C, 40 cycles of 45 seconds at 94°C, 90 seconds at 60°C, 2 minutes at 72°C, followed by a 10-minute extension

at 72°C. The primers used to detect TCRD rearrangements as well as their circular excision products locus are listed in Table 1. TCRG gene rearrangements were studied with the use of VyI-3', VyII-3', VyIII-3', VyIV-3', Jy1.1/2.1-3', Jy1.2-3', and Jy1.3/2.3-3' primers.<sup>33</sup> TCRB analysis was performed with VB2 and VB5A family primers (A.W.L., unpublished data, 2001) or DB1 and DB2 primers (T. Szczepanski, unpublished results, 2001) in combination with J $\beta$ 1(2) and J $\beta$ 2(2) primers.<sup>34</sup> Deletional rearrangements in the TCRD locus were studied with the use of 8REC-3' and  $\psi J \alpha$ -3' primers (Table 1).<sup>35</sup> In all reactions, proper positive controls for the various types of TCR rearrangements were included: well-defined leukemic cell DNA and/or total thymus DNA. We used mock-transfected BOSC 23 cells and/or HeLa genomic DNA as template containing negative controls. We analyzed 20 µL each PCR on a 2% agarose gel, followed by ethidium bromide staining. If TCR rearrangements were detected in the agarose gels, the remainder of the PCR products were subjected to heteroduplex analysis to discriminate between monoclonal and polyclonal rearrangements.<sup>36</sup> In short, heteroduplex analysis consisted of 5 minutes denaturation at 94°C and 60 minutes renaturation at 4°C prior to electrophoresis on 6% nondenaturing polyacrylamide gels (polyacrylamide to bisacrylamide, 29:1) in  $0.5 \times$  TBE buffer.<sup>36</sup> Ethidium bromide-stained homoduplex or heteroduplex PCR products were visualized with UV light.

#### Cloning of PCR products and sequencing

Following amplification, PCR products were purified by means of QIAquick PCR purification kits (Qiagen, Hilden, Germany) and cloned into pGEM-T Easy vector (Promega, Madison, WI) according to the manufacturer's instructions. Clones containing insert were sequenced on the ABI377 fluorescent sequencer, by means of the dye terminator cycle sequencing kit and Ampli*Taq* FS (Applied Biosystems).<sup>7</sup>

#### Real-time quantitative PCR of TCR rearrangements

Levels of particular rearranged *TCRD* and *TCRG* PCR products were quantified by real-time quantitative (RQ) PCR, by means of TaqMan technology on the ABI Prism 7700 Sequence Detection System (Applied Biosystems), as described earlier.<sup>37,39</sup> To this end, forward (F-DD2-KLON, F-VG8-KLON) and reverse primers (R-DD3-CONS4, R-JG13/23-KLON) (Table 1) were designed by means of Primer Express (Applied Biosystems) and Oligo6.2 (Dr W. Rychlik, Molecular Biology Insights, Cascade, CO) software to select melting temperature values of 58°C to 60°C and to exclude hairpin formation, dimer formation, and false priming. Design of the primers was performed so that the primers could be used with already present dual-labeled TaqMan probes (T-DD3-CONS2, T-JG13/23-CONS3) (Table 1). An albumin primer/probe RQ-PCR set (Applied Biosystems) was used to quantitate and normalize the amount of DNA used in the various transfections.<sup>37</sup>

#### Results

### E2A and HEB induce incomplete *TCRD* gene rearrangements in nonlymphoid cells

To study the effect of E-box proteins on recombination events in the various human TCR loci, we employed the model system described by Romanow et al.<sup>23</sup> Nonlymphoid BOSC 23 cells, which harbor their TCR loci in germline configuration, were transfected with E2A, splice variants (E12 or E47), or HEB, either alone or in combination with the RAG1/RAG2 proteins. PCR analysis of genomic DNA, isolated 3 days posttransfection, was performed by means of specific primers for the most frequently occurring types of incomplete and complete *TCRD* gene rearrangements (D $\delta$ 2-D $\delta$ 3, V $\delta$ 2-D $\delta$ 3, D $\delta$ 2-J $\delta$ 1, V $\delta$ 1-J $\delta$ 1, V $\delta$ 2-J $\delta$ 1, V $\delta$ 3-J $\delta$ 1 recombinations) (Figure 1A). Transfection of E2A or HEB alone, or mock transfection, did not result in activation of any of these rearrangements. However, D $\delta$ 2-D $\delta$ 3 and V $\delta$ 2-D $\delta$ 3 rearrangements were clearly induced upon transfection of either E2A or HEB in

Primer	Sequence (5'-3')	Reference
Coding joint PCR		
Vδ1-5′‴	ACT CAA GCC CAG TCA TCA GTA TCC	33
V82-5′	ACC AAA CAG TGC CTG TGT CAA TAG G	33
Vδ3-5′	GAC CAG ACG GTG GCG AGT GGC	33
Dδ2-5' RO2	CCA CAT TGG GAG TGT CAA CAT TT	This study
Dδ3-3' N	CTT CCT GCT ATC CCT TCC AGG	33
Dδ3-5′	CGC GTC GAC CAT ATA GTG GAA ACC GAG GGG	This study
Jδ1-3′	ACC TCT TCC CAG GAG TCC TCC	33
Jδ3-3′	CGC GTC GAC TCA AAT TAT CCC AGA AAT ATA GG	44
δREC-3′	GCA ACA TCA CTC TGT GTC TAG C	35
ψJα-3′	CCT GAA GCT TAA GGC ACA TTA GAA TCT CTC ACT G	35
Signal joint PCR		
Vδ2-sj 3′	CTG GTC AGT GGT TTT TGA GCT GCT	This study
Dô2-5' XBg	GTA GAT CTA GAA GAG GGT TTT TAT ACT GAT GTG	This study
Dδ2-lower	TCC CAA TGC TGA GAC ATA CAT	This study
D <sub>0</sub> 3-upper	CAG GGG CCA TAT AGT TGT GAA	This study
Dδ3-5' S	CGC GTC GA CCA TAT AGT GTG AAA CCG AGG GG	This study
Dδ3-3' N	CTT CCT GCT ATC CCT TCC AGG	33
Jδ1 sj 5′	GTC CCT ACC TGC AGA TGA TTA ACC	This study
Jδ3 sj 5′	CCC TTG GTC TCA TCA AGA GCA GC	This study
Real-time PCR		
F-DD2-KLON	GAA GAA GAG GGT TTT TAT ACT GAT GTG TT	This study
R-DD3-CONS4	TTG CCC CTG CAG TTT TTG TAC	This study
T-DD3-CONS2	ATA CGC ACA GTG CTA CAA AAC CTA CAG AGA CCT	This study
F-VG8-KLON	TCT ATT ACT GTG CCA CCT GGG ATA	This study
R-JG13/23-KLON	TTC CTG CCT TCC CTC TAT TAC CTT	This study
T-JG13/23-CONS3	TGT CAC AGG TAA GTA TCG GAA GAA TAC AAC ATT TCC	This study

### Table 1. Primers and probes used for polymerase chain reaction amplification of TCRD rearrangements and real-time quantitative polymerase chain reaction

PCR indicates polymerase chain reaction; sj, signal joint.

combination with the 2 *RAG* genes (Figure 1B-C). In contrast to V $\delta$ 2-D $\delta$ 3 rearrangements, D $\delta$ 2-D $\delta$ 3 recombination products were also detectable in transfectants with the 2 RAG genes only. However, quantification by RQ-PCR with the TaqMan technology revealed that the D $\delta$ 2-D $\delta$ 3 rearrangement levels were essentially higher (5- to 10-fold) in the HEB-plus-RAG transfectants than in the RAG1/RAG2–only transfectants. Further cloning and sequencing of the PCR products from the various transfectants showed heterogeneous V $\delta$ 2-D $\delta$ 3 junctional regions in all transfection combinations, with variable numbers of

deleted nucleotides at both sides and occasionally introduction of palindromic (P) nucleotides (Table 2). Strikingly, virtually all sequenced D&2-D&3 rearrangements of the transfected cells were found to be identical, showing complete deletion of the D&2 and D&3 gene segments and direct coupling of the upstream RSS of the D&2 segment to the downstream RSS of the D&3 segment (Table 2, Figure 2). This so-called signal joint, which is normally present in excision circles, was only occasionally found in normal thymocytes (Table 2, Figure 2).

In contrast to the D $\delta$ 2-D $\delta$ 3 and V $\delta$ 2-D $\delta$ 3 rearrangements, we did not





Table 2. Sequences of cloned TCRD recombination product
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Sample	5' gene segment*	P nucleotides	3' gene segment*
Vô2-Dô3 rearrangements			
Germline†	TGT GCC TGT GAC ACC		ACT GGG GGA TAC G cac agt g
E47/RR transfectant	0		-3
	0		-4
HEB/RR transfectant	-2		-3
	-1		-2
	-1		-3
	-2	т	0
	0		-3
Dδ2-Dδ3 rearrangements‡			
Germline§	c att gtg CCT TCC TAC		ACT GGG GGA TAC G cac agt g
E12/RR transfectant (4/4)#	c att gtg		cac agt g
E47/RR transfectant (10/11)#	c att gtg		cac agt g
E47/RR transfectant (1/11)#	c att gtg CCT TCC TAC		CT GGG GGA TAC G cac agt g
HEB/RR transfectant (7/7)#	c att gtg		cac agt g
Thymus	c att gtg CCT TCC TAC	N region	T GGG GGA TAC G cac agt g
	c att gtg CCT TCC TA		ACT GGG GGA TAC G cac agt g
	c att gtg		cac agt g
Dδ2-Dδ3 excision circles‡			
Germline¶	c act gtg		cac aca g
E12/RR transfectant	c act gtg ACT GGG GGA		T TCC TAC cac aca g
E47/RR transfectant	c act gtg ACT GGG GGA TAC G		C TAC cac aca g
HEB/RR transfectant	c act gtg ACT GGG GGA		TAC cac aca g
	c act gtg ACT GGG GGA TAC		cac aca g
Thymus	c act gtg		cac aca g
	c act gtg	N region	cac aca g
	c act gtg	N region	AC cac aca g
	c act	N region	cac aca g

RSS indicates recombination signal sequence; RR, RAG1 plus RAG2.

\*Indicated are the number of nucleotides deleted at the 5' and 3' gene segments. Lower case represents RSS nucleotides; upper case, nucleotides of gene segments.

 $\pm$  For 5' gene segment, this is V $\delta$ 2 (coding); for 3' gene segment, this is D $\delta$ 3 (coding + 3' RSS).

\$See Figure 2 for the D&2-D&3 rearrangement and excision circle composition, such as they occur in the classical and alternative recombination mechanisms.

§For 5' gene segment, this is D<sub>8</sub>2 (5' RSS + coding); for 3' gene segment, this is D<sub>8</sub>3 (coding + 3' RSS).

¶For 5' gene segment, this is D $\delta$ 3 (5' RSS); for 3' gene segment, this is D $\delta$ 2 (3' RSS).

#Indicated are the number of PCR products of the total number of sequenced PCR products.

detect the other 4 frequently occurring types of *TCRD* rearrangements, which all involve the J $\delta$ 1 gene segment (Table 3). D $\delta$ 2-D $\delta$ 3 recombinations can be formed only by direct rearrangement of the 2 neighboring D $\delta$ 2 and D $\delta$ 3 segments. The intermediate circular excision products or signal joints that are formed as a side product during this process (Figure



Figure 2. Scheme of classical V(D)J recombination and alternative recombination mechanisms resulting in D $\delta$ 2-D $\delta$ 3 rearrangement products and circular excision products. In classical recombination, D $\delta$ 2 and D $\delta$ 3 coding sequences are coupled to a coding joint, whereas the intervening sequence is coupled via the downstream D $\delta$ 2 RSS and the upstream D $\delta$ 3 RSS to a signal joint on the circular excision product. In the alternative mechanism that was observed in the transfected BOSC 23 cells on the basis of the sequencing of the rearrangements, a signal joint of the upstream D $\delta$ 2 RSS and the downstream D $\delta$ 3 RSS is formed, whereas the D $\delta$ 2 and D $\delta$ 3 coding sequences are deleted out on the excision circles. Primers mentioned are as follows: 1, D $\delta$ 2-5' RO2; 2, D $\delta$ 2-lower; 3, D $\delta$ 3-upper; 4, D $\delta$ 3-3' N.

3A) were indeed detected in the same transfection combinations (E2A or HEB plus RAG) in which the corresponding D&2-D&3 coding joints were found (Figure 3B). Sequencing of these excision circle products from transfected cells showed a coding joint-like configuration, confirming the less common mechanism of D82-D83 recombination (Table 2; Figure 2); similar products were occasionally found in thymocytes as well. V82-D83 rearrangements can be formed either by direct coupling or via a 2-step mechanism involving the D82-D83 joint as an intermediate. In the former case, V82-D83 signal joints should be detectable, whereas in the latter option both the D $\delta$ 2-D $\delta$ 3 signal joints as well as Vô2-Dô2 signal joints should be observed. PCR analysis did show Vô2-Dô3 signal joints in the E2A or HEB plus RAG transfectants (Figure 3A,C), whereas V82-D82 signal joints were not found, indicating that the observed V82-D83 recombination takes place via direct coupling of the 2 gene segments, rather than as a 2-step process in which  $D\delta 2$ - $D\delta 3$  coding joints act as intermediates.

### TCRD recombination involving Jô gene segments cannot be induced by E2A and HEB

From various studies, it is known that V $\delta$ -J $\delta$  rearrangements are formed via multiple (consecutive) couplings involving D $\delta$  segments, rather than as a direct joining of V $\delta$  to J $\delta$  gene segments. This even applies to D $\delta$ 2-J $\delta$ 1 joints that are known to be formed in 2 steps, given the presence of identifiable D $\delta$ 3 segment sequences in virtually all of these coding joints.<sup>40,41</sup> To determine whether E2A or HEB can induce recombination to J $\delta$ 1, we studied a rarer

Table 3.	TCRD	rearrangements	in various	E2A- or	HEB-transfected	BOSC 23 cells
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Rearrangement	E47	E47/RR	E12	E12/RR	HEB	HEB/RR	RR	BOSC-23*	Thymus
Dδ2-Dδ3	-	+	_	+	_	+	+/-	_	++
Vô2-Dô3	_	+	-	+	_	+	_	-	++
Dδ2-Jδ1	_	-	-	_	_	-	_	-	++
Vδ1-Jδ1	_	_	-	_	—	-	_	-	++
Vδ2-Jδ1	_	_	-	_	—	-	_	-	++
Võ3-Jõ1	_	_	-	_	—	-	_	-	++
Dδ2-Dδ3 signal joint	-	+	_	+	—	+	+/-	-	++
Vδ2-Dδ2 signal joint	_	_	-	_	—	-	_	-	++
Vδ2-Dδ3 signal joint	_	+	-	+	—	+	_	-	++
D83-J81	_	-	—	-	—	-	—	-	++
D83-J83	_	_	-	_	—	-	_	-	++
D <sup>8</sup> 3-J <sup>8</sup> 1 signal joint	_	_	-	_	—	-	_	-	++
Dδ3-Jδ3 signal joint	-	_	-	_	-	-	-	-	++

RR indicates RAG1 plus RAG2.

\*In some experiments, GFP-transfected cells were used as nonspecific template control.

type of TCRD rearrangement, D83-J81, which can occur only as a direct coupling and which is known to be present in human thymocytes (T. M. Breit et al, unpublished observations, 2001). Although readily detectable in thymocytes, this rearranged D83-J $\delta$ 1 product could not be detected in any of the transfected cell populations; the only PCR product apparent was the larger germline fragment encompassing the nonrearranged DS3 and JS1 segments that lie within 1 kb (Table 3). Even after nested PCR, no clear signs of D83-J81 rearrangements were found (data not shown), and also no Dδ3-Jδ1 signal joints were detectable (Table 3). To fully exclude recombination to J $\delta$  gene segments, we studied another type of TCRD rearrangement, Db3-Jb3, which is also formed in a 1-step reaction and occurs in thymocytes as well. Also this type of coupling (either coding joint or signal joint) could not be observed in any of the transfected combinations (Table 3). Collectively, these data illustrate that E2A and HEB have the ability to induce recombination in the TCRD locus, but that this concerns only 1-step rearrangements in the V $\delta$ 2-D $\delta$  region, and not in the more downstream Jδ region.

### V-J rearrangements in the *TCRG* locus are induced by E2A and HEB in cooperation with the RAG proteins

Although the TCRD locus is generally believed to be the first TCR locus that is rearranged during T-cell differentiation, we wished to ascertain whether in the transfected BOSC 23 cells recombination events would also be detectable in the TCRG locus, which starts rearranging later than TCRD during thymocyte differentiation, but earlier than the TCRB and TCRA genes.9-11 The human TCRG locus is composed of a limited set of  $V\gamma$  gene segments that are grouped in V $\gamma$  families and 5 J $\gamma$  gene segments clustered in the homologous  $J\gamma 1$  and  $J\gamma 2$  regions (Figure 4A). To analyze  $V\gamma$ - $J\gamma$  recombinations, we employed 4 V $\gamma$  family primers in combination with 3 primers known to recognize the  $J\gamma 1.1/2.1$ ,  $J\gamma 1.2$ , and  $J\gamma 1.3/2.3$ segments.  $V\gamma I$ -J $\gamma 1.3/2.3$  products were found to be induced by the basic HLH (bHLH) proteins E2A or HEB in the presence of RAG proteins as compared with RAG proteins only (Figure 4B). Similar to DS2-DS3 recombination levels, TaqMan RQ-PCR revealed an inducing effect (3- to 5-fold) in the E2A or HEB plus RAG



D<sub>8</sub>2-D<sub>8</sub>3 signal joint

Vδ2-Dδ3 signal joint

Figure 3. Analysis of intermediate excision circles or signal joints. Analysis of intermediate excision circles or signal joints illustrates that E2A and HEB induce singlestep TCRD recombinations in nonlymphoid cells. (A) In the example of a V82-D83 rearrangement, the coding and signal joints formed during direct coupling of the Vo2 and Do3 segments are shown. (B,C) Analysis of signal joints in BOSC 23 cells transfected with E47, HEB, RAG1, and RAG2 expression vectors. Genomic DNA (200 ng) of the various transfectants was used for PCR amplification by means of D<sub>8</sub>2-lower plus D<sub>8</sub>3-upper (panel B) or V<sub>8</sub>2-sj 3' plus D<sub>8</sub>3-5' S (panel C) primers that specifically detect Db2-Db3 and Vb2-Db3 signal joints, respectively. Thymus DNA was used as positive control, and GFP- or mock-transfected BOSC 23 DNA as nonspecific template control. PCR products were run on a 2% agarose gel and stained with ethidium bromide. Similar to the coding joints shown in Figure 1, Dô2-Dô3 and Vô2-Dô3 signal joints were observed upon combined transfection of either E47 or HEB together with RR. In line with the data on Do2-Do3 PCR products, D<sub>8</sub>2-D<sub>8</sub>3 signal joints were also detectable in genomic DNA derived from transfectants expressing RR only (lane RR). Importantly, transfection with 2 µg RAG expression vectors (lane RR2) did not result in detectable signal joints; this amount of RAG1/RAG2 vector leads to RAG activity levels that are more in line with the RAG activity levels in those cases in which 6 µg RAG1/RAG2 is cotransfected with E2A or HEB (W. J. Romanow, personal communication, 2001).



**Figure 4. Induction of V**<sub>Y</sub>-J<sub>Y</sub> **rearrangements by E2A or HEB in the presence of RAG1 and RAG2.** (A) Schematic diagram of the human *TCRG* locus. The human *TCRG* locus comprises 2 constant region gene segments (C<sub>Y</sub>) preceded by 2 or 3 joining gene segments (J<sub>Y</sub>) and at least 14 variable gene segments (V<sub>Y</sub>) belonging to 4 subgroups. Nine V<sub>Y</sub> gene segments (5 functional genes and 4 pseudogenes) belong to subgroup I, whereas subgroups II, III, and IV each consist of a single gene segment designated V<sub>Y</sub>9, V<sub>Y</sub>10, and V<sub>Y</sub>11, respectively. Two pseudogenes, V<sub>Y</sub>A and V<sub>Y</sub>B, located upstream of V<sub>Y</sub>9 and V<sub>Y</sub>11, respectively, belong to none of these subgroups. Five joining segments have been identified: J<sub>Y</sub>1.1, J<sub>Y</sub>1.2, and J<sub>Y</sub>1.3 upstream of C<sub>Y</sub>1; and J<sub>Y</sub>2.1 and J<sub>Y</sub>2.3 upstream of C<sub>Y</sub>2. In addition to the 6 functional V<sub>Y</sub> gene segments (solid blocks), 3 pseudogenes (shaded blocks) can also rearrange to J<sub>Y</sub> gene segments. (B) Following transfection of BOSC 23 cells, PCR amplification with 200 ng genomic DNA of the transfected cells was performed by means of V<sub>Y</sub>I-3' and J<sub>Y</sub>1.3' 2.3'-3'-specific primers. Thymus DNA and BOSC 23 DNA were used as positive and nonspecific template controls, respectively. Electrophoresis of PCR products on 2% agarose gels shows that E47 or HEB and RAG1 plus RAG2 proteins can promote V<sub>Y</sub>I-J<sub>Y</sub>1.3'/-3.2' arearrangements at high levels as compared with transfectant expressing RR only (lane RR) as quantitated by RQ TaqMan PCR analysis.

transfectants as compared with transfections with the RAG genes alone. Heteroduplex analysis of the  $V\gamma$ -J $\gamma$  PCR products to discriminate between polyclonal and clonal recombination products revealed some level of heterogeneity in the various transfection combinations (data not shown). Since the VyI gene family consists of many distinct  $V\gamma$  gene segments that can rearrange, we sequenced these  $V\gamma I$ -J $\gamma 1.3/2.3$  recombinations to study the diversity of V gene segment usage (Table 4). All (approximately 20) sequenced products were found to contain the  $J\gamma 2.3$  gene segment, which is discernible from  $J\gamma 1.3$  at a single nucleotide position. At the V side, 2 gene segments were identified:  $\psi V\gamma 7$  and  $V\gamma 8$ . Interestingly, within the  $V\gamma I$  cluster, these 2 segments are most proximal to the  $J\gamma$  segments (Figure 4A), which might explain their predominance. Rearrangements between VyIV (Vy11) and Jy1.3/ 2.3 gene segments were also observed in the E2A or HEB plus RAG transfectants, but they could be found at similar levels in the RAG1/RAG2 alone transfectants (data not shown). Apparently, the VyIV and Jy2.3 gene segments are relatively easily accessible to the action of the RAG proteins, even without the presence of lymphoid transcription factors. The position of the  $V\gamma IV$  gene segment just proximal to the  $J\gamma$  gene segments might explain the finding of the otherwise rare VyIV-Jy1.3/2.3 recombination. Never-

theless, rearrangements between the V $\gamma$ IV segment and the more proximal J $\gamma$  segments (J $\gamma$ 1.1 and J $\gamma$ 2.1) were not seen. Moreover, recombination products between any of the other V $\gamma$  gene segments and these J $\gamma$ 1.1 and J $\gamma$ 2.1 segments were not detectable at all in the BOSC 23 transfectants. The same was true for V $\gamma$ 9-J $\gamma$ 1.2 rearrangements, which are frequently found in peripheral blood TCR $\gamma$  $\delta^+$  T cells. The E2A and HEB bHLH proteins thus not only induce *TCRD* recombination, but also direct the RAG proteins to rearrange particular V and J gene segments within the *TCRG* locus.

## Absence of *TCRB* rearrangements and *TCRD* deletions in E2A and HEB transfectants

To further analyze potential effects of E2A and/or HEB on V(D)J recombination in the TCR loci that are normally rearranged in later stages of T-cell differentiation (late double-negative and immature single-positive stages), we first studied rearrangements in the human *TCRB* locus. Unlike the *TCRG* locus, the *TCRB* locus is built up of a large number of distinct V $\beta$  segments, clustered in approximately 25 V $\beta$  families, and 2 D $\beta$  segments each lying upstream of a cluster of 6 or 7 J $\beta$  gene segments. The presence of V, D, and J segments in the *TCRB* locus implies that both incomplete

	E12/RR			E47/RR			HEB/RR		
	5' del	P region	3' del	5' del	P region	3' del	5' del	P region	3' de
Vy7-Jy2.3 rearrangement									
	0	С	-4	-2		0	0	CC	-7
	-2		0				-9		-4
Vy8-Jy2.3 rearrangement									
	-3		-15	-1		0	0	С	-7
	-5		-2	-2	С	0	-2		-4
	-5		-3				-4		-4
							-4		-2
							-6		0

Table 4. Sequences of cloned TCRG recombination products

RR indicates RAG1 plus RAG2.

Indicated are the number of nucleotides deleted at the 5' and 3' gene segments

and complete rearrangements can occur. Incomplete *TCRB* rearrangements almost exclusively concern D $\beta$ -J $\beta$  recombinations, as incomplete V $\beta$ -D $\beta$  joints are rarely found in T cells. For this reason, we studied D $\beta$ 1-J $\beta$ 1, D $\beta$ 1-J $\beta$ 2, and D $\beta$ 2-J $\beta$ 2 rearrangements, which can readily be detected in thymocytes. However, we could not find either one of these products in any of the transfected cell populations, even after nested PCR reactions (data not shown). Given these results and also the results from the *TCRD* analysis concerning 2-step recombinations, we anticipated that V $\beta$ -J $\beta$  joints, which normally include D $\beta$  sequences, would not be detectable either. Employing J $\beta$ 1 or J $\beta$ 2 primers in combination with specific primers for the frequently used V $\beta$ 2 and V $\beta$ 5 families,<sup>42</sup> we indeed could not observe complete V $\beta$ -J $\beta$  joints (data not shown).

To further substantiate the absence of other mature TCR gene rearrangements, we studied *TCRD* deletional rearrangements mediated by the nonfunctional  $\delta REC$  and  $\psi J\alpha$  segments, which flank the coding elements of the *TCRD* locus (Figure 1). These *TCRD* deletions precede rearrangements in the human *TCRA* locus, and they are known to occur at a relatively late stage in T-cell differentiation. However, no  $\delta REC - \psi J\alpha$  rearrangements were identified in transfected BOSC 23 cells (data not shown). Taken together, these data do not provide any evidence for a role for E2A or HEB in inducing *TCRB* rearrangements or *TCRD* deletional rearrangements.

#### Discussion

In this study, we provide evidence that the bHLH transcription factors E2A and HEB play a role in the induction of V(D)J recombination in human TCRD and TCRG loci, employing a transfection-based model of nonlymphoid cells. E2A and HEB were found to induce immature D82-D83 and V82-D83 rearrangements upon cotransfection with RAG proteins in nonlymphoid cells. Rearrangements of VyI family (especially  $\psi$ Vy7 and Vy8) gene segments to  $J\gamma 2.3$  were also easily induced by the action of both transcription factors. All except the D&2-D&3 rearrangements were very similar to the majority of rearrangements found in lymphoid cells, with junctional regions characterized by nucleotide deletion and occasionally P nucleotides. The lack of N regions can be explained by the absence of terminal deoxynucleotidyl transferase, which mediates nontemplated nucleotide insertion. Remarkably, the junctions of virtually all D82-D83 rearrangements from the transfected cells were found to be identical, because of an unusual direct coupling of the intact upstream RSS of the D82 segment to the intact downstream RSS of the D83 segment. This unusual type of rearrangement shares similarity to atypical IGK gene recombinations between JK segments and an isolated RSSlike sequence in the J-Ck intron.43 In fact, both types of recombinations represent signal joints, which normally are not found on the

Table 5. Detection of cloned TCRD, cloned TCRG, and cloned TCRB gene rearrangements in E2A/HEB-transfected cells as compared with human thymocytes and acute lymphoblastic leukemias

	E2A/HEB	E2A/HEB Thymocytes				T-ALL, %*		
Rearrangement	transfectants	CD34+ 1a-	CD34+ 1a+	Total	CD3-	$TCR\gamma\delta^+$	$TCR\alpha\beta^+$	Precursor B-ALL, %*
TCRD								
D82-D83	+	++	++	+	4	_	_	13
Vδ2-Dδ3	+	+	+	+	6	4	_	67
Dδ2-Jδ1	-	+	++	+	10	15	7	_
Vδ1-Jδ1	-	-	+	+	24	56	13	_
Vδ2-Jδ1	_	-	+	+	10	6	20	_
Vδ3-Jδ1	-	-	+	+	5	8	7	_
Other	NT	NT	NT	NT	40	10	53	20
TCRG								
VγI-Jγ1.1/2.1	_	NT	NT	+	8	9	20	18
VγII-Jγ1.1/2.1	_	NT	NT	+	_	_	_	1
VγIII-Jγ1.1/2.1	_	NT	NT	+	1	_	1	_
VγIV-Jγ1.1/2.1	_	NT	NT	+	1	4	_	1
VγI-Jγ1.2	-	NT	NT	+	_	_	_	1
VγII-Jγ1.2	_	NT	NT	+	2	_	_	_
VγIII-Jγ1.2	NT	NT	NT	+	_	_	_	_
VγIV-Jγ1.2	NT	NT	NT	+	_	_	_	_
VγI-Jγ1.3/2.3	+	-	+	+	69	52	58	56
VγII-Jγ1.3/2.3	-	-	+	+	7	24	6	18
VγIII-Jγ1.3/2.3	-	-	+	+	4	7	9	_
VγIV-Jγ1.3/2.3	+/-	-	+	+	4	4	5	5
Other	NT	NT	NT	NT	4	1	_	_
TCRB								
Dβ1-Jβ1	-	-	+	+	12	16	9	_
Dβ1-Jβ2	-	-	+	+	4	14	3	16
Dβ2-Jβ2	-	-	+	+	11	5	17	17
Vβ-Jβ1	-	-	_	+	18	13	28	_
Vβ-Jβ2	_	-	_	+	47	45	30	53
Other	NT	NT	NT	NT	9	7	13	14

Data on thymocytes and acute lymphoblastic leukemia compiled from Southern blot and/or polymerase chain reaction data from this study and earlier reports from our laboratory.<sup>6,11,44,45,47</sup>

T-ALL indicates T-lineage acute lymphoblastic leukemia; B-ALL, B-lineage ALL; TCR, T-cell receptor; *TCRD*, TCRδ; *TCRG*, TCRγ; *TCRB*, TCRβ; NT, not tested. \*Frequencies defined as percentage of detected *TCRD* rearrangements (on the basis of TCRDJ1 probe hybridization), percentage of detected *TCRG* rearrangements, and percentage of *TCRB* rearrangements (involving both the Jβ1 and Jβ2 regions). genome, but rather on the excision circles that are formed during V(D)J recombination (see Figure 2).

The virtually identical patterns of TCR gene recombinations in both the E2A plus RAG- and the HEB plus RAG-transfected cells showed a striking similarity to the most immature types of rearrangements seen in human thymocyte subpopulations (Table 5). D $\delta$ 2-D $\delta$ 3 and V $\delta$ 2-D $\delta$ 3 rearrangements represent the earliest TCRD rearrangements in thymocytes, being clearly detectable via Southern blot analysis next to Dô2-Jô1 rearrangements in sorted cells of the most immature CD34<sup>+</sup>/CD1a<sup>-</sup> subset.<sup>11</sup> In the CD34<sup>+</sup>/ CD1a<sup>+</sup> fraction, complete V $\delta$ 1-J $\delta$ 1 and V $\delta$ 2-J $\delta$ 1 rearrangements become more prominent. However, rearrangements to  $J\delta$  segments, even simple single-step couplings like D $\delta$ 3-J $\delta$ 1 and D $\delta$ 3-J $\delta$ 3 as also seen in thymus, could not be induced in the transfected cells. TCRG rearrangements involving Jy1.3/2.3 segments and TCRB rearrangements (most probably, incomplete DB-JB joinings) were clearly detectable only in CD34<sup>+</sup>/CD1a<sup>+</sup> cells (Table 5), although low levels of (other) TCRD, TCRG, and/or TCRB recombinations in the CD34<sup>+</sup>/CD1a<sup>-</sup> subset cannot be fully excluded. The TCRD rearrangement pattern of the transfectants was also compared with TCR recombination data in human T-ALL (CD3<sup>-</sup>, TCR $\alpha\beta^+$ , and TCR $\gamma\delta^+$ ) and precursor B-ALL samples, as investigated in our laboratory (Table 5).6,44-47 In T-ALL, Jδ recombinations are predominant, especially in CD3<sup>+</sup> T-ALL, whereas D82-D83 and V82-D83 rearrangements are found only at low frequencies in immature CD3<sup>-</sup>T-ALL.<sup>6,41,44</sup> Interestingly, these latter 2 TCRD rearrangements are very characteristic in precursor B-ALL, where they constitute approximately 80% of the identified (cross-lineage) TCRD recombinations.<sup>6,44</sup> Rearrangements involving Jô have not been found in precursor B-ALL.<sup>6,44,48</sup> The data thus suggest that E2A or HEB can induce the very early TCRD recombinations that carry a less T-cell-specific character, as they are also readily observed in precursor B-ALL (Figure 5).

In the transfected cells,  $V\gamma I$ - $J\gamma 1.3/2.3$  rearrangements were readily observed; such recombinations are also frequent in both T-ALL and precursor B-ALL, albeit in the latter  $J\gamma 2.3$  is less frequently involved than  $J\gamma 1.3$ ;  $V\gamma IV$ - $J\gamma 1.3/2.3$  couplings are less prominent in both ALL subsets. Finally, incomplete D $\beta$ -J $\beta$  and complete V $\beta$ -J $\beta$  recombinations were not detected in the transfectants, but occur in all 3 T-ALL subtypes and at a lower frequency in human precursor B-cell leukemias (approximately 35% of cases).<sup>6</sup> Together, these data illustrate that the TCR gene recombination pattern as observed in the E2A or HEB plus RAG–transfected cells concerns the more immature types of TCR recombinations, as observed in CD34<sup>+</sup>/CD1a<sup>-</sup> thymocytes and also in precursor B-ALL, with the exception of the lack of *TCRB* rearrangements (Figure 5).



Figure 5. Compilation of *TCRD* recombination patterns. Indicated are the parts of the *TCRD* locus that can rearrange in the E2A and HEB transfectants described in this study, CD34<sup>+</sup>/CD1a<sup>-</sup> and CD34<sup>+</sup>/CD1a<sup>+</sup> thymocyte subsets, CD3<sup>-</sup> and TCR<sub>γ</sub>δ and TCR<sub>α</sub>β T-ALL, and precursor B-ALL. The pattern observed in the E2A/HEB-plus-RR–transfected cells is strikingly similar to that of the most immature thymocytes and the cross-lineage pattern of precursor B-ALL, illustrating that E2A and HEB can induce the immature and less T-cell–specific rearrangements in the *TCRD* locus.

Although recombination can be induced relatively easily in this transfection system, the efficiency seems to be too low to allow detection of recombination events that occur in 2 consecutive steps. This is best illustrated by our study on circular intermediate products that are formed during recombination. Upon study of V82-D83 recombinations, we exclusively found V82-D83 signal joints and no V82-D82 plus D82-D83 circular intermediates. Although we cannot formally exclude that, eg, 2-step V $\delta$ 1-J $\delta$ 1 or V82-J81 rearrangements can be induced at very low levels, the fact that the single-step  $D\delta 3$ -J $\delta 1$  and  $D\delta 3$ -J $\delta 3$  recombinations are also undetectable strengthens our view that Jo1 rearrangements are truly absent in E2A- or HEB-transfected cells rather than being an artifact in the model system. The same seems to be true for  $D\beta$ -J $\beta$ and V $\beta$ -J $\beta$  rearrangements, as well as  $\delta$ REC- $\psi$ J $\alpha$  TCRD deletional rearrangements. The TCRD recombination data thus suggest that E2A and HEB are able to regulate recombination of, especially, the Db3 gene segment, but that other factors are required to open up the downstream region of the J $\delta$  segments to the V(D)J recombinase. The area between D $\delta$ 3 and the J $\delta$  region might harbor important regulatory elements that can bind factors mediating the truly T-cell-specific types of TCRD recombinations.49 Interestingly, Vô-Dô intermediates are lacking in E2A<sup>-/-</sup> mice, but Dô-Jô intermediates do still occur,15 which indeed suggests the involvement of one or more yet unknown transcription factors. From E2A<sup>-/-</sup> mice, it is further deduced that E2A plays a role in regulation of  $V\gamma/V\delta$  subsets during ontogeny by activating and repressing V segments.15 These data fit perfectly with our model system, in which E2A can induce rearrangements of specific  $V\gamma$ gene segments and V $\delta$ -D $\delta$  rearrangements, but not incomplete  $D\delta$ -J $\delta$  or complete V $\delta$ -J $\delta$  rearrangements.

From the initial V(D)J recombination study in this nonlymphoid model system, it is known that induction of recombination by, eg, E2A is critically dependent on activation domains.<sup>23</sup> This has led to the hypothesis that E-box proteins such as E2A are involved in regulating chromatin accessibility by relieving the repressive effect of nucleosomes, most probably through recruitment of complexes containing histone acetyl transferase activity.<sup>23,50</sup> We therefore hypothesize that the interaction of E2A and HEB with chromatin leads to accessibility of RSS sequences of TCR loci as well, which is supported by the observation of  $V\gamma I$  and  $V\delta 2$  germline transcripts upon E2A or HEB transfection (J. K. Ghosh et al, personal communication, 2001). The proximity of gene segments, as in the case of D $\delta$ 2 and D $\delta$ 3 and also  $\psi V\gamma$ 7 and V $\gamma$ 8 within the V $\gamma$ I family, might be another relevant factor. Finally, the exact nucleotide sequence of the RSS might also influence recombination, although the RSSs of, eg, the various VyI family gene segments are equally perfect in that respect.

The observations in this study raise the issue of how E2A and HEB are involved in regulating TCR recombination during in vivo T-cell differentiation. Both E2A and HEB knock-out mice show a block, though not complete, in the early double-negative stages of T-cell differentiation. The phenotype of these mice thus illustrates the important role of both E2A and HEB in the earliest phases of T-cell differentiation, although the incompleteness of the differentiation arrest indicates some degree of redundancy by other regulatory factors. Using the in vitro model, we show that some types of immature TCR rearrangements can be induced by E2A or HEB, whereas the absence of these rearrangements in E2A knock-out mice shows that E2A and HEB are probably also involved in stimulating the immature TCR recombination in vivo. For induction of further (more mature) rearrangements, additional cofactors or transcription factors might be required or negative

regulatory elements might have to be downregulated at certain differentiation stages. Moreover, we cannot fully exclude a cooperative action of both E2A and HEB in particular recombinations in vivo, although we have not seen a synergistic effect of combined transfection of E2A and HEB with RAG proteins (data not shown).

Besides mediating physiological immunoglobulin/TCR gene rearrangements, V(D)J recombination is also thought to be involved in the formation of particular chromosome aberrations in human leukemias.<sup>25,26</sup> This especially concerns chromosome aberrations in T-ALL, in which TCRB or TCRD gene segments and their regulatory elements are translocated to oncogenes on partner chromosomes, resulting in activation of the oncogenes. Examples include translocations t(1;14) and t(11;14), involving the genes encoding the TAL1 and LMO1/LMO2 transcription factors, respectively.<sup>25,51</sup> As the breakpoint regions on these chromosomes have been found to be located near RSS or RSS-like sequences,<sup>25-27</sup> it is suggestive that accessibility of the involved oncogenes to RAG protein activity might be a critical step in the formation of these chromosome aberrations as well. We therefore also studied TAL1 deletions, which are V(D)J-like rearrangements that are found exclusively in T-ALL, particularly in T-ALL with TCRD deletions.52 However, such RSSmediated TAL1 deletions were not induced upon E2A or HEB transfection (data not shown). Given the relatively low frequency of V(D)J recombinations in nonlymphoid cells, it can be argued that detection of *TAL1* deletions in this model system would be very difficult. However, it might also be that transcription factors other than E2A and HEB regulate accessibility of the *TAL1* gene and that an aberrant combination of regulatory factors is required to induce these unwanted oncogenic *TAL1* deletions.

In summary, our data demonstrate a role for the bHLH proteins E2A and HEB in induction of TCR rearrangements. Comparison with the TCR-rearrangement patterns found in thymocyte subsets and in T- and B-lineage leukemias indicates that the identified rearrangements in, especially, the *TCRD* locus are immature and carry a less T-cell–specific character (Figure 5). A challenge for future studies remains the identification of factors that are involved in induction of more T-cell–specific TCR rearrangements, such as (complete) *TCRD* recombinations involving J $\delta$  couplings and/or *TCRB* recombinations. Furthermore, it will be interesting to see whether the here-described model system will be helpful in dissecting the molecular processes regulating RSS-mediated chromosome aberrations in human lymphoid leukemias.

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#### References

- 1. Tonegawa S. Somatic generation of antibody diversity. Nature. 1983;302:575-581.
- Schatz DG, Oettinger MA, Baltimore D. The V(D)J recombination activating gene, RAG-1. Cell. 1989;59:1035-1048.
- Oettinger MA, Schatz DG, Gorka C, Baltimore D. RAG-1 and RAG-2, adjacent genes that synergistically activate V(D)J recombination. Science. 1990;248:1517-1523.
- van Gent DC, McBlane JF, Ramsden DA, Sadofsky MJ, Hesse JE, Gellert M. Initiation of V(D)J recombination in a cell-free system. Cell. 1995; 81:925-934.
- Sleckman BP, Gorman JR, Alt FW. Accessibility control of antigen-receptor variable-region gene assembly: role of cis-acting elements. Annu Rev Immunol. 1996;14:459-481.
- Szczepanski T, Beishuizen A, Pongers-Willemse MJ, et al. Cross-lineage T-cell receptor gene rearrangements occur in more than ninety percent of childhood precursor-B-acute lymphoblastic leukemias: alternative PCR targets for detection of minimal residual disease. Leukemia. 1999;13: 196-205.
- Szczepanski T, Pongers-Willemse MJ, Langerak AW, et al. Ig heavy chain gene rearrangements in T-cell acute lymphoblastic leukemia exhibit predominant DH6–19 and DH7–27 gene usage, can result in complete V-D-J rearrangements, and are rare in T-cell receptor alpha beta lineage. Blood. 1999;93:4079-4085.
- Yancopoulos GD, Alt FW. Developmentally controlled and tissue-specific expression of unrearranged VH gene segments. Cell. 1985;40:271-281.
- Van Dongen JJM, Wolvers-Tettero ILM. Analysis of immunoglobulin and T-cell receptor genes, II: possibilities and limitations in the diagnosis and management of lymphoproliferative diseases and related disorders. Clin Chim Acta. 1991;198:93-174.
- Ktorza S, Blanc C, Laurent C, et al. Complete TCR-delta rearrangements and partial (D-J) recombination of the TCR-beta locus in CD34+7+ precursors from human cord blood. J Immunol. 1996;156:4120-4127.

- Blom B, Verschuren MCM, Heemskerk MHM, et al. TCR gene rearrangements and expression of the pre-T cell receptor complex during human T-cell differentiation. Blood. 1999;93:3033-3043.
- Alt FW, Yancopoulos GD, Blackwell TK, et al. Ordered rearrangement of immunoglobulin heavy chain variable region segments. EMBO J. 1984;3: 1209-1219.
- 13. Strominger JL. Developmental biology of T cell receptors. Science. 1989;244:943-950.
- Raulet DH, Garman RD, Saito H, Tonegawa S. Developmental regulation of T-cell receptor gene expression. Nature. 1985;314:103-107.
- Bain G, Romanow WJ, Albers K, Havran WL, Murre C. Positive and negative regulation of V(D)J recombination by the E2A proteins. J Exp Med. 1999;189:289-300.
- Stanhope-Baker P, Hudson KM, Shaffer AL, Constantinescu A, Schlissel MS. Cell type-specific chromatin structure determines the targeting of V(D)J recombinase activity in vitro. Cell. 1996;85: 887-897.
- Murre C, McCaw PS, Baltimore D. A new DNA binding and dimerization motif in immunoglobulin enhancer binding, daughterless, MyoD, and myc proteins. Cell. 1989;56:777-783.
- Bain G, Gruenwald S, Murre C. E2A and E2–2 are subunits of B-cell-specific E2-box DNA-binding proteins. Mol Cell Biol. 1993;13:3522-3529.
- Shen CP, Kadesch T. B-cell-specific DNA binding by an E47 homodimer. Mol Cell Biol. 1995;15: 4518-4524.
- Bain G, Maandag EC, Izon DJ, et al. E2A proteins are required for proper B cell development and initiation of immunoglobulin gene rearrangements [comment appears in Cell. 1994;79:751-753]. Cell. 1994;79:885-892.
- Zhuang Y, Soriano P, Weintraub H. The helixloop-helix gene E2A is required for B cell formation. Cell. 1994;79:875-884.
- Schlissel M, Voronova A, Baltimore D. Helix-loophelix transcription factor E47 activates germ-line immunoglobulin heavy-chain gene transcription and rearrangement in a pre-T-cell line. Genes Dev. 1991;5:1367-1376.

- Romanow WJ, Langerak AW, Goebel P, et al. E2A and EBF act in synergy with the V(D)J recombinase to generate a diverse immunoglobulin repertoire in nonlymphoid cells. Mol Cell. 2000;5: 343-353.
- Zhuang Y, Cheng P, Weintraub H. B-lymphocyte development is regulated by the combined dosage of three basic helix-loop-helix genes, E2A, E2–2, and HEB. Mol Cell Biol. 1996;16:2898-2905.
- 25. Rabbitts TH. Translocations, master genes, and differences between the origins of acute and chronic leukemias. Cell. 1991;67:641-644.
- Schatz DG, Oettinger MA, Schlissel MS. V(D)J recombination: molecular biology and regulation. Annu Rev Immunol. 1992;10:359-383.
- Tycko B, Sklar J. Chromosomal translocations in lymphoid neoplasia: a reappraisal of the recombinase model. Cancer Cells. 1990;2:1-8.
- Hwang LY, Baer RJ. The role of chromosome translocations in T cell acute leukemia. Curr Opin Immunol. 1995;7:659-664.
- Pear WS, Nolan GP, Scott ML, Baltimore D. Production of high-titer helper-free retroviruses by transient transfection. Proc Natl Acad Sci U S A. 1993;90:8392-8396.
- Kee BL, Murre C. Induction of early B cell factor (EBF) and multiple B lineage genes by the basic helix-loop-helix transcription factor E12. J Exp Med. 1998;188:699-713.
- 31. Takebe Y, Seiki M, Fujisawa J, et al. SR alpha promoter: an efficient and versatile mammalian cDNA expression system composed of the simian virus 40 early promoter and the R-U5 segment of human T-cell leukemia virus type 1 long terminal repeat. Mol Cell Biol. 1988;8:466-472.
- Roman CA, Cherry SR, Baltimore D. Complementation of V(D)J recombination deficiency in RAG-1(-/-) B cells reveals a requirement for novel elements in the N-terminus of RAG-1. Immunity. 1997;7:13-24.
- 33. Pongers-Willemse MJ, Seriu T, Stolz F, et al. Primers and protocols for standardized detection

of minimal residual disease in acute lymphoblastic leukemia using immunoglobulin and T cell receptor gene rearrangements and *TAL1* deletions as PCR targets: report of the BIOMED-1 CON-CERTED ACTION: investigation of minimal residual disease in acute leukemia. Leukemia. 1999;13:110-118.

- Kneba M, Bolz I, Linke B, Hiddemann W. Analysis of rearranged T-cell receptor beta-chain genes by polymerase chain reaction (PCR) DNA sequencing and automated high resolution PCR fragment analysis. Blood. 1995;86:3930-3937.
- Breit TM, Verschuren MCM, Wolvers-Tettero ILM, Van Gastel-Mol EJ, Hählen K, van Dongen JJM. Human T cell leukemias with continuous V(D)J recombinase activity for TCR-delta gene deletion. J Immunol. 1997;159:4341-4349.
- Langerak AW, Szczepanski T, van der Burg M, Wolvers-Tettero ILM, van Dongen JJM. Heteroduplex PCR analysis of rearranged T cell receptor genes for clonality assessment in suspect T cell proliferations. Leukemia. 1997;11:2192-2199.
- Pongers-Willemse MJ, Verhagen OJHM, Tibbe GJM, et al. Real-time quantitative PCR for the detection of minimal residual disease in acute lymphoblastic leukemia using junctional regions specific TaqMan probes. Leukemia. 1998;12: 2006-2014.
- Bruggemann M, Droese J, Bolz I, et al. Improved assessment of minimal residual disease in B cell malignancies using fluorogenic consensus probes for real-time quantitative PCR. Leukemia. 2000;14:1419-1425.
- Verhagen OJ, Willemse MJ, Breunis WB, et al. Application of germline IGH probes in real-time quantitative PCR for the detection of minimal re-

sidual disease in acute lymphoblastic leukemia. Leukemia. 2000;14:1426-1435.

- Breit TM, Wolvers-Tettero ILM, Hählen K, van Wering ER, van Dongen JJM. Extensive junctional diversity of γδ T-cell receptors expressed by T-cell acute lymphoblastic leukemias: implications for the detection of minimal residual disease. Leukemia. 1991;5:1076-1086.
- Langerak AW, Wolvers-Tettero ILM, van den Beemd MWM, et al. Immunophenotypic and immunogenotypic characteristics of TCR<sub>γ</sub>δ<sup>+</sup> T cell acute lymphoblastic leukemia. Leukemia. 1999; 13:206-214.
- Van den Beernd R, Boor PPC, van Lochem EG, et al. Flow cytometric analysis of the Vbeta repertoire in healthy controls. Cytometry. 2000;40:336-345.
- Feddersen RM, Martin DJ, Van Ness BG. Novel recombinations of the IG kappa-locus that result in allelic exclusion. J Immunol. 1990;145:745-750.
- Breit TM, Wolvers-Tettero ILM, Beishuizen A, Verhoeven M-AJ, van Wering ER, van Dongen JJM. Southern blot patterns, frequencies and junctional diversity of T-cell receptor-δ gene rearrangements in acute lymphoblastic leukemia. Blood. 1993;82:3063-3074.
- Langerak AW, Wolvers-Tettero ILM, van Dongen JJM. Detection of T cell receptor beta (TCRB) gene rearrangement patterns in T cell malignancies by Southern blot analysis. Leukemia. 1999; 13:965-974.
- Moreau E, Langerak AW, van Gastel-Mol EJ, et al. Easy detection of all T cell receptor gamma (TCRG) gene rearrangements by Southern blot

analysis: recommendations for optimal results. Leukemia. 1999;13:1620-1626.

- 47. Szczepanski T, Langerak AW, Willemse MJ, Wolvers-Tettero ILM, van Wering ER, van Dongen JJM. T cell receptor gamma (TCRG) gene rearrangements in T cell acute lymphoblastic leukemia reflect "end-stage" recombinations: implications for minimal residual disease monitoring. Leukemia. 2000;14:1208-1214.
- Yokota S, Hansen-Hagge TE, Bartram CR. T-cell receptor delta gene recombination in common acute lymphoblastic leukemia: preferential usage of V delta 2 and frequent involvement of the J alpha cluster. Blood. 1991;77:141-148.
- Krangel MS, Band H, Hata S, McLean J, Brenner MB. Structurally divergent human T cell receptor γ proteins encoded by distinct Cγ genes. Science. 1987;237:64-67.
- Massari ME, Grant PA, Pray-Grant MG, Berger SL, Workman JL, Murre C. A conserved motif present in a class of helix-loop-helix proteins activates transcription by direct recruitment of the SAGA complex. Mol Cell. 1999;4:63-73.
- Rabbitts TH, Lefranc MP, Stinson MA, et al. The chromosomal location of T-cell receptor genes and a T-cell rearranging gene: possible correlation with specific translocations in human T-cell leukemia. EMBO J. 1985;4:1461-1465.
- Breit TM, Mol EJ, Wolvers-Tettero IL, Ludwig WD, van Wering ER, van Dongen JJ. Site-specific deletions involving the tal-1 and sil genes are restricted to cells of the T cell receptor alpha/beta lineage: T cell receptor delta gene deletion mechanism affects multiple genes. J Exp Med. 1993;177:965-977.