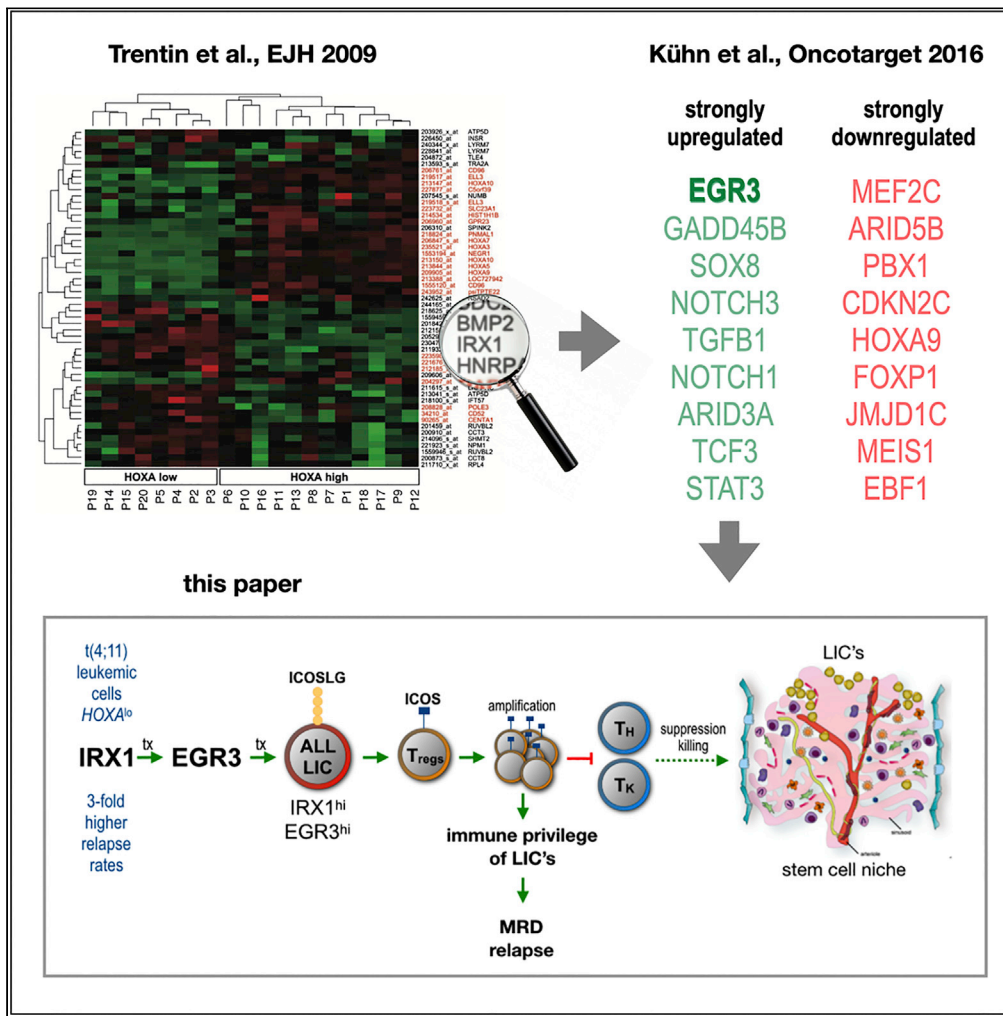


Article

The immune checkpoint ICOSLG is a relapse-predicting biomarker and therapeutic target in infant t(4;11) acute lymphoblastic leukemia



Marius Külp, Anna Lena Siemund, Patrizia Larghero, ..., Halvard Bonig, Claus Meyer, Rolf Marschalek

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Highlights

Early growth response 3 (EGR3) is a direct transactivator of the immune checkpoint gene ICOSLG

high ICOSLG expression at diagnosis is predictive for ALL relapse

EGR3 and ICOSLG expressions are relapse-associated

expression of ICOSLG on t(4;11) ALL cells leads to the rapid expansion of Tregs



## Article

## The immune checkpoint ICOSLG is a relapse-predicting biomarker and therapeutic target in infant t(4;11) acute lymphoblastic leukemia

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## SUMMARY

**The most frequent genetic aberration leading to infant ALL (iALL) is the chromosomal translocation t(4;11), generating the fusion oncogenes *KMT2A:AFF1* and *AFF1:KMT2A*, respectively. *KMT2A-r* iALL displays a dismal prognosis through high relapse rates and relapse-associated mortality. Relapse occurs frequently despite ongoing chemotherapy and without the accumulation of secondary mutations. A rational explanation for the observed chemo-resistance and satisfactory treatment options remain to be elucidated. We found that elevated *ICOSLG* expression level at diagnosis was associated with inferior event free survival (EFS) in a cohort of 43 patients with t(4;-11) iALL and that a cohort of 18 patients with iALL at relapse displayed strongly increased *ICOSLG* expression. Furthermore, co-culturing t(4;11) ALL cells (*ICOSLG*<sup>hi</sup>) with primary T-cells resulted in the development of T<sub>regs</sub>. This was impaired through treatment with a neutralizing *ICOSLG* antibody. These findings imply *ICOSLG* (1) as a relapse-predicting biomarker, and (2) as a therapeutic target involved in a potential immune evasion relapse-mechanism of infant t(4;11) ALL.**

## INTRODUCTION

*KMT2A*-rearrangements (*KMT2A-r*) occur in 70-80% of all infant acute lymphoblastic leukemias (iALL) (Pieters et al., 2007, 2019). Among these, the translocation t(4;11) generating the fusion oncogenes *KMT2A::AFF1* and *AFF1::KMT2A* is the most frequent, accounting for about 49% of all *KMT2A-r* iALL cases (Meyer et al., 2018). *KMT2A-r* is a negative risk factor in iALL with a 4-year event-free survival (4-year-EFS) of only 37% among *KMT2A-r* cases and 36% among t(4;11) cases, respectively. By comparison, patients with *KMT2A*-germline achieve a 4-year-EFS of 74% (Pieters et al., 2007). The typical clinical picture of *KMT2A-r* iALL is that patients initially reach complete remission (CR) during induction therapy but relapse months later despite continuous chemotherapy (Pieters et al., 2007). This results in a 3-year post-relapse overall survival (3-year-OS) of only 17% for patients with t(4;11), with OS defined as the time from relapse to death (Driessen et al., 2016). Designated studies indicate the absence of significant accumulation or selection of secondary mutations in the dominant leukemic clone between the timepoints of diagnosis and relapse (Andersson et al., 2015; Agraz-Doblas et al., 2019). Thus, an explanation for how therapy resistance is achieved remains an open question.

Our group previously identified the C2H2 zinc finger transcription factors *Early growth response 1, 2, and 3* (*EGR1, EGR2, EGR3*) as downstream mediators of the transcription factor *Iroquois homeobox 1* (*IRX1*) (K  hn et al., 2016). Relatively higher transcription of *IRX1* was observed in patients with iALL presenting a low *HOXA* gene expression (Trentin et al., 2009). This *IRX1*<sup>hi</sup>/*HOXA*<sup>lo</sup> patient group displayed an increased relapse rate with dismal outcome (Stam et al., 2010; Kang et al., 2012). *EGR1, EGR2, EGR3*, and their murine homologs have been described to be involved in cell cycle inhibition and stemness of hematopoietic stem cells (HSC) (Min et al., 2008; Cheng et al., 2015; Li et al., 2017). Additionally, *EGR3* expression in murine T- and B-cells is known to restrict immune activity and prevent autoimmunity (Safford et al., 2005; Anderson et al., 2006; Collins et al., 2008; Li et al., 2012; Parkinson et al., 2014).

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Continued



We were able to demonstrate that the zinc finger transcription factor *Early growth response 3* (EGR3) binds directly to the promoter of the *ICOSLG* gene and upregulates ICOSLG protein. ICOSLG is an immune checkpoint expressed by antigen-presenting cells (APC) as well as non-lymphoid cells including mesenchymal stem cells, endothelial cells, fibroblasts, and tumor cells (Swallow et al., 1999; Khayyamian et al., 2002; Martin-Orozco et al., 2010; Zhang et al., 2016). ICOSLG belongs to the B7 family of costimulatory immune receptors with partial sequence identity of CD80/CD86 and binds ICOS (Coyle and Gutierrez-Ramos, 2001). ICOS belongs to the CD28 superfamily of immune receptors, is expressed at low levels in resting memory T-cell subsets, but becomes upregulated through ICOSLG binding (Hutloff et al., 1999; Khayyamian et al., 2002). The interaction between ICOSLG and ICOS has been shown to lead to the development of T<sub>regs</sub> in healthy bone marrow as well as in several tumor microenvironments with clinical implications in melanoma, glioblastoma, breast cancer and acute myeloid leukemia (AML) (Martin-Orozco et al., 2010; Faget et al., 2012; Lee et al., 2017; Han et al., 2018; Iwata et al., 2019).

In summary, we report here that high gene expression of *ICOSLG* at initial diagnosis was associated with inferior EFS in a cohort of 43 infant patients with t(4;11) ALL and that a cohort of 18 patients with infant ALL (iALL) at relapse displayed strongly increased *EGR3* and *ICOSLG* transcription levels. Upregulation of *ICOSLG* expression in t(4;11) ALL cells caused increased development of regulatory T-cells (T<sub>regs</sub>) when co-cultured with primary T-cells. The development of T<sub>regs</sub> was impaired through the treatment with a neutralizing  $\alpha$ -ICOSLG antibody.

Thereby, we provide a potential relapse mechanism that could explain how KMT2A-r leukemia cells reach therapy resistance. Consequently, our study not only suggests *ICOSLG* as a relapse-predicting prognostic biomarker, but identifies the ICOSLG/ICOS interaction as a putative therapeutic target in infant t(4;11) ALL.

## RESULTS

### Early growth response 3 upregulates *ICOSLG* through direct promoter binding in an early growth response 3-overexpressing t(4;11) SEM cell model

The initial observation that *IRX1* is upregulated in patients with t(4;11) iALL that display a *HOXA*<sup>lo</sup> signature prompted us to investigate the role of *IRX1* in further detail. Overexpression of *IRX1* resulted in the transcriptional upregulation of *HOXB4* and the *Early growth response genes EGR1-3* (Kühn et al., 2016). We cloned the *EGR3* open reading frame (ORF) fused to a C-terminal FLAG tag into our sleeping beauty vector system (Kowarz et al., 2015). This vector construct was stably integrated into the genome of the t(4;11) cell line SEM (SEM::EGR3), and transgene expression was induced by the addition of Doxycycline. An empty vector was used for the generation of a stable control cell line (SEM::mock). This cell culture model system enabled us to investigate the effects of *EGR3* overexpression in a t(4;11) pro-B cellular context. Quantitative real-time PCR (qPCR) confirmed transgene expression and revealed strong upregulation of *ICOSLG* by *EGR3* ( $\Delta\Delta C_T = 3.24 \pm 0.16$ ) at the transcription level (Figure 1A). We used Western blot experiments to confirm the *EGR3* protein expression and the *ICOSLG* upregulation on the protein level (Figure 1B).

Considering that *EGR3* is a zinc finger transcription factor we aimed to investigate whether *EGR3* binds directly to the *ICOSLG* promoter or acts as an indirect *ICOSLG* activator. For that purpose, we performed chromatin immunoprecipitation (ChIP) followed by qRT-PCR and ChIP-Seq. ChIP-qRT-PCR and percent input calculation of the SEM::EGR3 cell model demonstrated that *EGR3* binds directly to the promoter of the *ICOSLG* gene (Figure 1C). This initial result was later corroborated by ChIP-Seq (Figure 1D).

### Iroquois homeobox 1, early growth response 3, and *ICOSLG* expressions correlate strongly in a cohort of 50 infant patients with t(4;11) pro-B ALL

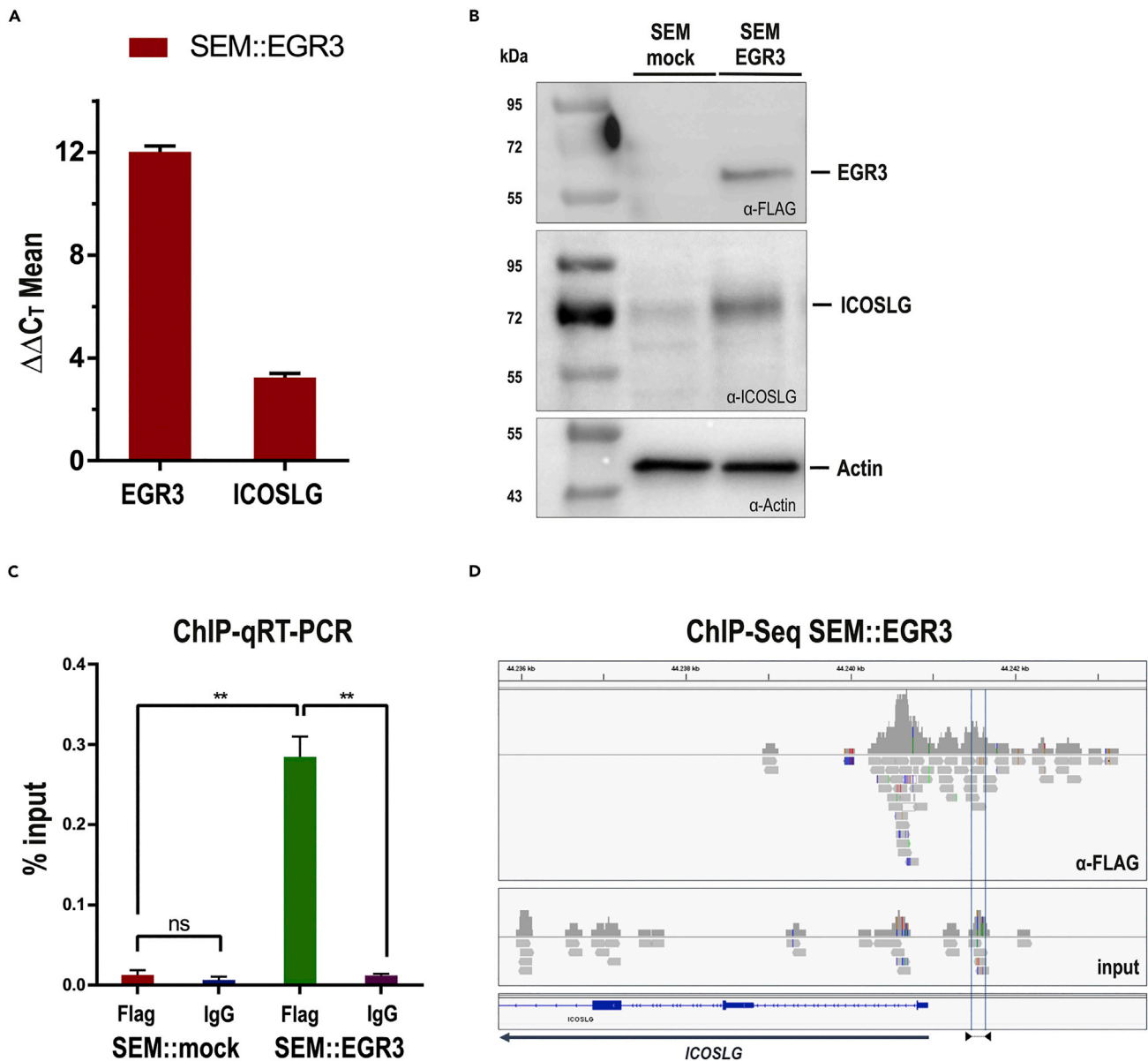
In order to validate the findings of our experimental model system, we examined the gene expression of 50 infant patients with t(4;11) pro-B ALL using qRT-PCR. Patient characteristics are summarized in Table 1. The age distribution of the cohort showed that no distinct age group was overrepresented (Figure 2A). Plotting of the  $\Delta C_T$  values and subsequent Pearson correlation testing for the *IRX1*, *EGR3*, and *ICOSLG* expressions revealed that the *ICOSLG* expression strongly correlated with the expression of *EGR3* and *IRX1* (Figures 2B–2F). The highest Pearson R value for the *EGR3-ICOSLG* relation (Figure 2E) was in line with the observation that *EGR3* binds directly to the *ICOSLG* promoter in the SEM cell model system (Figures 1C and 1D). According to our studies, *IRX1* does not directly bind the *EGR3* or *ICOSLG* promoter, in agreement with the relatively lower Pearson r values for these correlations (Figures 2D and 2F). In summary, gene expression analysis of patient samples verified that *IRX1* upregulates *EGR3*, which subsequently transactivates *ICOSLG* in infant t(4;11) ALL, as suggested by our SEM::EGR3 cell culture model system.

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<https://doi.org/10.1016/j.isci.2022.104613>



**Figure 1. EGR3 upregulates ICOSLG through direct promoter binding in an EGR3-overexpressing t(4;11) SEM cell model**

(A) qRT-PCR proved the transcriptional upregulation of EGR3 ( $\Delta\Delta C_T = 12.03 \pm 0.22$ ) and ICOSLG ( $\Delta\Delta C_T = 3.24 \pm 0.16$ ) 48h after the transgene induction of the SEM::EGR3 cell model.

(B) Western blotting of 20  $\mu$ g protein lysate confirmed the upregulation of EGR3 (FLAG-tagged) and ICOSLG on the protein level. SEM::mock showed a slight ICOSLG band owing to the basal expression of ICOSLG on SEM cells.

(C and D) Chromatin immunoprecipitation followed by qRT-PCR (C) and sequencing (D) displayed the direct binding of EGR3 towards the ICOSLG promoter area. ChIP-qRT-PCR was analyzed through percent input calculation followed by two-tailed unpaired t tests with Welch's correction to compare the percent input values of SEM:EGR3  $\alpha$ -FLAG with SEM:EGR3 IgG ( $p = 0.0082$ ), SEM::mock  $\alpha$ -FLAG with SEM::mock IgG ( $p = 0.4452$ ), and SEM::mock  $\alpha$ -FLAG with SEM:EGR3  $\alpha$ -FLAG ( $p = 0.0063$ ). The location of the primer pair used for ChIP-qRT-PCR is displayed as blue lines in the ICOSLG promoter area and indicated below with triangles accordingly.

### Patients with *HOXA9*<sup>lo</sup> show slightly inferior EFS than patients with *HOXA9*<sup>hi</sup> in a cohort of 43 infant patients with t(4;11) pro-B ALL

Patients with t(4;11) iALL were reported to cluster into a *HOXA*<sup>hi</sup> and a *HOXA*<sup>lo</sup> group (Trentin et al., 2009), with higher relapse rates in the *HOXA*<sup>lo</sup> group (Stam et al., 2010; Kang et al., 2012; Symeonidou and Ottersbach, 2021). We aimed to verify this finding and could demonstrate the clustering of patients with t(4;11)

**Table 1. Patient characteristics of the dx cohort (n = 50)**

Patient no.	Patient age [months]	Treatment protocol	sex	Time to event (last follow-up) [months]	HOXA9 $\Delta C_T$ Mean	IRX1 $\Delta C_T$ Mean	EGR3 $\Delta C_T$ Mean	ICOSLG $\Delta C_T$ Mean
1	5.0	Interfant-06	F	(40.1)	13.3592	6.1705	8.1850	7.6499
2	6.0	/	F	/	3.9204	7.7667	11.2758	8.0223
3	2.0	Interfant-06	F	(23.6)	12.9546	7.1581	9.3618	7.1255
4	6.0	Interfant-06	M	(17.1)	11.5783	7.9774	11.1637	7.8868
5	10.0	/	M	/	3.5974	8.3317	8.8214	6.0364
6	2.0	Interfant-06	M	(83.6)	4.9698	9.5555	11.5618	6.8416
7	3.0	Interfant-06	F	(5.6)	11.0093	5.1167	8.5196	5.6658
8	3.0	Interfant-06	F	(1.2)	4.9163	6.4184	8.2271	4.8055
9	3.0	Interfant-06	M	(2.5)	4.5976	9.0338	8.0114	9.8380
10	6.0	Interfant-06	F	9.5 (10.7)	5.0818	8.4812	12.3136	9.3984
11	0.1	/	F	/	8.6964	8.0250	11.2989	9.9469
12	1.0	/	M	/	12.3443	6.0576	11.6236	8.4245
13	12.0	/	F	/	2.2824	7.9734	9.1826	7.9894
14	4.0	/	F	/	11.4825	6.3166	11.4695	9.0590
15	2.0	Interfant-06	M	6.0 (6.0)	4.6973	5.9447	13.3440	8.7451
16	5.5	Interfant-06	M	9.5 (15.0)	6.2129	6.5399	9.5330	8.2498
17	1.5	Interfant-06	F	10.0 (13.0)	6.0703	4.9900	6.2132	4.0517
18	3.0	Interfant-06	F	6.0 (11.5)	2.0906	8.4791	9.4723	8.5023
19	2.0	Interfant-06	F	10.5 (19)	7.8537	3.1429	7.2056	5.2518
20	1.5	Interfant-06	M	26.5 (41.0)	4.4627	8.3685	11.2026	7.5564
21	6.5	Interfant-06	M	24.0 (32.8)	11.2673	5.3156	8.7817	8.6026
22	6.0	Interfant-06	F	19.5 (23.5)	11.5891	7.0636	10.0553	7.8697
23	12.0	Interfant-06	F	32.0 (60.6)	11.1226	4.2582	8.0600	7.3822
24	9.0	Interfant-06	M	(84.9)	8.7665	2.2034	7.0569	5.8906
25	2.0	Interfant-06	F	(118.8)	4.6017	7.1138	10.2925	7.1171
26	3.0	Interfant-06	M	8.0 (108.5)	13.0872	5.9083	8.7375	8.5604
27	5.5	Interfant-06	M	(9.0)	5.9626	8.1536	9.1105	8.6464
28	5.0	Interfant-06	M	17.0 (33.0)	5.9922	4.8934	5.0456	3.8081
29	2.0	Interfant-06	F	56.0 (58.6)	5.3297	8.4461	9.3271	8.9127
30	1.5	Interfant-06	F	(146.1)	5.8358	8.0518	9.8910	8.1978
31	8.3	Interfant-06	F	(109.3)	2.2195	6.2132	6.5075	6.2880
32	8.2	Interfant-06	M	(95.9)	1.7726	7.4956	7.2074	7.4143
33	10.7	Interfant-06	M	8.8 (8.8)	2.6953	3.0935	2.4138	4.9064
34	1.7	Interfant-06	F	(91.5)	3.8579	7.1200	6.2540	7.6448
35	11.7	Interfant-06	F	(73.2)	2.9311	7.9152	10.9273	7.2855
36	10.4	Interfant-06	F	(75.9)	5.7541	8.8581	10.4144	9.1282
37	2.4	Interfant-06	F	(73.8)	13.3331	6.1426	9.8438	9.0200
38	4.5	Interfant-06	M	16.1 (63.5)	4.6562	8.9152	11.4891	8.7484
39	1.1	Interfant-06	F	3.3 (15.7)	12.9071	6.3552	11.9464	9.3496
40	5.3	Interfant-06	M	(58.7)	5.4284	8.4380	10.3705	8.3493
41	6.9	Interfant-06	F	(63.2)	9.2472	8.5374	9.7053	6.0532
42	5.3	Interfant-06	M	(60.7)	3.4737	7.4371	9.5640	7.1599
43	0.6	Interfant-06	F	5.0 (11.9)	7.9122	6.2711	8.6364	8.0174

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Table 1. Continued

Patient no.	Patient age [months]	Treatment protocol	sex	Time to event (last follow-up) [months]	<i>HOXA9</i> $\Delta C_T$	<i>IRX1</i>	<i>EGR3</i>	<i>ICOSLG</i>
					Mean	$\Delta C_T$ Mean	$\Delta C_T$ Mean	$\Delta C_T$ Mean
44	1.5	Interfant-06	M	17.3 (18.3)	8.1333	8.0008	9.5745	7.1672
45	4.4	Interfant-06	M	6.6 (10.8)	4.2993	8.0016	10.2528	8.4346
46	9.1	Interfant-06	F	39.7 (45.2)	3.1663	7.7915	9.3985	8.2159
47	0.5	Interfant-06	F	8.7 (13.3)	6.2701	6.5276	9.9358	7.8396
48	6.1	AIEOP-BFM ALL 2017	F	6.6	12.6734	4.2609	9.4741	4.7534
49	6.1	AIEOP-BFM ALL 2017	F	8.3	4.5847	8.1496	10.5529	7.6922
50	10.6	/	M	/	5.0774	9.4705	5.1199	5.4875

All samples were taken from PB at the day of diagnosis. Gene expression values are indicated as the  $\Delta C_T$  mean of three technical replicates. F = female, M = male.

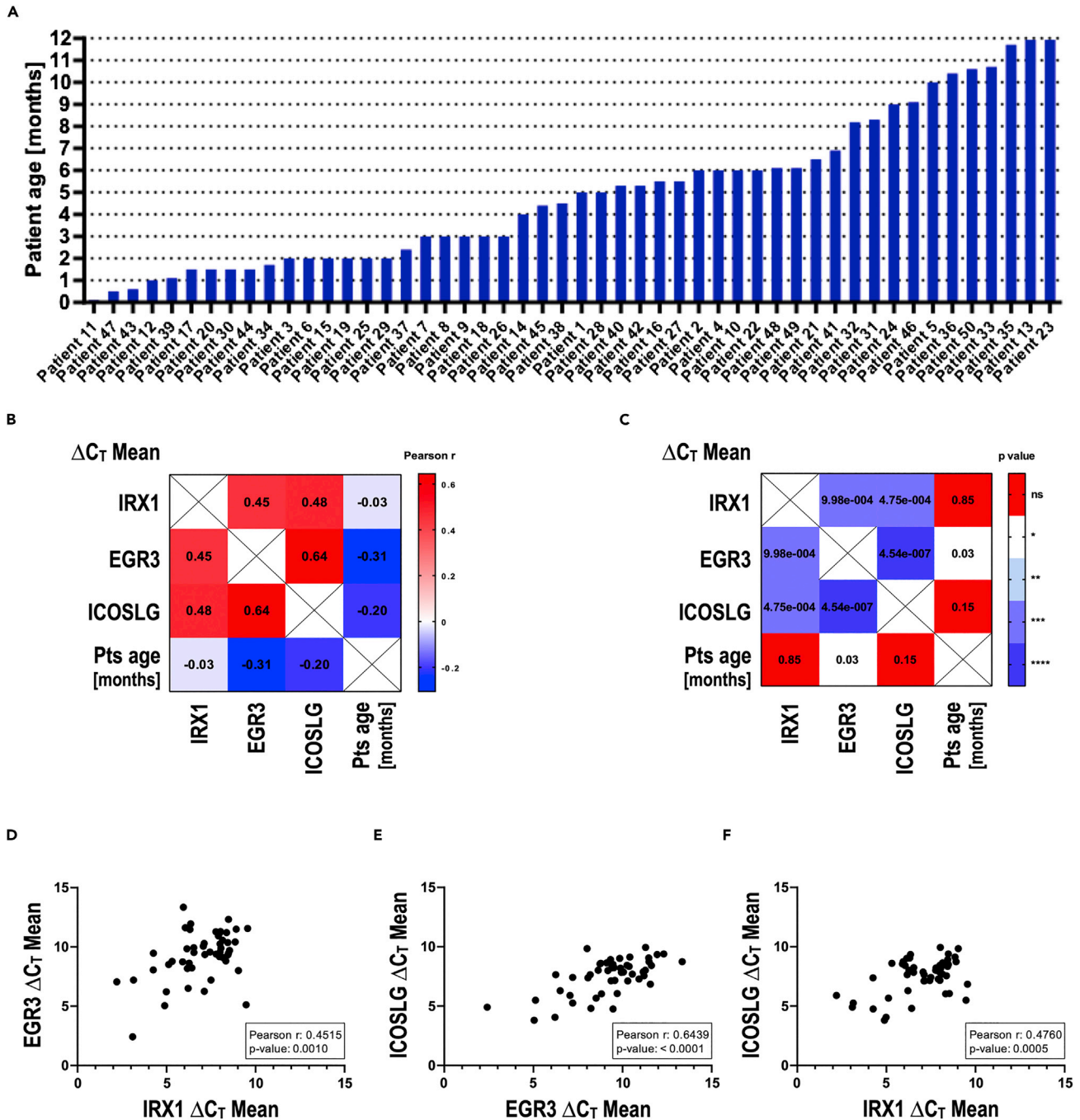
iALL into these two subgroups when analyzing the *HOXA9* transcriptional profiles (19 pts with *HOXA9*<sup>lo</sup>, 31 pts with *HOXA9*<sup>hi</sup>). Within the cohort of 50 patients, outcome data were available for 43 pts. We divided the patients into a group that achieved no event within 5 years from diagnosis (n = 12), and a group that achieved an event within the same time frame (n = 23). The assignment of these patients to the respective *HOXA9* transcription level revealed that patients with *HOXA9*<sup>lo</sup> were found in both groups (Figure 3A). Subsequent Kaplan-Meier analysis of all patients with the available outcome data (n = 43) demonstrated that the EFS disadvantage of patients with *HOXA9*<sup>lo</sup> was quite modest and did not reach statistical significance in a log rank test (Figure 3B). Consistent with the above-mentioned studies, patients with *HOXA9*<sup>lo</sup> displayed a higher median *IRX1* expression level than patients with *HOXA9*<sup>hi</sup> (p = 0.0012) (Figure 3C). However, this was not the case for the *EGR3* expression level (p = 0.5397) (Figure 3D) although we showed that the *IRX1* and *EGR3* expressions correlate strongly among the 50 patients (Figure 2D).

### High *ICOSLG* expression is associated with inferior EFS in a cohort of 43 infant patients with t(4;11) pro-B ALL

We then focused on *ICOSLG* expression, hypothesizing it could contribute to therapy resistance and relapse rates. For that purpose, we assigned the patients of the “no event group” (n = 12) and the “event group” (n = 23) to their respective *ICOSLG* transcription levels and found the clustering of patients into an *ICOSLG*<sup>lo</sup> and an *ICOSLG*<sup>hi</sup> group (Figure 4A). Interestingly, the “no event group” consisted only of patients with *ICOSLG*<sup>lo</sup>, whereas the “event group” included an *ICOSLG*<sup>lo</sup> and an *ICOSLG*<sup>hi</sup> cluster. The Kaplan-Meier analysis of all patients with available outcome data (n = 43) clearly demonstrated inferior EFS of the *ICOSLG*<sup>hi</sup> group (n = 7; 2-year-EFS = 0.0%) compared to the *ICOSLG*<sup>lo</sup> group (n = 36; 2-year-EFS = 58.8 ± 8.5%) (Figure 4B). A log rank test confirmed that both curves were significantly different (p value = 0.0107). However, this analysis could be biased owing to the difference in group sizes, but was not biased owing to patient age as the *ICOSLG*<sup>hi</sup> group is composed of patients younger and older than 6 months (Figure 4C). In agreement with our previous data, we showed that the *ICOSLG*<sup>hi</sup> group (n = 8) of all patients (n = 50) possessed significantly elevated *IRX1* and *EGR3* transcription levels. (Figures 4D and 4E).

### High *EGR3* and *ICOSLG* expressions are relapse associated

To examine whether the high expression of *ICOSLG* at diagnosis could be associated with relapse formation, we assessed the *HOXA9*, *IRX1*, *EGR3*, and *ICOSLG* gene expressions in a cohort of 18 relapsed pro-B iALL patient samples with different *KMT2A-r* partner genes taken at the time of relapse. These patients were an independent cohort, not related to the samples taken at diagnosis and patient characteristics are summarized in Table 2. We compared the gene expression data of the cohort at initial diagnosis (“dx”) with that one at relapse diagnosis (“rel”). To enable an appropriate comparison of both groups, we included only those patients of the dx cohort that achieved an event (n = 23). Thereby, we found that the rel cohort possessed a generally higher expression level of *EGR3* and *ICOSLG* when compared to the dx cohort (Figure 5A). Notably, the *ICOSLG* expression of the rel cohort was mainly at the same level as the *ICOSLG*<sup>hi</sup> group of the dx cohort (Figure 5B). The higher *EGR3* and *ICOSLG* expressions of the rel cohort compared to the dx cohort were statistically significant in unpaired t tests (p < 0.0001) (Figures 5B and 5C), whereas no clear difference became obvious regarding the *IRX1* and *HOXA9* expression comparison of both cohorts (Figures 5D and 5E). Subsequently, we performed a principal component

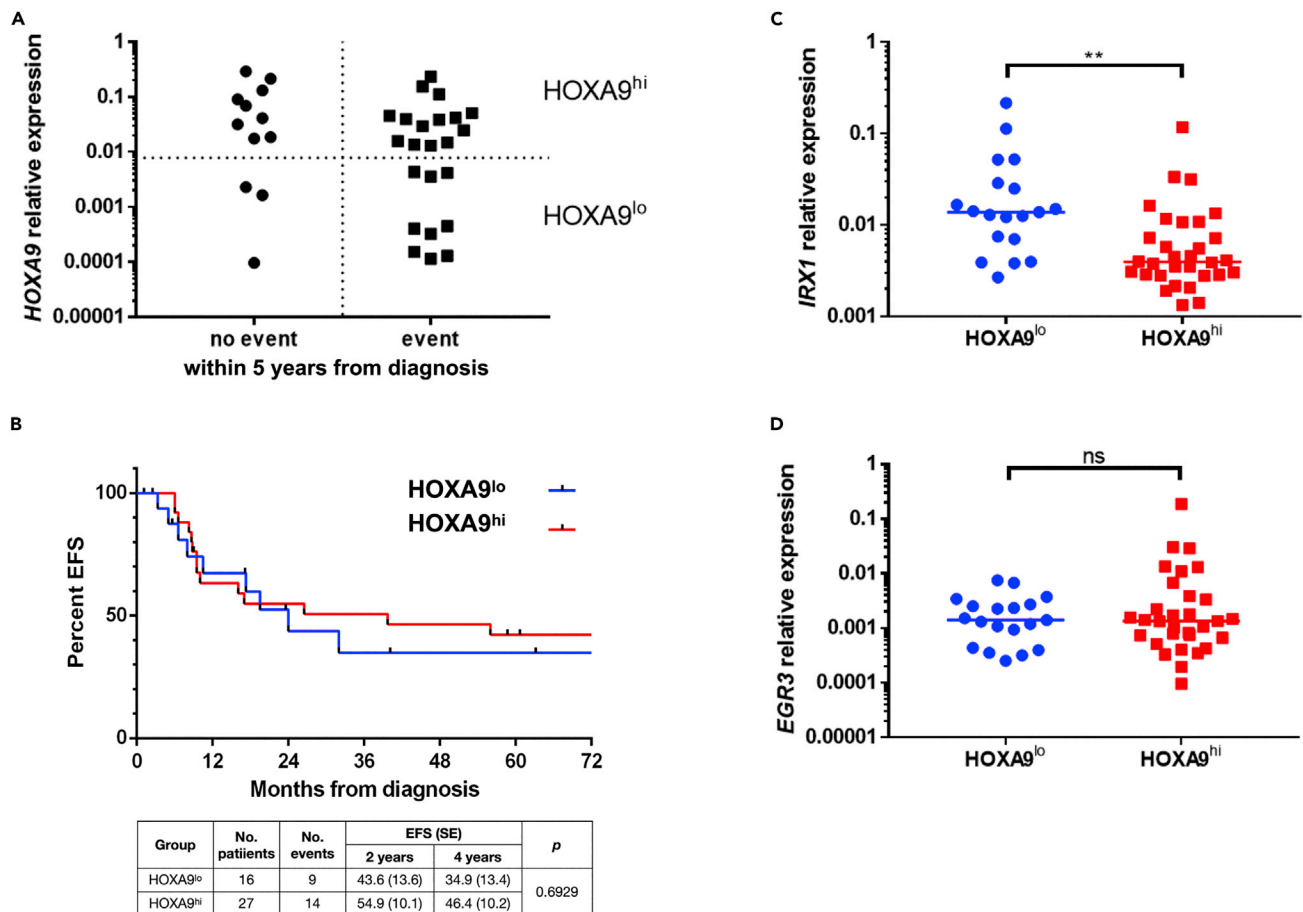


**Figure 2. *IRX1*, *EGR3*, and *ICOSLG* expressions correlate strongly in a cohort of 50 infant patients with *t*(4;11) pro-B ALL**

(A) Patient age distribution. Patients of all infant age groups are represented, and no age group is overrepresented.

(B–F) Pearson correlation matrix visualizes the Pearson *r* values (B) and corresponding *p* values (C) for all correlations tested. The strong positive correlations between the expressions of *EGR3*→*IRX1* (D), *ICOSLG*→*EGR3* (E) and *ICOSLG*→*IRX1* (F) are obvious mapping each patient sample in the correlation plots.

analysis (PCA) of the *IRX1*, *EGR3*, and *ICOSLG* expression data of the dx and rel cohorts using ClustVis (Met-salu and Vilo, 2015). Patients were assigned to the dx *ICOSLG*<sup>hi</sup> group, dx *ICOSLG*<sup>lo</sup> group, or the rel cohort to display the clustering of the patients representing the grade of similarity of their gene expression profiles (Figure 5F). The dx *ICOSLG*<sup>hi</sup> group (red) clustered together and overlapped completely with the rel cohort (green), whereas the dx *ICOSLG*<sup>lo</sup> group (blue) clustered besides with only a marginal overlap. The



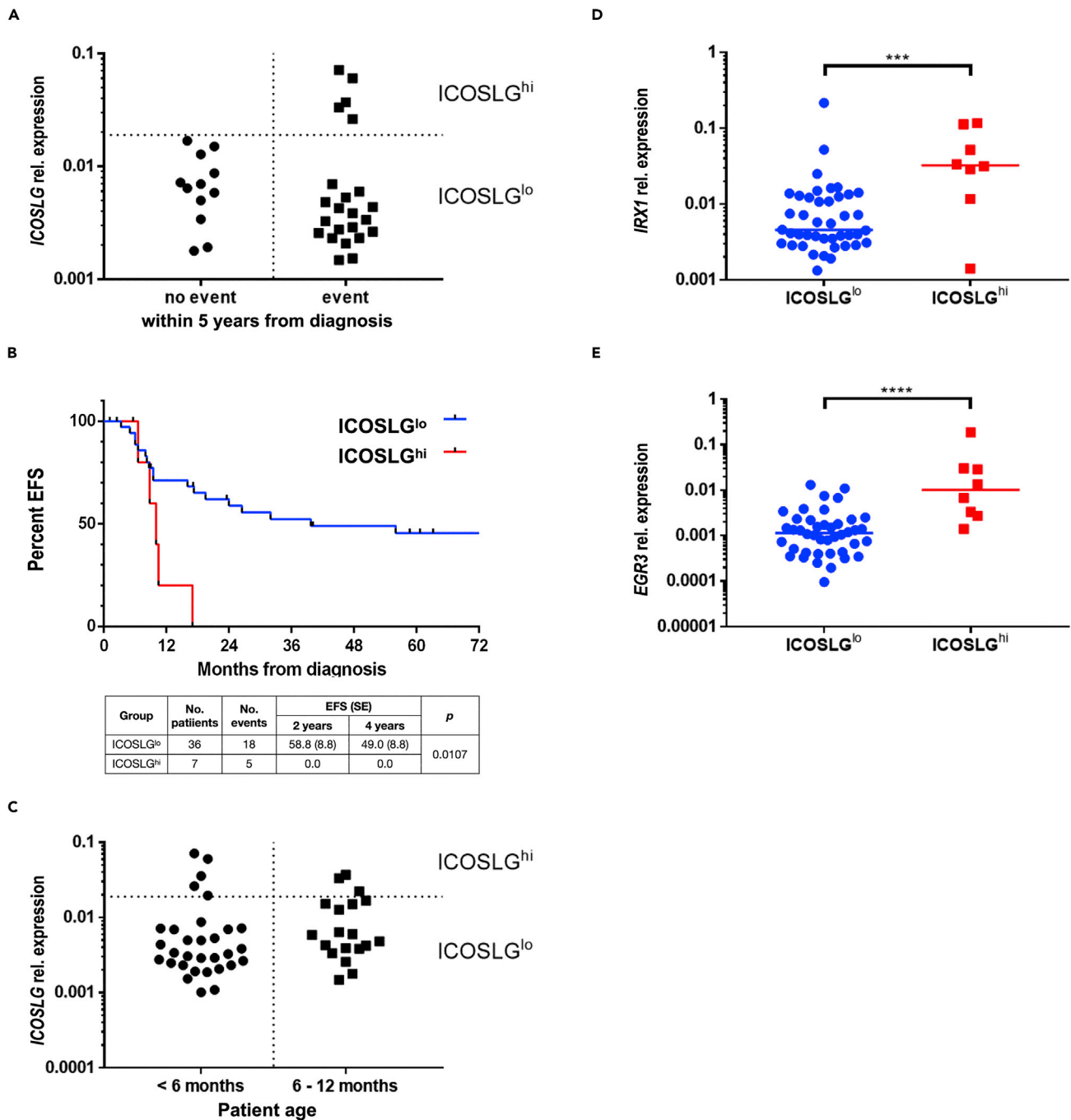
**Figure 3. Patients with  $HOXA9^{lo}$  show slightly inferior EFS than patients with  $HOXA9^{hi}$  in a cohort of 43 infant patients with t(4;11) pro-B ALL**  
 (A) Patients were divided into a group that displayed no event within 5 years from diagnosis ( $n = 12$ ), and a group with an event within the same time frame ( $n = 23$ ). Subsequently, patients were assigned to the respective  $HOXA9$  transcription level and divided into a  $HOXA9^{lo}$  and a  $HOXA9^{hi}$  subgroup.  
 (B) Kaplan-Meier curves of the patients with  $HOXA9^{lo}$  vs  $HOXA9^{hi}$  visualize the slightly inferior EFS of patients with  $HOXA9^{lo}$  which did not reach statistical significance in a log rank test ( $p = 0.6929$ ).  
 (C and D) Patients with  $HOXA9^{lo}$  have a higher median  $IRX1$  ( $p = 0.012$ ) (C) but not  $EGR3$  gene expression ( $p = 0.5397$ ) (D) than patients with  $HOXA9^{hi}$ . Significance was tested with two-tailed unpaired t tests.

coherent clustering of the dx  $ICOSLG^{hi}$  group with the rel cohort against dx  $ICOSLG^{lo}$  became also obvious regarding the PCA heatmap (Figure 5G). Thus, PCA revealed a high grade of similarity regarding  $IRX1$ ,  $EGR3$ , and  $ICOSLG$  expressions between the group of patients with high  $ICOSLG$  expression at diagnosis and the relapsed cohort. Taken together, these results strongly suggest that the  $IRX1$ - $EGR3$ - $ICOSLG$  axis is involved in relapse formation, and that high  $ICOSLG$  expression level at diagnosis can be described as a predictor for subsequent relapse.

### ICOSLG upregulation in SEM::EGR3 cells leads to the rapid development of $T_{regs}$ upon co-culture with primary T-cells

We then speculated that the causal relationship between  $ICOSLG$  expression and relapse in t(4;11) iALL could be the  $ICOSLG$ -mediated induction of  $T_{reg}$  development which was previously described in different physiological and pathological contexts (Martin-Orozco et al., 2010; Lee et al., 2017; Han et al., 2018; lwata et al., 2019). Accordingly, we examined this question in the t(4;11) pro-B cellular context using the established SEM::EGR3 cell line. For this purpose, primary human T-cells were co-cultured with either SEM::EGR3 or SEM::mock cells for 48h (50% SEM cells with 25%  $CD4^{+}$  and 25%  $CD8^{+}$  T-cells). The T-lymphocytes were isolated from peripheral blood mononuclear cells (PBMC) of healthy MHC-unmatched donors and were stimulated with coated  $\alpha$ -CD2/-CD3/-CD28 beads for 16h before co-culture. This stimulation mimicked T-cell receptor (TCR) activation, and thus, enabled a TCR-independent immune activation.





**Figure 4. High ICOSLG expression is associated with inferior EFS in a cohort of 43 infant patients with t(4;11) pro-B ALL**

(A) Patients of the “no event group” (n = 12) and the “event group” (n = 23) were assigned to their respective ICOSLG transcription levels and divided into an ICOSLG<sup>lo</sup> and an ICOSLG<sup>hi</sup> subgroup.

(B) Kaplan-Meier curves and log rank test of the patients with ICOSLG<sup>lo</sup> vs ICOSLG<sup>hi</sup> visualize the clearly inferior EFS of patients with ICOSLG<sup>hi</sup> (p = 0.0107).

(C) Patients separated by age were assigned to the respective ICOSLG transcription level illustrating that the ICOSLG<sup>hi</sup> group was composed of patients younger and older than 6 months.

(D and E) The ICOSLG<sup>hi</sup> group (n = 8) of all patients (n = 50) displayed significantly elevated IRX1 (p = 0.0006) (D) and EGR3 (p < 0.0001) (E) transcription levels in two-tailed unpaired t tests.

**Table 2. Patient characteristics of the rel cohort (n = 18)**

Patient no.	Age at primary diagnosis [months]	KMT2A-r partner	sex	HOXA9 $\Delta C_T$ Mean	IRX1 $\Delta C_T$ Mean	EGR3 $\Delta C_T$ Mean	ICOSLG $\Delta C_T$ Mean
REZ1	1.3	AFF1	F	2.016418	5.156435	2.832994	6.612465
REZ2	2.9	MLL1	M	5.517492	6.952302	4.17735	7.647544
REZ3	3.8	AFF1	F	10.23312	2.528003	4.301431	5.388622
REZ4	2.5	AFF1	F	7.767597	2.621157	6.657438	4.662416
REZ5	1.2	AFF1	F	3.034274	6.716536	1.18892	5.421249
REZ6	10.7	MLL1	M	7.759405	4.871906	5.178097	5.714579
REZ7	4.3	MLL1	M	-0.47106	4.344864	3.729839	2.899311
REZ8	3.6	MLL3	F	6.206228	6.106342	4.742364	5.892561
REZ9	9.1	AFF1	F	10.72401	4.103871	2.276168	4.287584
REZ10	0.7	AFF1	F	8.609946	2.540371	1.63076	4.492456
REZ11	4.4	AFF1	M	3.047612	5.159082	6.565317	4.748623
REZ12	4.2	AFF1	M	1.789008	6.10047	5.030182	5.842515
REZ13	2.0	AFF1	F	10.53802	1.708041	2.584866	5.500404
REZ14	3.1	AFF1	F	11.73027	9.246894	8.439681	4.853207
REZ15	7.6	AFF1	F	7.390529	6.94658	7.833634	2.701138
REZ16	6.7	AFF1	M	7.300869	9.940318	9.541564	4.663118
REZ17	5.3	AFF1	M	4.871099	8.645488	9.86666	3.962195
REZ18	7.2	AFF1	M	14.54123	9.570634	6.724747	3.354452

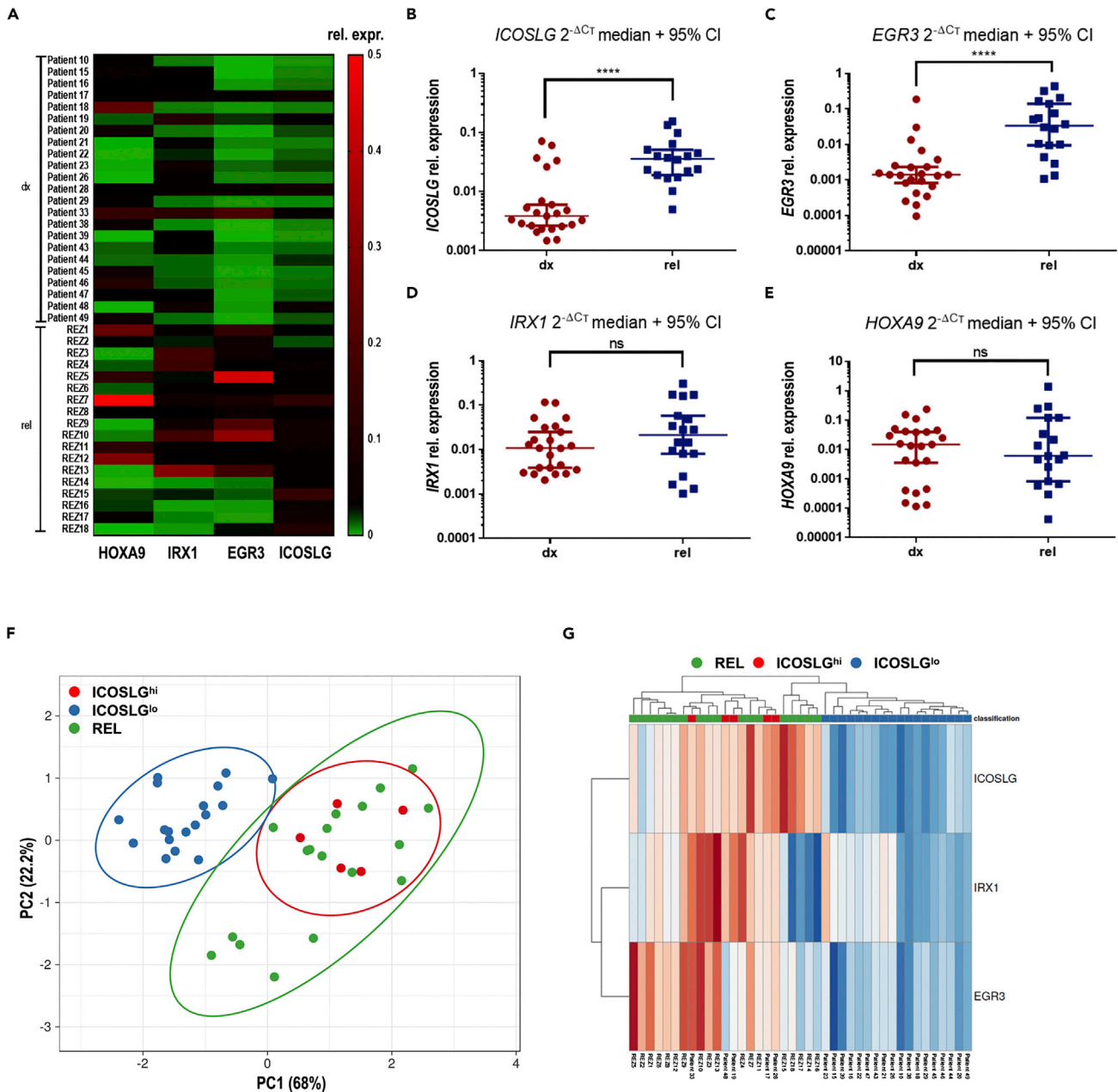
All samples were taken from PB on the day of relapse diagnosis. Gene expression values are indicated as the  $\Delta C_T$  mean of three technical replicates. F = female, M = male

After 48h of co-culture, the percentage of CD4<sup>+</sup>CD25<sup>+</sup>FOXP3<sup>+</sup> cells ( $T_{regs}$ ) among CD4<sup>+</sup> cells was quantified using flow cytometry. With six independent healthy donors (HD1-6), we observed an increase in the percentage of CD4<sup>+</sup>CD25<sup>+</sup>FOXP3<sup>+</sup>  $T_{regs}$  in the SEM::EGR3 co-culture between 7.96 and 23.94% (geometric mean:  $1.14 \pm 0.009991$ , p value = 0.0023) compared to the SEM::mock co-culture (Figure 6B). Noteworthy, SEM::mock cells expressed low amounts of ICOSLG but were not ICOSLG-negative (see Figure 1B).

To prove whether the ICOSLG/ICOS interaction was responsible for  $T_{reg}$  development, we performed the experiment under the addition of 20  $\mu$ g/mL mouse IgG1 isotype control or neutralizing monoclonal mouse  $\alpha$ -human ICOSLG antibody to the media (HD1-3).  $\alpha$ -ICOSLG but not IgG control antibody impaired the EGR3-mediated  $T_{reg}$  development during co-culture. (Figure 6C). These experimental observations suggest that EGR3-expressing t(4;11) ALL cells rapidly induced the development of  $T_{regs}$ . This induction was most likely caused by the EGR3-mediated transactivation of ICOSLG, as the effect was not observed when the immune checkpoint was selectively blocked by a specific checkpoint inhibitor antibody. We also assessed the cytokine levels in the supernatant of the co-culture using appropriate ELISA tests. These experiments revealed that ICOSLG blockade strongly decreased the IL-10 level of both the SEM::mock and SEM::EGR3 co-culture, whereas the IL-2 levels were not affected (Figures 6D and 6E). All this indicated that ICOSLG blockade impaired the development of  $T_{regs}$ . However, for HD1 and HD3 we observed also a strong increase of  $T_{regs}$  in the SEM::mock co-culture through ICOSLG antibody blockade which anyway did not affect the decreased IL-10 level through ICOSLG blockade.

### **$T_{regs}$ exhibit a higher ICOS surface expression than other CD4<sup>+</sup> cells after co-culture**

We hypothesized that a higher ICOS surface expression on  $T_{regs}$  compared to other T-cell subsets could be the reason for the preferential induction of  $T_{regs}$  through ICOSLG expression. This has already been reported for  $T_{regs}$  in the PB of healthy individuals (Duhon et al., 2012). Flow cytometry analyses revealed that CD25<sup>+</sup>FOXP3<sup>+</sup> cells displayed a higher ICOS surface expression than CD25<sup>+</sup>FOXP3<sup>-</sup> and CD25<sup>-</sup>FOXP3<sup>-</sup> cells within the CD4<sup>+</sup> subset (Figure 7). This was neither influenced by the co-cultured cell model (SEM::mock vs SEM::EGR3) nor by different antibody treatments (IgG vs  $\alpha$ -ICOSLG). As a result, the higher ICOS surface expression of  $T_{regs}$  in comparison to other T-cell subsets was likely the explanation for the induction of  $T_{regs}$  through ICOSLG expression.



**Figure 5. High *EGR3* and *ICOSLG* expressions are relapse associated**

(A) *HOXA9*, *IRX1*, *EGR3*, and *ICOSLG* expression levels are displayed in a heatmap for the dx cohort (patients with event; n = 23) and the rel cohort (n = 18). High  $2^{-\Delta CT}$  values indicating high expression are displayed in red.

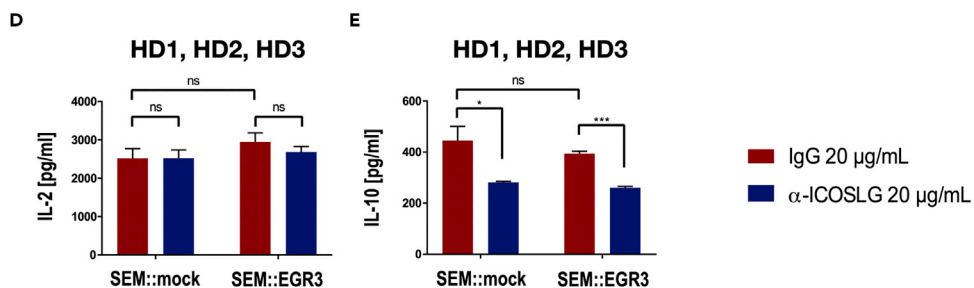
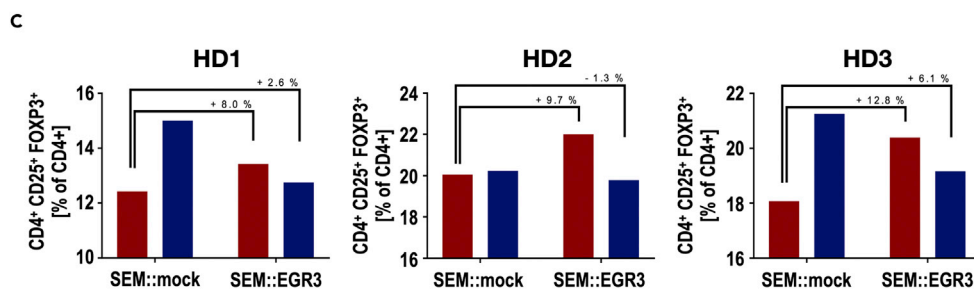
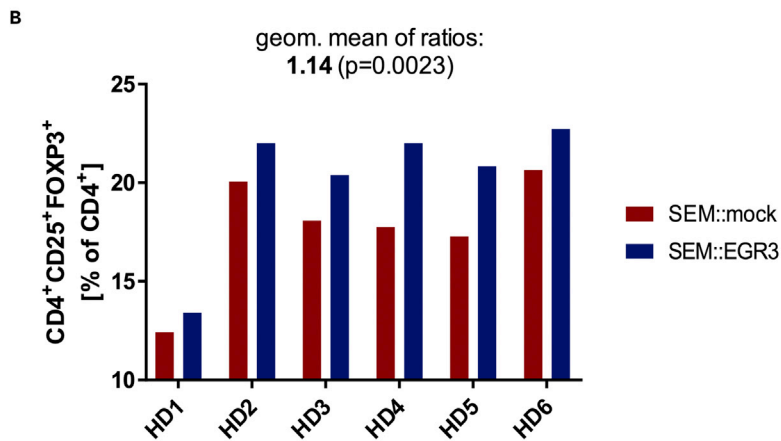
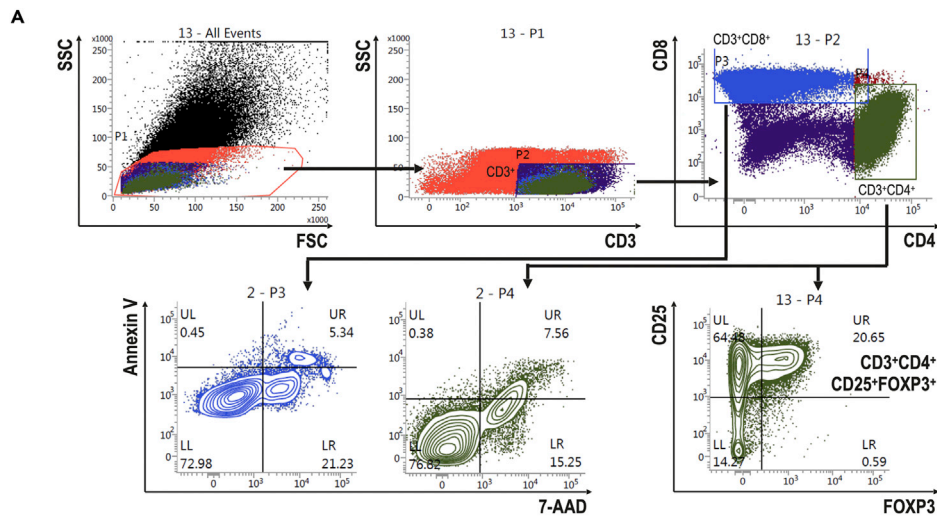
(B–D) Unpaired t tests indicated significantly higher *ICOSLG* ( $p < 0.0001$ ) (B) and *EGR3* ( $p < 0.0001$ ) (C) expression levels of the rel cohort compared to the dx cohort, whereas the *HOXA9* and *IRX1* expression levels did not significantly differ between both cohorts ( $p_{HOXA9} = 0.8374$ ;  $p_{IRX1} = 0.2737$ ) (E and D).

(F) Principal component analysis among *IRX1*, *EGR3*, and *ICOSLG* expression data with patients assigned to the rel cohort (green), or to the *ICOSLG*<sup>hi</sup> (red) or *ICOSLG*<sup>lo</sup> (blue) group of the dx cohort. PC1 represents 68%, PC2 22.2% of total variance, respectively.

(G) Heatmap of the PCA confirms the clustering of the *ICOSLG*<sup>hi</sup> group at diagnosis with the relapse cohort. High  $2^{-\Delta CT}$  values indicating high transcription levels are displayed in red.

## DISCUSSION

Infant t(4;11) ALL is one of the cancers with the lowest mutational burden with approximately 1.3 non-silent mutations in the dominant leukemic clone besides the chromosomal translocation (Andersson et al., 2015; Agraz-Doblas et al., 2019; Brown et al., 2019). Secondary mutations are present in some patients at



**Figure 6. ICOSLG upregulation in SEM::EGR3 cells leads to the rapid development of T<sub>regs</sub> upon co-culture with primary T-cells**

(A) Flow cytometry gating strategy of the co-cultured cells. Cells with low granularity (low SSC) were gated out of all cells. CD3<sup>+</sup> cells were selected to subsequently separate CD4<sup>+</sup> and CD8<sup>+</sup> cells. Viability after co-culture of these populations was assayed using Annexin V and 7-AAD staining to ensure that the majority of T-cells were alive throughout co-culture. CD25<sup>+</sup>FOXP3<sup>+</sup> cells (T<sub>regs</sub>) were quantified out of the CD4<sup>+</sup> population.

(B) Percentages of CD4<sup>+</sup>CD25<sup>+</sup>FOXP3<sup>+</sup> T<sub>regs</sub> among CD4<sup>+</sup> T-cells after co-culturing T-cells from six healthy donors (HD1-HD6) with SEM::mock or SEM::EGR3. SEM::EGR3 co-culture led to 7.96%-23.94% more T<sub>regs</sub> than SEM::mock co-culture (geometric mean: 1.14 ± 0.009991, p = 0.0023 in a two-tailed ratio paired t test).

(C) Replication of the co-culture experiment of HD1-HD3 under the addition of 20 µg/mL mouse IgG1 isotype control or neutralizing monoclonal mouse α-human ICOSLG antibody demonstrated that EGR3-mediated T<sub>regs</sub> induction could be impaired through ICOSLG antibody blockade.

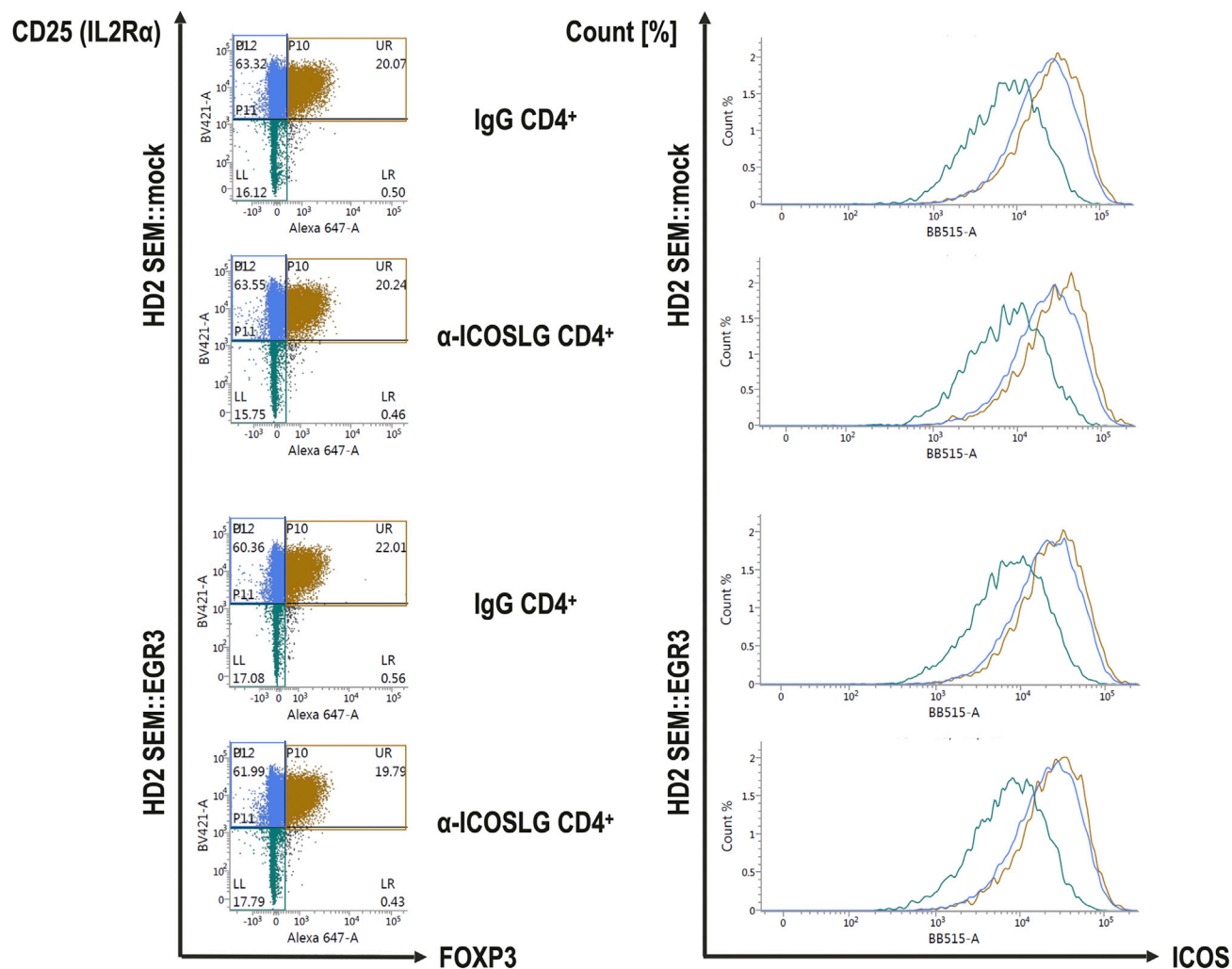
(D and E) ELISAs of the co-culture supernatants depicted that ICOSLG antibody blockade has no effect on the IL-2 level (D) but strongly decreased the IL-10 level (SEM::mock: p = 0.0424; SEM::EGR3: p = 0.0002) (E) of the media. Significance was tested with multiple t tests.

diagnosis but are the result of the dramatic cycling activity of the dominant clone and disappear once leukemia re-emerges as a relapse, indicating that the translocation is the only necessary leukemic driver (Agraz-Doblas et al., 2019). Based on our results, we propose that high ICOSLG expression at diagnosis and inferior EFS are causally linked, in that the ICOSLG-mediated induction of T<sub>reg</sub> development counteracts the immune recognition of leukemia cells. Through this mechanism, ALL cells could acquire a protective immune privilege in their bone marrow niche, enabling them to re-emerge as relapse despite ongoing chemotherapy. This mechanism is a possible explanation for relapse occurrence without the accumulation of additional mutations and clonal evolution.

This hypothesis implies that “fusion derived neoantigens” (FDNA) lead to T-cell priming and specific immune responses against the leukemic cells. Although cancers with a low mutational burden are generally regarded as barely immunogenic, *ETV6::RUNX1* fusion-specific CD8<sup>+</sup> T-cell responses have been reported in pediatric ALL (Zamora et al., 2019). Moreover, gene fusions are known to render T-cell responses in head and neck tumors with low mutational burden and minimal immune infiltration resulting in clinical response to immune checkpoint blockade (Yang et al., 2019). Therefore, it could be possible that also t(4;11)-derived neoantigens are immunogenic although this has not yet been demonstrated experimentally.

In pediatric ALL, the bone marrow (BM) is the predominant compartment where relapse originates (Tallen et al., 2010; Driessen et al., 2016). The BM provides the specialized niche in which HSC reside and give rise to all subsets of blood cells throughout life (Morrison and Scadden, 2014; Chen et al., 2016). In AML, leukemia stem cells (LSC) occupy the HSC niche and outcompete their healthy counterparts (Bonnet and Dick, 1997; Boyd et al., 2014; Medyouf, 2017; Kumar et al., 2018). In ALL, chemotherapy induces the formation of a protective niche by leukemia-initiating cells (LIC) that perturb hematopoiesis and lead to apoptosis of healthy HSC (Duan et al., 2014; Tang et al., 2018; Burt et al., 2019). In case of t(4;11) ALL, fetal liver-derived CD34<sup>+</sup>CD19<sup>-</sup> primitive B lineage-negative but lymphoid-restricted progenitor/stem cells have been identified as the most likely cellular origin of transformation (Hotfilder et al., 2005; Rice et al., 2021). However, it has been shown that not only CD34<sup>+</sup> but also CD34<sup>-</sup> ALL cells including blasts at different stages of differentiation possess stem cell capabilities and may act as LIC when serially transplanted in immunocompromised mice (le Viseur et al., 2008; Aoki et al., 2015). Accordingly, in t(4;11) ALL stemness seems not to be restricted to a certain cell type, suggesting that leukemia cells at different stages of differentiation can home to and reside in the BM niche and subsequently contribute to relapse. Therefore, therapeutic approaches that interfere with leukemic niche formation hold promise to prevent relapse development, particularly in infant t(4;11) ALL.

Importantly, the BM is not only a primary but also a secondary immune organ as it is part of the lymphocyte recirculation network (Osmond, 1994). CD4<sup>+</sup> and CD8<sup>+</sup> T-cells enter the BM from the vasculature and become primed through BM-residing dendritic cells (DC) resulting in primary immune responses (Feuerer et al., 2001, 2003). Besides CD4<sup>+</sup> and CD8<sup>+</sup> T-cells, T<sub>regs</sub> migrate to and accumulate in the BM through CXCL12/CXCR4 interactions (Zou et al., 2004). These T<sub>regs</sub> have been shown to colocalize with HSC and provide an immune privilege to the niche (Fujisaki et al., 2011). Additionally, niche-associated T<sub>regs</sub> are known to maintain HSC quiescence (Hirata et al., 2018, 2019). As healthy HSC depend on the colocalization with T<sub>regs</sub>, it is highly likely that LSC/LIC do as well, considering especially that the recruitment of immune



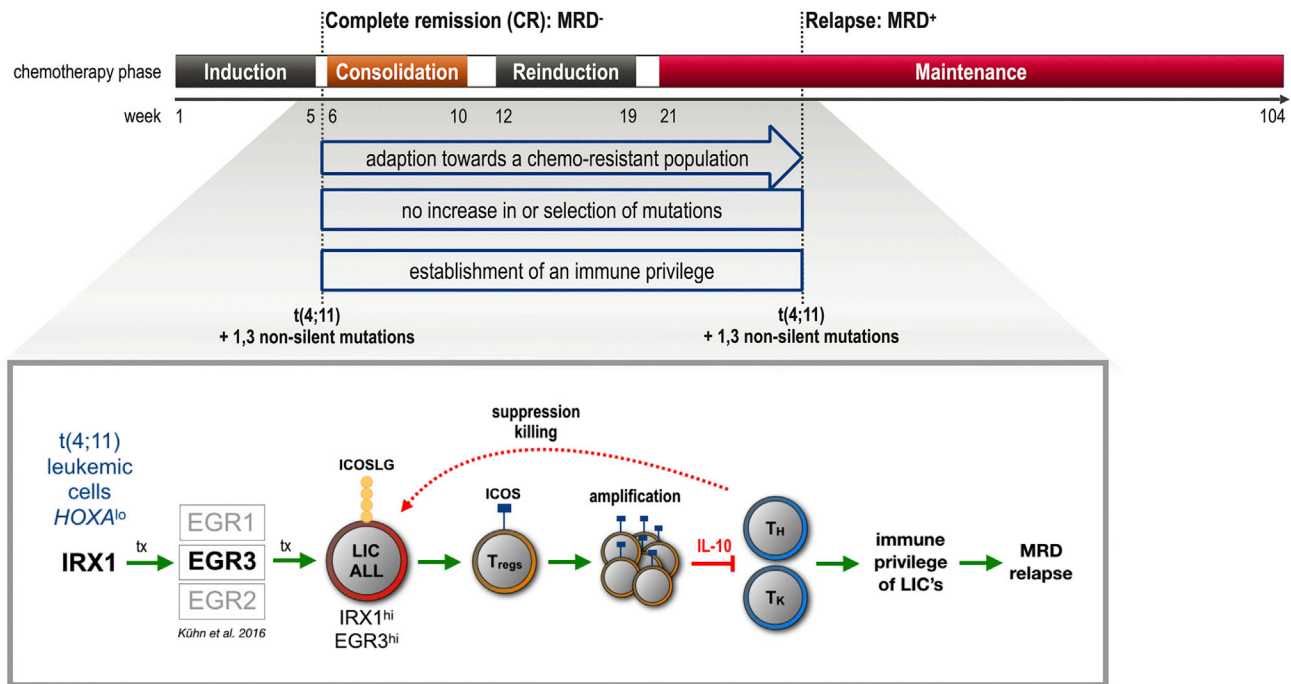
**Figure 7.  $T_{reg}$ s exhibit a higher ICOS surface expression than other  $CD4^+$  cells after co-culture**

Shown are the representative CD25- $\rightarrow$ FOXP3 dot plots of the  $CD3^+CD4^+$  subset of HD2.  $CD25^+FOXP3^+$ ,  $CD25^+FOXP3^-$  and  $CD25^-FOXP3^-$  cells were gated and their respective ICOS surface expression were displayed in a comparative plot.  $CD25^+FOXP3^+$  Tregs showed the highest ICOS expression, followed by  $CD25^+FOXP3^-$  cells.  $CD25^-FOXP3^-$  cells clearly display the lowest ICOS surface expression.

suppressive cells to avoid immune destruction is a common measure of several solid tumors and a hallmark of cancer (Hanahan and Weinberg, 2011).

Therefore, we speculate that ALL cell expression of ICOSLG facilitates the recruitment of  $T_{reg}$ s to the BM niche. The ICOSLG/ICOS interaction is known to affect  $CD4^+$  T-cells to produce IL-10 resulting in immunosuppression (Figure 8) (Hutloff et al., 1999; Akbari et al., 2002; Löhning et al., 2003). Furthermore, the development of  $T_{reg}$ s in the healthy bone marrow as well as in melanoma, glioblastoma, breast cancer and AML has been shown to strongly depend on the ICOSLG/ICOS interaction (Martin-Orozco et al., 2010; Faget et al., 2012; Lee et al., 2017; Han et al., 2018; lwata et al., 2019). Accordingly, the here described identification of EGR3-mediated ICOSLG-expressing ALL cells, enabling increased  $T_{reg}$  development is in line with previous observations.

In our study, co-culture of SEM::EGR3 with adult T-cells leads to an enrichment of  $ICOS^+FOXP3^+$   $T_{reg}$ s. In mice, the high surface expression of ICOS among  $T_{reg}$ s determines a higher proliferative capability, longer survival, and stronger suppression compared to  $ICOS^-$   $T_{reg}$ s (Chen et al., 2012). The higher suppressive activity of  $ICOS^+$   $T_{reg}$ s has also been observed in human studies (Vocanson et al., 2010). ICOS is expressed at low levels in all T-cell subsets but becomes upregulated upon ICOSLG binding (Hutloff et al., 1999;



**Figure 8. Proposed immune-evasion relapse mechanism in t(4;11) ALL**

Upper panel: Scheme of ALL chemotherapy. Although complete remission (CR) is frequently achieved after induction therapy, relapse occurs prevalently during consolidation, reinduction, or maintenance phase. Bottom panel: Infant t(4;11) ALL cells characterized by low *HOXA* transcription overexpress *IRX1*, which in turn causes the upregulation of several EGR transcription factors. EGR3 was shown to cause the transcriptional upregulation of *ICOSLG*. The interaction of *ICOSLG* on ALL cells with *ICOS* on T-cells initiates the development of regulatory T-cells ( $T_{regs}$ ). The accumulation of  $T_{regs}$  in the bone marrow niche could provide ALL cells with an immune privilege allowing them to acquire therapy resistance and to develop minimal residual disease (relapse).

Khayyamian et al., 2002). The  $T_{regs}$  which developed in our co-culture setting showed the highest *ICOS* surface expression among the  $CD4^+$  T-cells mirroring the results of previous studies (Duhon et al., 2012). Therefore, we assume that the higher and constitutional *ICOS* surface expression of  $T_{regs}$  compared to non- $T_{regs}$  could be a reason for their preferential proliferation during their co-culture with *ICOSLG*-expressing SEM cells. It is known that  $ICOS^+$   $T_{regs}$  suppress other T-cells mainly through *IL-10* secretion, whereas  $ICOS^-$   $T_{regs}$  secrete transforming growth factor- $\beta$  (*TGF- $\beta$* ) leading to weaker immunosuppression (Ito et al., 2008). In concordance, we show that *ICOSLG* antibody blockade leads to a strong decrease in the *IL-10* level of the co-culture media supernatant. Intracellularly, *ICOS* activation promotes *FOXP3* transcription, and in turn, *FOXP3*- and *NFAT*-dependent transactivation of *IL10* and *TGFB1* (Li and Xiong, 2020). Additionally,  $ICOS^+FOXP3^+$   $T_{regs}$  play an important role during fetal development and pregnancy as both cord blood and the newborn's thymus already contain  $ICOS^+FOXP3^+$   $T_{regs}$  (Takahata et al., 2004; Ito et al., 2008). Furthermore, the newborn's thymus contains more  $ICOS^+FOXP3^+$   $T_{regs}$  than adult peripheral blood and these  $T_{regs}$  express the thymus emigration marker *CD31* which implies that the  $ICOS^+$   $T_{reg}$  subset is derived directly from the thymus and does not develop out of  $ICOS^-FOXP3^+$   $T_{regs}$  (Takahata et al., 2004; Ito et al., 2008). The presence of  $ICOS^+$   $T_{regs}$  during fetal development is of special interest because it has been demonstrated by twin studies that the translocation t(4;11) arises *in utero*. (Gale et al., 1997; Wiemels et al., 1999; Greaves and Wiemels, 2003; Greaves et al., 2003). During pregnancy, the fetus is protected from the mother's semioforeign immune system through placental trophoblast cells which surround fetal tissues. *ICOSLG* is highly expressed throughout pregnancy by the extravillous trophoblast cells (EVT) (Petroff et al., 2005) and has been validated as a mediator of tolerance at the fetomaternal interface during pregnancy (Riella et al., 2013). Furthermore, EVTs mediate the invasion and differentiation of decidual  $T_{regs}$  which suppress T-cell responses during pregnancy through the secretion of *IL-10* (Aluvihare et al., 2004; Tilburgs et al., 2008; Du et al., 2014; Svensson-Arvelund et al., 2015; Salvany-Celades et al., 2019). Taken together, it is highly likely that the *ICOSLG/ICOS* interaction also provides an immune privilege to the fetomaternal interface. Recombining this knowledge with the results of our study leads to the suggestion that t(4;11) iALL depends on an immune privilege firstly to occur *in utero* and secondly to relapse in the BM. Accordingly, the *ICOSLG*-mediated  $T_{reg}$  development in the leukemic BM niche could reflect the immune

privilege during pregnancy, and thus could be a prerequisite for relapse development. Correspondingly, switching the immune-privileged leukemic BM niche toward immune-susceptibility holds promise to prevent relapse or to treat relapse-associated therapy resistance.

Here, we have shown that elevated ICOSLG expression at diagnosis is predictive for relapse development, and that targeting ICOSLG with a neutralizing mAb prevents the ICOSLG-mediated  $T_{reg}$  induction *in vitro*. Based on these findings, we propose the clinical investigation of ICOSLG targeting therapies such as the  $\alpha$ -ICOSLG fully human mAb Prezalumab. Prezalumab (AMG 557/MEDI 5872) has been tested clinically for the treatment of autoimmune diseases such as systemic lupus erythematosus (SLE) and active lupus arthritis in three phase I trials: NCT02391259, NCT00774943, and NCT01683695 (Sullivan et al., 2016; Cheng et al., 2018). In the first and second trial, 112 adult patients with mild SLE were enrolled and the safety and tolerability of single (1.8-210 mg s.c. or 18 mg i.v.) vs. multiple (6-210 mg s.c. every other week) dose administration of Prezalumab have been shown. Importantly, only non-neutralizing  $\alpha$ -Prezalumab antibodies were observed. The third trial demonstrated the long-term (23 weeks) safety and tolerability of Prezalumab among 10 patients. However, Prezalumab failed to demonstrate clinical improvement in a phase IIa study (NCT02334306) for the treatment of primary Sjögren's syndrome halting all further clinical investigation (Mariette et al., 2019; Pontarini et al., 2020).

Unfortunately, up to now, no studies were conducted to evaluate the safety of Prezalumab in infants or children with ALL. Regarding checkpoint inhibition in general, an interim analysis is available of the KEYNOTE-051 trial (NCT02332668), a phase I/II study evaluating the safety and efficacy of Pembrolizumab ( $\alpha$ -PD-L1) in children >6 months with advanced melanoma or PD-L1 positive solid tumor or lymphoma (Geoerger et al., 2020). Three patients aged 6 months-2 years, 22 aged 2-5 years, 25 aged 6-9 years, and 104 aged 10-17 years were enrolled. The study proved Pembrolizumab checkpoint inhibition safe in children as serious adverse events occurred only in 9% of patients (pyrexia, hypertension, and pleural effusion). Besides, another phase I/II trial proved the safety and tolerability of the PD-1 checkpoint inhibitor Nivolumab in children >1 year with lymphoma (Davis et al., 2020). These studies demonstrate that PD-1/PD-L1 immune checkpoint inhibition is safe in young children; however, the safety of ICOSLG-directed mAbs remains to be demonstrated in infants and children.

In summary, we provide a molecular rationale for the clinical evaluation of immune-based therapies targeting the ICOSLG/ICOS interaction for the treatment and prevention of relapse in infants with t(4;11) ALL. Further studies are needed to clarify whether t(4;11)-derived neoantigens lead to immune responses in infants and if that explains the potential demand of the disease for an immune-privileged site firstly to originate *in utero* and secondly to relapse in the BM. Even though these questions remain unanswered, we communicate our findings to enable a rapid translation into the clinical investigation.

## LIMITATIONS OF THE STUDY

The conclusion of this study has been obtained by *in vitro* studies, and thus, an appropriate mouse model is still missing. Also, the co-culture experiments with genetically modified t(4;11) cells, overexpressing the ICOSLG, were performed *in vitro* with a limited time window (48 h) and without MHC matching. In addition, we used a neutralizing commercial antibody against ICOSLG and not the clinically tested Prezalumab (AMGEN) because this antibody was not available to us.

## STAR★METHODS

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- Flow cytometry experiments
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**ACKNOWLEDGMENTS**

We thank all our collaborators for fruitful discussions and Birthe Fedders for sample composition and sending. This work has been funded in part by DFG grants MA 1876/12-1 and MA 1876/13-1.

**AUTHOR CONTRIBUTIONS**

MK and RM wrote the article; MK, HB, and RM planned and interpreted experiments; MK generated cell lines, performed qRT-PCR, Western blot, co-culture experiments, flow cytometry and ELISA; ALS performed plasmid cloning; CM, PL, and MK performed ChIP-Seq; CM and PL helped to write the article; AD and MK performed ChIP-qRT-PCR; JA, GC, CE, ACE, HC, MB, GiC, PDL, and MGV provided patient samples and outcome data, and helped to analyze clinical data; LD and HB provided blood samples and contributed to T-cell isolation; RM supervised the study and provided funding; all authors reviewed and approved the article.

**DECLARATION OF INTERESTS**

The authors declare that they have no competing interests.

Received: March 17, 2022

Revised: April 14, 2022

Accepted: June 1, 2022

Published: July 15, 2022

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## STAR★METHODS

### KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
<b>Antibodies</b>		
Goat Anti-Rabbit IgG, polyclonal antibody, Horseradish Peroxidase conjugated (1:10000; WB)	abcam	Abcam Cat# ab6721, RRID: AB_955447
DYKDDDDK Tag (D6W5B) Rabbit, monoclonal antibody, unconjugated (1/1,000 WB and ChIP)	Cell Signaling Technology	Cell Signaling Technology Cat# 14793, RRID: AB_2572291
Rabbit Anti-Actin, polyclonal antibody, unconjugated (1/1,000 WB)	Sigma-Aldrich	Sigma-Aldrich Cat# A2066, RRID: AB_476693
Rabbit Recombinant Anti-ICOS Ligand/ICOSL, monoclonal antibody, unconjugated (1/1,000 WB)	abcam	ab209262 RRID not found
Mouse Anti-Human CD3, monoclonal antibody, APC-H7 conjugated (flow cytometry)	BD	BD Biosciences Cat# 560176, RRID: AB_1645475
Mouse Anti-Human CD4, monoclonal antibody, PE-Cy7 conjugated (flow cytometry)	BD	BD Biosciences Cat# 557852, RRID: AB_396897
Mouse Anti-Human CD8, monoclonal antibody, BV510 conjugated (flow cytometry)	BD	BD Biosciences Cat# 563919, RRID: AB_2722546
Mouse Anti-Human CD25 (IL-2 Receptor $\alpha$ ), monoclonal antibody, BV421 conjugated (flow cytometry)	BD	BD Biosciences Cat# 564033, RRID: AB_2738555
Mouse Anti-Human FoxP3, monoclonal antibody, AF647 conjugated (flow cytometry)	BD	BD Biosciences Cat# 561184, RRID: AB_10584328
Mouse Anti-Human CD278 (ICOS), monoclonal antibody, BB515 conjugated (flow cytometry)	BD	BD Biosciences Cat# 564549, RRID: AB_2738840
Mouse Anti-Human B7-h2, monoclonal antibody, neutralizing, unconjugated (neutralization assays)	RnD Systems	R and D Systems Cat# MAB165, RRID: AB_2122734
Mouse IgG1 kappa isotype control, functional grade (neutralization assays)	Thermo Fisher	Thermo Fisher Scientific Cat# 16-4714-85, RRID: AB_470162
<b>Bacterial and virus strains</b>		
NEB® 5-alpha competent <i>E. coli</i> ; DH5 $\alpha$ ™ derivative	NEB	NEB Cat# C2987
<b>Biological samples</b>		
see Tables		
<b>Chemicals, peptides, and recombinant proteins</b>		
Q5® High-Fidelity DNA Polymerase	NEB	NEB Cat# M0491L
SuperScript™ II Reverse Transcriptase	Invitrogen	Invitrogen 18064071
Annexin V Pacific Blue	Invitrogen	Invitrogen A35122
SfiI restriction enzyme	NEB	NEB Cat# R0123L
T4 DNA ligase	NEB	NEB Cat# M0202S
Clarity Max Western ECL Substrate	Bio-Rad	Bio-Rad Cat# 1705062
Protease inhibitory cocktail tablets	Roche	Roche 4693116001
<b>Critical commercial assays</b>		
Amata™ Cell Line Nucleofector™ Kit R	Lonza	VCA-1001
Pierce™ BCA Protein Assay Kit (23225, thermo scientific)	Thermo Scientific	Thermo Scientific 23225
IL-10 (human) ELISA Kit	Enzo	Enzo ADI-900-036
IL-2 ELISA Kit (human)	Aviva	Aviva OKBB00179

(Continued on next page)

**Continued**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Accel-NGS® 2S plus DNA library kit	Swift Biosciences	Swift Biosciences 21024
Swift Unique Dual Indexing Primer Kit	Swift Biosciences	Swift Biosciences X9096
Accel-NGS 2S Truncated Adapters	Swift Biosciences	Swift Biosciences 28196

**Deposited data**

All Chip-Seq data have been deposited in the NCBI Geobus database	GSE205652
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**Experimental models: Cell lines**

SEM	DSMZ	ACC 546
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**Oligonucleotides**

GAPDH_fwd_short 5'-GGTCACCAGGGCTGC-3'	this paper	N/A
GAPDH_rev_short 5'-CGTTCTCAGCCTTGA-3'	this paper	N/A
ICOSLG_fwd 5'-TGACATCGGAGAGAGACAAG-3'	this paper	N/A
ICOSLG_rev 5'-ACGCTTTTGAAGGGCCTCA-3'	this paper	N/A
EGR3.SfiI_fwd 5'-GGCCTCTGAGGCCACCATGACCGCAAACCTCGCC-3'	this paper	N/A
EGR3cFLAG.SfiI_rev 5'-GGCCTGACAGGCCTTACTTATCGTCATCGTCCTTGTAGTC-3'	this paper	N/A
EGR3.SF1_fwd 5'-GGCCTAGCAAGACACCGC-3'	this paper	N/A
HOXA9_fwd_short 5'-CCACGCTTGACACTCACACT-3'	this paper	N/A
HOXA9_rev_short 5'-AGTTGGCTGCTGGGTTATTG-3'	this paper	N/A
IRX1_fwd_short 5'-GGATCTCAGCCTTCTTCG-3'	this paper	N/A
IRX1_rev_short 5'-GTGGAGACTGCGTGAGG-3'	this paper	N/A
EGR3_fwd_short 5'-CAGCGACTCGGTAGTCCATT-3'	this paper	N/A
EGR3_rev_short 5'-TAGGTCACGGTCTTGTGCC-3'	this paper	N/A
ICOSLG_fwd_short 5'-AGTCCGGAGACAGAGCTCAC-3'	this paper	N/A
ICOSLG_rev_short 5'-CAGTCTGGGAGTCCATGCTC-3'	this paper	N/A
ICOSLG_fwd_ChIP1 5'-ACCGGGACCCATGGCA-3'	this paper	N/A
ICOSLG_rev_ChIP1 5'-CTCCCTCCTTCCAGCGTTC-3'	this paper	N/A
ICOSLG_fwd_ChIP2 5'-GTGAGCCGGGAAGGA-3'	this paper	N/A
ICOSLG_rev_ChIP2 5'-ATGGGTCCCGGTGCCA-3'	this paper	N/A
ICOSLG_fwd_ChIP3 5'-GCAGAGCCGAACCTTCCG-3'	this paper	N/A
ICOSLG_rev_ChIP3 5'-TGGAGGCAGCCGTGTC-3'	this paper	N/A
ICOSLG_fwd_ChIP4 5'-CCTCAGAGCCAGTGTTAAGC-3'	this paper	N/A
ICOSLG_rev_ChIP4 5'-CGCATGCATGAATGGAAGGA-3'	this paper	N/A
ICOSLG_fwd_ChIP5 5'-GCCCTGCCAGCTCTT-3'	this paper	N/A
ICOSLG_rev_ChIP5 5'-CAGGCCACGCTGAGAGAC-3'	this paper	N/A
ICOSLG_fwd_ChIP6 5'-GCTCTCCAAGGCCAGCTG-3'	this paper	N/A
ICOSLG_rev_ChIP6 5'-AAGCTGCTACTCCTGGTTCA-3'	this paper	N/A

**Recombinant DNA**

pSBtet_Puro sleeping beauty vector backbone	<a href="#">Kowarz et al. (2015)</a>	AddGene #60507
EGR3 open reading frame	this paper	HEK293T cDNA
Sleeping beauty transposase plasmid pcGLobin-SB100 <sub>xco</sub>	<a href="#">Moudgil et al. (2020)</a>	AddGene #154887

**Software and algorithms**

GraphPad Prism 7.01	GraphPad Software Inc.	GraphPad Prism 7.01
ClustVis ( <a href="#">Metsalu and Vilo, 2015</a> )	open source	N/A

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**Continued**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
TrimGalore	open source	N/A
Bowtie2	open source	N/A
SAMtools	open source	N/A
MACS3	open source	N/A
R package "ChIPseeker"	open source	N/A
R package "encodeChIPqc"	open source	N/A
python tool "deepTools"	open source	N/A
Integrative Genomics Viewer (IGV)	open source	N/A
<b>Other</b>		
Low Fluorescence PVDF Western Membrane	Abcam	Abcam Cat# ab133411
Precision Plus Protein™ All Blue Standards; marker used for Western Blots with fluorescence detection	Bio-Rad	Bio-Rad Cat# 1610373EDU
Color Protein Standard Broad Range	NEB	NEB P7719S

**RESOURCE AVAILABILITY**

**Lead contact**

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Rolf Marschalek ([Rolf.marschalek@em.uni-frankfurt.de](mailto:Rolf.marschalek@em.uni-frankfurt.de)).

**Materials availability**

All unique/stable reagents generated in this study are available from the Lead Contact with a completed Materials Transfer Agreement.

**Data and code availability**

All experimental data or any additional information required to reanalyze the data reported in this paper are available from the [Lead contact](#). This paper does not report original code. Any additional information required to reanalyze the data reported in this paper is available from the [Lead contact](#) upon request.

**EXPERIMENTAL MODEL AND SUBJECT DETAILS**

**Animals**

Not applicable.

**Cell culture**

SEM cells (ACC 546) and isolated primary T-cells were cultivated under sterile conditions. SEM cells were maintained in RPMI 1640 (RPMI-HA, Capricorn Scientific) supplemented with 10% FBS (FBS-11A, Capricorn Scientific), 2 mM L-glutamine (STA-B, Capricorn Scientific), 100 U/ml penicillin and 100 µg/mL streptomycin (PS-B, Capricorn Scientific), preheated to 37°C prior to use. Primary T-cells were maintained in DMEM high glucose (DMEM-HPA, Capricorn Scientific) supplemented with 10% human plasma (derived from each donor), 2 mM L-glutamine (STA-B, Capricorn Scientific), 100 U/ml penicillin and 100 µg/ml streptomycin (PS-B, Capricorn Scientific), preheated to 37°C prior to use. Cells were kept at 37°C in 5% CO<sub>2</sub> and a relative humidity of 95%. SEM cells were passaged twice a week with complete media exchange maintaining a density of ~1 x 10<sup>6</sup> cells/mL. For this, cell suspension was centrifuged at 200 x g for 5 min and subsequently cell pellet was resuspended in new prewarmed media. T-cells were immediately activated following isolation as described below and underwent co-culture experiments, thus no passaging was necessary.

**Co-culture study**

1x10<sup>6</sup> SEM::EGR3 or SEM::mock were co-cultured for 48h with 5x10<sup>5</sup> CD4<sup>+</sup> T-cells and 5x10<sup>5</sup> CD8<sup>+</sup> T-cells in 1 mL of DMEM-HPA (Capricorn Scientific) supplemented with 10% human plasma (derived from each donor), 2 mM L-glutamine (STA-B, Capricorn Scientific), 100 U/ml penicillin and 100 µg/mL streptomycin (PS-B, Capricorn Scientific), preheated to 37°C prior to use. T-lymphocytes were isolated from the

peripheral blood mononuclear cells (PBMC) of healthy donors using magnetic bead separation as described below. Prior to co-culture, isolated T-cells were stimulated with the T-cell Activation/Expansion Kit human (130-091-441, Miltenyi Biotec) for 24h according to the manufacturer's recommendations. SEM::EGR3 and SEM::mock were induced with 1  $\mu\text{g}/\text{mL}$  Doxycycline 24h prior to co-culture and 20  $\mu\text{g}/\text{mL}$  neutralizing monoclonal mouse  $\alpha$ -human ICOSLG antibody (RnD Systems®) or mouse IgG1 isotype control (Invitrogen™) were added 2h prior to co-culture. After 48h of co-culture cells were collected and stained with FACS antibodies and supernatant immediately frozen in liquid nitrogen and stored for ELISA experiments.

## METHOD DETAILS

### Plasmid cloning

EGR3 open reading frame (ORF) was amplified from HEK293T cDNA with the primers EGR3.Sfil\_fwd and EGR3cFLAG.Sfil\_rev using Q5® High-Fidelity DNA Polymerase (NEB, M0491L). EGR3.Sfil\_fwd adds the 5'-Sfil restriction site GGCCTCTGAGGCC upstream of ATG and EGR3cFLAG.Sfil\_rev adds a FLAG (DYKDDDDK) tag to the ORF and the 3'-Sfil restriction site GGCCTGTCAGGCC downstream of TAA. Equipped with the Sfil restriction sites, the ORF was ligated into Sfil-digested pSBtet\_Puro backbone (R0123L, NEB) using T4 DNA ligase (M0202S, NEB). The yielded plasmid pSBtet\_EGR3cFLAG\_Puro was sequence validated.

### Establishment of the stable cell lines SEM::mock and SEM::EGR3

Electroporation of SEM cells was performed using the Nucleofector™ 2b device (AAB-1001, Lonza) together with the Amaxa™ Cell Line Nucleofector™ Kit R (VCA-1001) and the nucleofection program T-016 according to the manufacturer's recommendations. 0.1  $\mu\text{g}$  sleeping beauty transposase plasmid pcGLobin-SB100<sub>XCO</sub> together with 1.9  $\mu\text{g}$  sleeping beauty plasmid (pSBtet\_ohneLuc\_Puro = SEM::mock; pSBtet\_EGR3cFLAG\_Puro = SEM::EGR3; pSBtet\_Luc\_GP = green fluorescence positive control) were electroporated. pSBtet\_ohneLuc\_Puro and pSBtet\_EGR3cFLAG\_Puro do not have a fluorescence marker in their backbone to avoid interference in multicolor flow cytometry. To monitor transfection and selection efficiency with a fluorescence microscope, pSBtet\_Luc\_GP containing a GFP marker in the backbone was treated equally and served as a reporter. Cells were selected six times (once to twice per week) with Puromycin 2  $\mu\text{g}/\text{mL}$  for 24-48h until all reporter cells displayed green fluorescence. Proof of transgene expression was performed using PCR and subsequent gel electrophoresis, qRT-PCR, and western blot 48h after induction of transgene expression with Doxycycline 1  $\mu\text{g}/\text{mL}$ .

### Human T-cell isolation

PBMC were isolated from the blood of healthy adult donors using Ficoll-Paque density gradient centrifugation. Therefore, 15 mL blood were diluted with 20 mL PBS (PBS-1A, Capricorn Scientific) supplemented with 0.5% BSA and overlaid with Lymphocyte Separation Media (MS00ZZ1002, Biowest) followed by 30 min centrifugation at 400 g without break leading to separation of plasma and PBMC from remaining blood cells. Plasma was stored for T-cell culture media preparation. PBMC layer was moved to another tube and resuspended in 50 mL PBS supplemented with 0.5% BSA followed by centrifugation at 400 g for 5 min with break. Pellet was resuspended in 10 mL RBC lysis buffer for human liquid (J62990, Alfa Aesar) and 10 min incubated at room temperature to lyse remaining erythrocytes. Afterwards, cell suspension was washed with 30 mL PBS supplemented with 0.5% BSA and isolated PBMC were counted. CD4<sup>+</sup> and CD8<sup>+</sup> cells were magnetically separated using CD4 Micro-Beads, human (130-045-101, Miltenyi Biotec) or CD8 MicroBeads, human (130-045-201, Miltenyi Biotec), MS columns (130-042-201, Miltenyi Biotec) and OctoMACS™ separator (130-042-109, Miltenyi Biotec) according to the manufacturer's protocols. Success of T-cell isolation was checked with flow cytometry prior to activation.

### Gene expression analysis using qRT-PCR

RNA was extracted from  $\sim 5 \times 10^7$  SEM::mock and SEM::EGR3 48h after transgene induction with 1  $\mu\text{g}/\text{mL}$  Doxycycline. cDNA was synthesized out of 2  $\mu\text{g}$  RNA using random hexamer N6 primers and SuperScript™ II Reverse Transcriptase (18064071, Invitrogen). PCR primers used in the study are summarized in [STAR Methods](#). qRT-PCR was performed with the StepOnePlus system.  $\Delta C_T$  mean values were calculated out of triplicates using GAPDH expression as a reference,  $\Delta\Delta C_T$  values are related to the respective expression of SEM::mock.



### Protein expression analysis using Western blot experiments

Protein was extracted from  $\sim 5 \times 10^7$  SEM::mock and SEM::EGR3 48h after transgene induction with 1  $\mu\text{g}/\text{mL}$  Doxycycline. Cells were lysed in 1 mL protein extraction buffer (150 mM NaCl, 10 mM Tris-HCl, 1 mM EDTA, 1 mM EGTA, 0.2 mM sodium ortho-vanadate, 0.2 mM PMSF, 1% Triton-X-100, 0.5% NP-40, freshly added protease inhibitory cocktail (4693116001, Roche) under constant agitation for 30 min at 4°C. Centrifugation at 400 g, 4°C enabled separation of the supernatant (= protein lysate) from cell debris. Protein lysate was quantified using Pierce™ BCA Protein Assay Kit (23225, Thermo Scientific) according to the manufacturer's recommendations. 20  $\mu\text{g}$  protein lysate were electrophoretically separated using Mini-PROTEAN TGX Stain-Free Gels 4-15% (4568085, Bio-Rad) and Color Protein Standard Broad Range (P7719S, NEB). Proteins were blotted onto Low Fluorescence Western Membranes PVDF (ab133411, Abcam) in Towbin buffer (25 mM Tris, 192 mM Glycin, 15% methanol, 0.01% SDS, pH = 8.3) at 110 V for 70 min. Membranes were blocked using 5% BSA in TBS-T for 1h at room temperature and primary antibody incubation (1/1,000) performed o/n at 4°C. Membranes were washed with TBS-T and incubated with HRP-conjugated secondary antibody (1/10,000) for 1h at room temperature. Chemiluminescence detection was performed using Clarity™ Western ECL Substrate (170-5061, Bio-Rad) and ChemiDoc™ XRS+ system (1708265, Bio-Rad). Antibodies used for western blot are listed in [STAR Methods](#).

### Flow cytometry experiments

Cells were blocked and stained with FACS antibodies (BD Biosciences) according to the manufacturer's recommendations. Half of the cell suspension was stained with  $\alpha$ -CD3, -CD4, -CD8, Annexin V and 7-AAD to assess the viability of the T-cells after co-culture. The remaining cells were stained with  $\alpha$ -CD3, -CD4, -CD8, -CD25, -ICOS, fixed, permeabilized and stained with  $\alpha$ -FOXP3. Cells were analysed with a BD FACSVerser™. The gating strategy is explained in [Figure 6A](#). All antibodies used for this strategy are described in [STAR Methods](#).

### ELISA experiments

The IL-2 and IL-10 concentrations of the co-culture supernatants were assessed using IL-10 (human) ELISA Kit (ADI-900-036, Enzo) and IL-2 ELISA Kit (human) (OKBB00179, Aviva) according to the manufacturer's recommendations.

### Chromatin immunoprecipitation qRT-PCR and sequencing

$2 \times 10^7$  SEM::EGR3 and SEM::mock were induced with 1  $\mu\text{g}/\text{mL}$  Doxycycline for 48h followed by ChIP procedure as described by [Kühn et al., \(2016\)](#). The transcription factor binding site prediction tool JASPAR ([Sandelin et al., 2004](#)) was used to detect potential EGR3 binding sites 1 kb up- and downstream of the transcription start site of *ICOSLG*. Six primer pairs were used to cover the nine highest scoring potential binding sites in qRT-PCR. Primers used for ChIP-qRT-PCR are summarized in [STAR Methods](#). Antibodies used for ChIP are described in [STAR Methods](#). The ChIP-Seq library was prepared using the Accel-NGS® 2S plus DNA library kit (21024, Swift Biosciences) together with Swift Unique Dual Indexing Primer Kit (X9096, Swift Biosciences) and Accel-NGS 2S Truncated Adapters (28196, Swift Biosciences) according to the manufacturer's recommendations with indexing by PCR (10 cycles). NGS was performed using the MiSeq system (SY-410-1003, Illumina) and the MiSeq Reagent Kit v2 (MS-102-2003, Illumina). The ChIP-Seq analysis was performed in several steps. As a first step, in order to remove adapters contamination, raw Illumina paired-end reads were processed with TrimGalore. In a second step, we mapped paired-end sequence reads against the human reference genome GRCh38 with Bowtie2. SAMtools was then used to convert the resulting sam files to bam files and to generate the index files. Once the chromatin immunoprecipitate (IP) and control (input) libraries have been aligned, we proceeded with the peak calling in order to determine the regions in which the IP signal is significantly higher than the input signal (=background). We performed the peak calling with the tool MACS3 with the option BAMPE active because of paired end reads. The result of the peak calling (narrowPeak file) was then used with the R package "ChIPseeker" to annotate the peaks to neighboring genes and genomics regions. We then performed a number of routine quality controls such as PCR bottleneck coefficient (PBC), fraction of reads in peaks (FriP) using the R package "encodeChIPqc" and fingerprint plot using "plotFingerprint" from the python tool "deepTools". In the final step we used the functions "bamCoverage" and "bamCompare" from "deepTools" to generate signal files in the bigwig format. We generated either separate bigwig files for IP and control with "bamCoverage" or a single bigwig file in which we subtracted the background noise from the IP with "bamCompare". Finally, the signal files, along with narrowPeak files and bam files, can be visualized using the Integrative Genomics

Viewer (IGV). Data have been deposited in the NCBI Geobus database with the accession number GSE205652.

### Patient gene expression study

Informed consent was obtained for all patients by the respective study centers that provided patient RNA. All patients displayed a pro-B phenotype and were diagnosed between 0 and 12 months of age (infants). RNA was extracted from blood at day of initial diagnosis (dx cohort, n = 50) or relapse diagnosis (rel cohort, n = 18) by the respective study centers. The dx cohort is solely composed of t(4;11) iALL cases, the rel cohort is composed of different translocations as indicated in Table 2. cDNA was synthesized out of 1 µg RNA as described above. *HOXA9*, *IRX1*, *EGR3* and *ICOSLG* gene expression was measured as triplicates using qRT-PCR (StepOnePlus system) and  $\Delta C_T$  mean values were calculated using *GAPDH* expression as a reference. Relative expressions were calculated as ratio (reference/target) =  $2^{CT(GAPDH) - CT(target)} = 2^{-\Delta C_T}$ . Pearson correlation was calculated for the gene expression datasets *IRX1*- > *EGR3*, *IRX1*- > *ICOSLG* and *EGR3*- > *ICOSLG* using GraphPad Prism software. Clinical outcome of the patients was provided by the study centers.

### QUANTIFICATION AND STATISTICAL ANALYSIS

Statistical tests were performed using GraphPad Prism 7.01 and settings are indicated in figure legends. Principal component analysis (PCA) was performed using ClustVis (Metsalu and Vilo, 2015) with patient sample  $\Delta C_T$  mean value assigned to the dx *ICOSLG*<sup>hi</sup>, dx *ICOSLG*<sup>lo</sup> or rel cohort, respectively. PCA was performed as singular value decomposition (SVD) and unit variance scaling for rows was applied. The PCA heatmap was generated with *correlation* as clustering distance for rows, *average* as clustering method for rows and columns, and *Manhattan* as clustering distance for columns.

Kaplan-Meier plotting and log rank test were performed using GraphPad Prism 7.01. EFS was defined as the time from diagnosis to first failure (induction failure, relapse, death, or second malignant neoplasm). Time was censored at last follow-up if no events had been observed. Curves were estimated with Kaplan-Meier, with SEs calculated according to Greenwood, and compared using the log rank test.