T-BET and EOMES Accelerate and Enhance Functional Differentiation of Human Natural Killer Cells

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T-bet and Eomes are transcription factors that are known to be important in maturation and function of murine natural killer (NK) cells. Reduced T-BET and EOMES expression results in dysfunctional NK cells and failure to control tumor growth. In contrast to mice, the current knowledge on the role of T-BET and EOMES in human NK cells is rudimentary. Here, we ectopically expressed either T-BET or EOMES in human hematopoietic progenitor cells. Combined transcriptome, chromatin accessibility and protein expression analyses revealed that T-BET or EOMES epigenetically represses hematopoietic stem cell quiescence and non-NK lineage differentiation genes, while activating an NK cell-specific transcriptome and thereby drastically accelerating NK cell differentiation. In this model, the effects of T-BET and EOMES are largely overlapping, yet EOMES shows a superior role in early NK cell maturation and induces faster NK receptor and enhanced CD16 expression. T-BET particularly controls transcription of terminal maturation markers and epigenetically controls strong induction of KIR expression. Finally, NK cells generated upon T-BET or EOMES overexpression display improved functionality, including increased IFN-γ production and killing, and especially EOMES overexpression NK cells have enhanced antibody-dependent cellular cytotoxicity. Our findings reveal novel insights on the regulatory role of T-BET and EOMES in human NK cell maturation and function, which is essential to further understand human NK cell biology and to optimize adoptive NK cell therapies.

Keywords: human NK cells, transcription factors, T-BET, EOMES, CD16 expression, antibody-dependent cellular cytotoxicity, NK cell biology, NK cell therapy
INTRODUCTION

Natural killer (NK) cells can provide anti-tumor effects by the production of proinflammatory cytokines and by direct target lysis (1–3). NK cells, like other lymphocytes, originate from CD34+ hematopoietic stem cells (HSC) in the bone marrow that differentiate through a common lymphoid progenitor stage. In secondary lymphoid tissues, human NK cell development is pursued, whereby the cells sequentially develop into stage 1 (CD34+CD45RA+CD117+CD94+) pro-NK cells, followed by stage 2 or pre-NK cells (CD34+CD45RA+CD117+CD94+). Stage 1 and stage 2 cells are multipotent as they also have T-cell and dendritic cell developmental potential. Stage 3 cells (CD34+CD117+CD94+CD16+) are committed NK cell precursors since they can no longer develop into T-cells or dendritic cells. Stage 4 (CD34+CD56brightCD94+CD16+) and stage 5 (CD34−CD56dimCD94+CD16−) are mature NK cells (4–7). Differentiation and functional maturation of NK cells is a complex molecular process tightly regulated by transcription factors. Many essential factors have been identified in the transcriptional control of murine NK cell differentiation, thanks to the generation of transcription factor-deficient mice (8). In contrast to mice, the current knowledge on the role of transcription factors in human NK cell differentiation is extremely limited.

Because of their intrinsic anti-tumor effects, NK cells are promising agents for cancer immunotherapy. Although different approaches using NK cells in cancer therapy have already been used, there are still major limitations. It has been shown that NK cell function after allogeneic hematopoietic stem cell transplantation (HSCT) in leukemia patients is impaired (9). The expression level of the transcription factors T-box expressed in hematopoietic cells and is known as a master (14, 15). T-bet-decient (T-bet−/−) mice show reduced numbers of NK cells in liver, spleen and peripheral blood. In contrast, the number of NK cells in the bone marrow is slightly higher in T-bet−/− mice and these NK cells have an immature phenotype (16, 17). Eomesbox/boxVav-Cre+ mice show a more substantial decrease of NK cell numbers in spleen and peripheral blood, but not in liver. Eomes-deficient NK cells also show an immature phenotype. Mice lacking both T-bet and Eomes completely fail to develop NK cells in all organs (17). These knockout mouse models show that both T-bet and Eomes are indispensable for NK cell development and terminal NK cell maturation. In parallel to mice, human peripheral blood and spleen NK cells are characterized by a T-BET and EOMES gradient. As NK cells progress from stage 3 to stage 5, they downregulate EOMES and upregulate T-BET, highlighting their reciprocal relationship. This illustrates that the T-BET and EOMES gradient follows the pattern of NK cell maturation, whereby EOMESlow T-BETHigh cells are considered as terminal mature NK cells (18, 19).

As low T-BET and EOMES expression levels in NK cells from tumor patients negatively impacts the anti-tumor effects, we here studied the effects of either T-BET or EOMES overexpression in cord blood-derived hematopoietic progenitor cells (HPC) on NK cell differentiation and function. Transcriptome and chromatin accessibility profiling demonstrate that T-BET or EOMES overexpression in human HPC epigenetically regulates activation of an NK cell transcriptome, leading to drastic acceleration of NK cell differentiation. Furthermore, the early arising NK cells have a mature phenotype and are enriched in CD16 expression. In-depth analysis of mature NK cells generated from T-BET- or EOMES-overexpressing HPC shows that terminal maturation of these NK cells is regulated at the epigenome level, wherein T-BET plays a predominant role. Additionally, NK cells generated from T-BET- or EOMES-overexpressing HPC are functional, whereby EOMES overexpression NK cells display enhanced antibody-dependent cellular cytotoxicity (ADCC). Altogether, these findings give new insights in the regulatory role of T-BET and EOMES in human NK cell differentiation and function that can be used to optimize adoptive NK cell therapies.

MATERIALS AND METHODS

Retroviral Overexpression Constructs

Human T-BET and EOMES cDNA (Source BioScience, Nottingham, UK; T-BET cDNA: IRATp970D0558D; EOMES cDNA: IRAKp961A1269Q) were ligated separately into the LZRS-IRES-eGFP retroviral vector (20). The empty LZRS-IRES-eGFP vector was used as control. Retrovirus was generated as previously described (21).

Isolation of CD34+ HPC From Umbilical Cord Blood

Umbilical cord blood (UCB) was obtained from the Cord Blood Bank, Ghent University Hospital, Ghent, Belgium. Mononuclear cells were obtained by Lymphoprep density gradient centrifugation and CD34+ HPC were subsequently enriched using Magnetic Activated Cell Sorting (Direct CD34+ HSC MicroBead Kit, Miltenyi Biotech, Leiden, The Netherlands) according to the manufacturer’s guidelines. Enriched CD34+ HPC were stored in liquid nitrogen until usage.

In Vitro NK Cell Differentiation Co-Culture

In vitro NK cell differentiation co-cultures on the murine embryonic liver cell line EL08.1D2 were performed as described
by Cichoki and Miller (22) with minor adaptations. EL08.1D2 feeder cells were maintained in 50% Myelocult M5300 medium (Stem Cell Technologies, Grenoble, France), 35% α-MEM, 15% FCS and 10 μM β-mercaptoethanol (Sigma-Aldrich, Saint Louis, MO), supplemented with 100 U/mL penicillin, 100 μg/mL streptomycin, 2 mM glutamine (all from Life Technologies, Carlsbad, CA) on 0.1% gelatin-coated plates at 33°C and 5% CO2. EL08.1D2 cells were inactivated by adding 10 μg/mL mitomycin C (Sigma-Aldrich) to the culture medium for 2-3 hours. Thereafter, cells were thoroughly rinsed before harvesting using trypsin-EDTA. Cells were plated at a density of 5x10⁴ cells per well on a 0.1% gelatin-coated tissue culture-treated 24-well plate at least 24 h before adding HPC or before transfer of differentiated NK cells on day 14 and day 21 of culture.

**Retroviral Transduction of HPC and NK Cell Differentiation**

Isolated UCB-derived CD34⁺ HPC were cultured in complete IMDM containing 10% FCS and supplemented with thrombopoietin (TPO) (20 ng/mL), stem cell factor (SCF) (100 ng/mL) (all from Peprotech, London, UK.) and FMS-like tyrosine kinase 3 ligand (FLT3-L) (100 ng/mL, R&D Systems, Minneapolis, MN) for 48 hours. Subsequently, these cells were harvested, transferred to RetroNectin (Takara Bio, Saint-Germain-en-Laye, France)-coated plates and viral supernatant was added. Additional cytokines were added to keep the concentrations constant after virus addition. The plates were centrifuged at 950 g and 32°C during 90 min. After 48 hours, eGFP⁺ “lineage-” CD34⁺ HPC were sorted by BD FACSAria™ Fusion cell sorter (BD Biosciences, San Jose, CA). Sorted HPC were co-cultured with mitomycin-inactivated EL08.1D2 cells for 3, 7, 14 or 21 days in Dulbecco’s modified Eagle medium plus Ham’s F-12 medium (2:1 ratio), supplemented with 100 U/mL penicillin, 100 μg/mL streptomycin, 2 mM glutamine, 10 mM sodium pyruvate (all from Life Technologies), 20% of heat-inactivated human AB serum (BioWest, Nuaille, France), 24 μM β-mercaptoethanol, 20 μg/mL ascorbic acid and 50 ng/mL sodium selenite (all from Sigma-Aldrich). The following cytokines were added: IL-3 (5 ng/mL, first week only, R&D systems), IL-7 (20 ng/mL), IL-15 (10 ng/mL) (all from Miltenyi Biotec), SCF (20 ng/mL), and Flt3-L (10 ng/mL). Alternatively, to test the necessity of IL-15 in NK cell differentiation upon T-BET and EOMES transduction, IL-15 was not included in the cytokine mix. Culture medium was refreshed on day 7 by addition of the same volume of fresh medium with cytokines. At day 14 the non-adherent cells were harvested and transferred to new mitomycin C-treated EL08.1D2 feeder cells.

**Library Preparation, RNA Sequencing, and qPCR Confirmation**

For transcriptome analysis, day 0 eGFP⁺ “lineage-” CD34⁺ HPC, day 21 stage 4 (eGFP⁺CD45⁺CD56⁻CD94⁺CD16⁻) and stage 5 NK cells (eGFP⁺CD45⁺CD56⁻CD94⁺CD16⁻) were sorted and RNA was isolated (RNeasy micro kit, Qiagen, Hilden, Germany). The concentration and quality of the extracted RNA was checked using the ‘Quant-it ribogreen RNA assay’ (Life Technologies) and the RNA 6000 nano chip (Agilent Technologies, Santa Clara, CA, U.S.A), respectively. Subsequently, 71 ng and 59 ng of RNA from HPC samples and NK cell samples, respectively, was used to perform an Illumina sequencing library preparation using the QuantSeq 3’ mRNA-Seq Library Prep Kit (Lexogen, Vienna, Austria) according to manufacturer’s protocol. Libraries were quantified by qPCR, according to Illumina’s protocol ‘Sequence Library qPCR Quantification protocol guide’, version February 2011. A High sensitivity DNA chip (Agilent Technologies) was used to check the library’s size distribution and quality. Sequencing was performed on a high throughput Illumina NextSeq 500 flow cell generating 75 bp single-end reads. Per sample, on average 5.3x10⁶ ± 1.7x10⁵ and 7.32x10⁶ ± 1.78x10⁵ reads of HPC and NK cells samples, respectively, were generated, whereby quality control of these reads was performed with FastQC (23). Reads were then mapped against the Homo sapiens GRCh38.90 reference genome using STAR version 2.4.2 and gencode version 25 as guide gtf. Gene quantification was performed on the fly by STAR (24). To explore if the samples from different treatment groups clustered together and to detect outlier samples, a Principal Component Analysis (PCA) on vst transformed counts was performed using the R statistical computing software (25). No outliers among the samples were detected. Differential gene expression analysis was performed using DESeq2 (26) with Wald test, whereby T-BET- or EOMES-overexpressing HPC were compared to control-transduced HPC, or stage 4 and stage 5 NK cells generated upon T-BET or EOMES overexpression were compared to control NK cells. Genes with an FDR<0.1 were considered significantly differential.

GSEA was performed using the GSEA software tool v4.1.0 of the Broad Institute (27, 28). The ‘GSEAPreranked’ module was run using standard parameters and 1000 permutations.

To perform qPCR analysis, RNA was converted to cDNA using Superscript RT III (Life Technologies, Carlsbad, CA). PCR reactions were executed using the LightCycler 480 SYBR Green I Master mix (Roche, Diegem, Belgium) on a LightCycler 480 real-time PCR system (Roche) using the primers indicated in Table 1.

**Library Preparation and Fast-ATAC Sequencing**

Chromatin landscape analysis was performed on the same cell populations as used for RNA sequencing. Library preparation was performed as previously described (29, 30). Briefly, 5x10⁶ HPC or day 21 stage 4 and stage 5 NK cells were sorted using a BD FACSAria™ Fusion cell sorter, washed 2 times with cold PBS and pelleted by centrifugation at 500g for 5 minutes at 4°C. After removal of the supernatant, 50 μl of transposase mixture (25 μl 2x TD buffer, 2.5 μl of TDE1 and 22 μl nuclease free water) (all from Illumina, San Diego, CA) containing 0.5 μl of 1% digitonin (Sigma-Aldrich) was added to the cells. Transposition reactions were incubated at 37°C for 30 min with agitation at 250 rpm. Transposed DNA was then purified using the QiAgen MinElute PCR purification kit (Qiagen, Hilden, Germany) according to the manufacturer’s instructions. Thereafter, dsDNA was qualified and quantified with Qubit™ dsDNA HS Assay Kit and Qubit™ Fluometer (both from Thermo
Motif enrichment analysis was executed with Homer findMotifsGenome.pl with default settings, whereby a bed file of all detected ATAC peaks was used as background. Homer findMotifsGenome.pl with -find option was used to recover individual motif locations (31).

Flow Cytometry
Cells were examined using flow cytometry (LSRII flow cytometer, BD Biosciences) and data were analyzed with FACSDiva Version 6.1.2 (BD Biosciences) and/or FlowJo_V10 (Ashland, OR, U.S.A) software. Utilized antibodies are listed in Table 3. Before staining, cells were blocked with anti-mouse FcγRII/III (unconjugated, clone 2.4G2, kindly provided by Dr J. Unkeless, Mount Sinai School of Medicine, New York, NY) and anti-human FcR blocking reagent (Milltenyi Biotec). Propidium iodide or the LIVE/DEAD Fixable Aqua Dead Cell Stain Kit (Life Technologies) was used to discriminate live and dead cells. FoxP3/Transcription Factor Staining buffer set (eBioscience, Thermo Fisher Scientific, Waltham, MA) was used to perform transcription factor staining.

To determine absolute cell numbers of HPC or NK cell subpopulations, cultured cells were harvested and counted with trypan blue in a Bürker counting chamber to determine the total number of viable cells. These counted viable cell numbers were normalized to a starting cell number of 1000 cells on day 0 for each condition. FACS analysis was used to determine the percentages of the different subpopulations and the normalized viable cell numbers were then multiplied by the corresponding percentage of each subpopulation.

Cytospins

eGFP⁺CD45⁺CD56⁺CD94⁺ NK cells were sorted from day 3 and 7 T-BET or EOMES overexpression cultures and from day 19 control cultures. Cytospins were made using Shandon Cytospin™ 4 (Thermo Fisher Scientific, Carlsbad, CA) according to the manufacturer’s guidelines. For reliable microscopic discrimination of individual cells, less than 500 cells per µl in a volume of 250 µl was added to the sample chamber of the cytospins. Dry cytoslides were subsequently Wright-Giemsa stained for microscopic evaluation.
**TABLE 3 | Utilized antibodies for flow cytometry.**

| Antibody | Alternative name | Conjugated fluorochrome | Clone | Supplier |
|----------|------------------|-------------------------|-------|----------|
| CD117    | CD117            | Phycoerythrin-Cyanin7 (PECy7) | 104D2 | eBioscience™; Thermofisher Scientific |
| CD11a    | LFA-1            | Phycoerythrin-Cyanin7 (PECy7) | H111  | Biolegend |
| CD11a    | LFA-1            | Phycoerythrin (PE)        | H111  | BD Pharmingen |
| CD11a    | LFA-1            | Allophycocyanin (APC)    | H111  | Biolegend |
| CD14     | CD14             | Allophycocyanin (APC)    | REA599 | Miltenyi Biotec |
| CD158b/h | KIR2DL1/DS1      | Phycoerythrin (PE)       | REA1010 | Miltenyi Biotec |
| CD158b/h | KIR2DL2/DS2/DS3 | Phycoerythrin (PE)       | GL183  | Beckman Coulter |
| CD158b/h | KIR2DL1/DS1      | Phycoerythrin (PE)       | Z27.3,7 | Beckman Coulter |
| CD158b/h | KIR2DL4          | Phycoerythrin (PE)       | FES172 | Beckman Coulter |
| CD159a   | NKG2A            | Allophycocyanin (APC)    | REA110 | Miltenyi Biotec |
| CD16     | CD16             | Allophycocyanin (APC)    | B73.1  | Biolegend |
| CD16     | CD16             | Phycoerythrin (PE)       | B73.1  | Biolegend |
| CD19     | CD19             | Allophycocyanin (APC)    | SJ25C1 | eBioscience™; Thermofisher Scientific |
| CD226    | DNAM-1           | Alexa Flur 647           | DX11   | BD Pharmingen |
| CD3      | CD3              | Allophycocyanin (APC)    | SK7    | Biolegend |
| CD314    | NKG2D            | Allophycocyanin (APC)    | 1D111  | Biolegend |
| CD335    | NKp46            | Phycoerythrin-Cyanin7 (PECy7) | 9E2    | Biolegend |
| CD335    | NKp44            | Allophycocyanin (APC)    | 44.189 | eBioscience™; Thermofisher Scientific |
| CD337    | NKp30            | Phycoerythrin (PE)       | P80-15 | Biolegend |
| CD34     | CD34             | VioBlue (VB)             | AC136  | Miltenyi Biotec |
| CD34     | CD34             | Phycoerythrin (PE)       | AC136  | Miltenyi Biotec |
| CD45     | CD45             | Allophycocyanin (APC)    | 2D1    | Biolegend |
| CD45RA   | CD45RA           | Allophycocyanin (APC)    | H100   | Biolegend |
| CD56     | NCAM1            | Pacific Blue (PB)        | 5.1H11 | Biolegend |
| CD56     | NCAM1            | Allophycocyanin (APC)    | 5.1H11 | Biolegend |
| CD94     | KLRD1            | Peridinin Chlorophyll Protein- Cyanin5.5 (PerCP-Cy5,5) | DX22  | eBioscience™; Thermofisher Scientific |
| EOMES    | EOMES            | Phycoerythrin (PE)       | WD1928 | eBioscience™; Thermofisher Scientific |
| Granyme B| Granyme B        | Phycoerythrin (PE)       | GB11   | eBioscience™; Thermofisher Scientific |
| IFN-γ    | Interferon-γ     | eFluor 660               | 4S.B3  | eBioscience™; Thermofisher Scientific |
| IL-15    | HELIOS           | Allophycocyanin (APC)    | 22F6   | eBioscience™; Thermofisher Scientific |
| Perforin | Perforin         | Phycoerythrin (PE)       | delta G9 | Biolegend |
| T-BET    | T-BET            | Phycoerythrin (PE)       | 4B10   | eBioscience™; Thermofisher Scientific |

**Functional Assays**  
**IFN-γ Production**  
For intracellular IFN-γ detection, 10⁵ cells from day 21 T-BET or EOMES overexpression and control cultures were stimulated with 50 ng/ml phorbol myristate acetate (PMA) and 1 µg/ml ionomycin (both from Sigma-Aldrich) or with 10 ng/ml IL-12 (PeproTech) and 10 ng/ml IL-18 (R&D Systems) with or without 10 ng/ml IL-15 (Miltenyi Biotec) for 24 h. The final 4 h, brefeldin A (BD GolgiPlug, 1/1000, BD Biosciences) was added. Intracellular IFN-γ staining was performed using Cytofix/ Cytoperm Kit (BD Biosciences) and analyzed on gated NK cells by flow cytometry.

**Cytotoxicity Assays**  
For cell specific killing K562, Daudi, RL, Raji (all from ATCC, Manassas, VA) and Nalm-6 (DSMZ, Braunschweig, Germany) target cells were challenged with sorted (eGFP™CD45™CD56+) NK cells of day 21 control, T-BET and EOMES overexpression, eGFP™CD45™CD56+ HPC or naive CD34+ HPC were challenged with sorted (eGFP+CD45+CD56+) early effects of T-BET and EOMES overexpression, eGFP+CD45+CD56+ (hereafter referred to as day 0) and transcriptome and epigenetic Regulation by T-BET or EOMES results in an NK Cell Transcriptome

**Statistical Analysis**  
Statistical analysis was performed with GraphPad Prism 9.0.0 software (San Diego, CA) using Wilcoxon signed rank test or t-test. P-values of p<0.05 were considered as statistically significant.

**RESULTS**  
**Epigenetic Regulation by T-BET or EOMES Overexpression in HPC Results in an NK Cell Transcriptome**  
Separate overexpression of T-BET or EOMES was obtained by retroviral transduction in human umbilical cord blood (UCB)-derived CD34⁺ HPC, in parallel to the control vector, only containing the eGFP-reporter gene. To evaluate the direct and early effects of T-BET and EOMES overexpression, eGFP⁺ lineage CD34⁺ HPC were sorted after 48 h of transduction (hereafter referred to as day 0) and transcriptome and chromatin accessibility profiling were performed by RNA- and Fast-ATAC sequencing, respectively. The gating strategy of the HPC sort is depicted in **Supplementary Figure 1A**. Flow cytometric analysis confirmed the expression of both proteins in the transduced cells (**Supplementary Figure 1B**). mRNA expression analysis revealed that genes expressed in HPC or
involved in quiescence of hematopoietic stem cells were downregulated, including \textit{CD34} and \textit{HOPX} (33). Also, genes involved in erythropoiesis, platelet formation or myeloid differentiation, including \textit{TAL1}, \textit{LM02}, \textit{MMRN1} and \textit{MPO} (34–36), were downregulated in T-BET- and/or EOMES-overexpressing HPC. In contrast, a large set of genes encoding transcription factors with a proven role in murine and/or human NK cell development was upregulated, including \textit{ETS-1}, \textit{IKZF2}, \textit{IKZF3}, \textit{ZNF683}, \textit{IRF8}, \textit{TOX}, \textit{TOX2}, \textit{TXNIP}, \textit{TCF7} and \textit{ZBTB16} (3, 8, 32, 37–39). In addition, genes encoding NK cell developmental and/or cytotoxicity markers were upregulated as well, including \textit{IL2RB}, \textit{KLRD1}, \textit{SLAMF7}, \textit{ITGAL}, \textit{NCR3}, \textit{CD244}, \textit{PRF1}, \textit{GZMB}, \textit{GZMH} and \textit{FASLG} (Figure 1A). Differential gene expression of selected genes was confirmed by RT-qPCR or flow cytometry (Supplementary Figures 1C, D). The transcriptome profiles of T-BET- and EOMES-overexpressing HPC were partially overlapping, yet subtle differences exist, e.g. \textit{MMRN1}, \textit{ETS-1}, \textit{TOX2}, \textit{PRF1} and \textit{FASLG} were exclusively differentially regulated by EOMES overexpression, while \textit{TCF7} and \textit{ZBTB16} were only upregulated with T-BET overexpression (Figure 1A; Supplementary Table 1). Overall, more differentially expressed genes were identified in EOMES-overexpressing (727 UP, 698 DOWN) compared to T-BET-overexpressing HPC (333 UP, 239 DOWN) (Figure 1B, upper part). The same trend was observed in the number of differential ATAC sites, whereby EOMES-overexpressing HPC had noticeably more differential ATAC sites (18283 UP, 10591 DOWN) compared to T-BET-overexpressing HPC (12817 UP, 5978 DOWN) (Figure 1B, lower part). The majority of the T-BET-induced up- and downregulated ATAC regions were similarly regulated upon EOMES overexpression. However, a relatively larger part of EOMES-induced up- and downregulated ATAC regions was not differentially regulated by T-BET overexpression (Figure 1C). To investigate whether T-BET or EOMES overexpression in HPC regulates the overall chromatin accessibility of NK cell-linked genes, we first defined the NK cell-linked ATAC regions by comparing control stage 4 NK cells to control HPC, that were then compared to differential ATAC sites of T-BET- or EOMES-overexpressing HPC versus control HPC. The results show indeed that the majority of more and less accessible NK cell-linked ATAC regions corresponded to up- and down-regulated ATAC regions, respectively, upon T-BET or EOMES overexpression in HPC (Figure 1D).

Analysis of the overlap between the transcriptome and chromatin accessibility profile of HPC showed that the majority of the upregulated genes upon T-BET or EOMES overexpression had either non-differential or upregulated ATAC sites, or contained both up- and downregulated ATAC sites. A minority of these genes contained downregulated ATAC sites only. Whereas a higher fraction of genes with downregulated mRNA expression displayed downregulated ATAC sites, a relatively high fraction also contained non-differential or upregulated ATAC sites (Figure 1E). Mainly T-BOX motifs, followed by RUNX and ETS motifs, were present in upregulated ATAC regions upon T-BET or EOMES overexpression, whereas mainly ETS motifs, followed by GATA and RUNX motifs, were prevalent in downregulated ATAC regions (Figures 1F, G and Supplementary Table 2)

We inspected the chromatin landscape of selected genes that were up- or downregulated in HPC upon T-BET or EOMES overexpression, and combined this analysis with available H3K27ac ChIP-seq data (40, 41). Somewhat unexpected, HPC-linked genes downregulated upon T-BET or EOMES overexpression, like \textit{CD34} and \textit{HOPX}, showed particular DNA regions that became more accessible and these open regions were also present in stage 4 control NK cells. This indicates that these ATAC regions are presumably important for the repression of \textit{CD34} and \textit{HOPX} transcription. Other downregulated genes involved in non-NK lineage differentiation showed less accessible DNA regions in T-BET- or EOMES-overexpressing HPC, whereby these particular DNA regions were closed in stage 4 control NK cells and the majority of these regions also lacked a T-BOX DNA binding motif (Figure 1H; upper part). Genes of several NK cell-linked transcription factors and developmental markers showed upregulated ATAC sites in HPC upon T-BET or EOMES overexpression, and these DNA regions were also accessible in stage 4 control NK cells. Comparison to available mature NK cell-derived H3K4me3 and H3K27ac histone-ChIP data (41), that respectively reveal active promoters and poised or active promoters and enhancers, shows that promoters and/or enhancers were activated upon T-BET or EOMES overexpression. The large majority of these promoters and enhancers contained a T-BOX DNA binding motif (Figure 1H; middle part), in sharp contrast to the less accessible DNA regions, as mentioned above (Figure 1H; upper panel). Gene set enrichment analysis (GSEA) further demonstrated that the transcriptome of T-BET or EOMES overexpressing HPC was highly enriched for mature CD56dim NK cell specific transcripts (Figure 1I).

Altogether, these results indicate that T-BET and EOMES overexpression in HPC results in epigenetic changes in the chromatin landscape of NK cell-linked genes and induces an NK-cell specific transcriptome. The effects of T-BET or EOMES overexpression on the transcriptome and chromatin accessibility of HPC are largely overlapping, in which EOMES plays a more pronounced role.

**NK Cell Differentiation Upon T-BET or EOMES Overexpression in Human HPC Is Drastically Accelerated**

Transduced and eGFP+-sorted HPC were cultured in NK cell differentiation conditions for 21 days. Kinetic analysis of T-BET or EOMES expression in NK cells generated from control versus T-BET- or EOMES-overexpressing HPC demonstrated that T-BET and EOMES expression was maintained throughout the 21 day culture period (Supplementary Figure 2A). After 3 days of culture, nearly no HSC (\textit{CD34}linage \textit{CD45RA}+), stage 1 (\textit{CD34}+ \textit{CD45RA}+\textit{CD117}+) and stage 2 (\textit{CD34}+\textit{CD45RA}+\textit{CD117}+) cells remained in the T-BET and EOMES overexpression cultures, whereas these populations were still clearly present in control transduced cultures. In addition, significantly less stage 3 (\textit{CD34}+\textit{CD117}+\textit{CD94}) cells were found upon EOMES overexpression in comparison to the control (Figure 2A). In sharp contrast, mature
**FIGURE 1** | Epigenetic regulation by T-BET or EOMES overexpression in human HPC results in an NK cell transcriptome. Human eGFP+CD34+ HPC transduced with control, T-BET or EOMES constructs were sorted at day 0 (after 48 h of transduction) and RNA-seq (n=5) and Fast-ATAC-seq (n=3) was performed. (A) Fold change plots illustrating differential mRNA expression of T-BET or EOMES transduced versus control transduced HPC. Blue and red dots indicate T-BET- and EOMES-specific differential genes, respectively. Orange dots indicate genes differentially regulated by both overexpression constructs. Selected differential genes are highlighted. (B) Bar charts showing the number of differential events of RNA- and ATAC-seq for T-BET- or EOMES-overexpressing versus control HPC. (C) Overlap analysis of differential ATAC sites in HPC overexpressing T-BET versus EOMES. (D) NK cell-linked ATAC sites were defined as differential ATAC sites of stage 4 control NK cells versus control HPC. These were then compared to the differential ATAC regions of T-BET- or EOMES-overexpressing versus control HPC. Bar charts show relative numbers of ATAC sites. (E) Overlap analysis of up- or downregulated mRNA and the indicated subgroups of ATAC sites for T-BET- or EOMES-overexpressing HPC. Relative numbers of ATAC sites are shown in the bar chart. (F, G) Motif enrichment analysis of up- or downregulated, or non-differential ATAC sites of T-BET- or EOMES-overexpressing HPC. (F) The top 5 best matched motifs are indicated. (G) The relative presence of the indicated motif families is shown. (H) Representative Genome Browser views of selected gene loci, showing tracks of ATAC-seq for the indicated HPC samples and stage 4 control NK cells, and H3K27ac and/or H3K4me3 ChIP-seq of HPC and mature NK cells. Differentially regulated ATAC regions are indicated in grey or orange, whereby the orange colored regions contain a T-BOX motif. (I) Gene set enrichment analysis (GSEA) shows a significant enrichment of CD56dim signature genes in T-BET- or EOMES-overexpressing vs. control HPC. Normalized enrichment scores (NES) and q-values are indicated. See also Supplementary Figure 1.
FIGURE 2 | NK cell differentiation upon T-BET or EOMES overexpression in human HPC is drastically accelerated. T-BET, EOMES or control transduced HPC were cultured in the *in vitro* NK cell differentiation co-culture. Cultures were analyzed by flow cytometry on pre-gated eGFP^+CD45^+ cells on different time points as indicated. (A) Absolute cell numbers of HSC (CD34^+CD45RA^−), stage 1 (CD34^+CD45RA^+CD117^−), stage 2 (CD34^+CD45RA^+CD117^+) and stage 3 cells (CD34^−CD117^+CD94^−) at day 3 of culture (mean ± SEM; n = 5–6). (B) Representative dot plots of CD11a^+ gated cells at the indicated days of culture. Cells in the upper right quadrant represent the CD56^+CD94^+ NK cell population. The numbers in the plots indicate the percentages. Absolute cell numbers of stage 4 (CD16^−) and stage 5 (CD16^+) NK cells are shown in the bar charts (mean ± SEM; n = 6–9). (C) Percentages of stage 5 (CD16^+) of total NK cells is shown (mean ± SEM; n = 6–9). (D) Microscopic images of sorted CD56^+CD94^+ NK cells on day 3 and 7 of overexpression cultures and on day 19 of control cultures. Arrows indicate cytotoxic granules. 100× magnification. Scale bar = 5μm. (E) Representative dot plots of CD11a^+ gated cells from cultures in the absence of exogenous IL-15. (F) Representative histograms of the indicated NK cell markers of gated NK cells (CD11a^+CD56^+CD94^+), whereby the overexpression conditions at different days of culture are compared to day 21 control NK cells. The fluorescence minus one (FMO) is included as background fluorescence. Percentages or MFI of the indicated marker is presented in the bar charts (mean ± SEM; n = 6–12). *, **, *** and **** indicate significant difference of the specified overexpression condition compared to the control, with p<0.05, <0.01, <0.001, <0.0001, respectively. §, §§ and §§§ indicate significant difference of T-BET vs. EOMES overexpression, with p<0.05, <0.01, 0.001, respectively. See also Supplementary Figure 2.
stage 4 (CD56"CD94+CD16") NK cells already appeared at day 3 of T-BET and EOMES overexpression cultures, whereas these cells merely become detectable at day 14 of control cultures (Figure 2B). In addition, maturation towards stage 5 (CD56"CD94+CD16") NK cells was accelerated and increased throughout the overexpression cultures, where these cells were clearly present from day 7 compared to few cells at day 14 in control cultures (Figure 2B and Supplementary Figure 2B). Throughout the culture, the percentage of CD16-expressing cells with T-BET overexpression was significantly increased compared to the control condition, but was highest with EOMES overexpressing cells (Figure 2C and Supplementary Figure 2B). Microscopic imaging of sorted CD56"CD94" NK cells from T-BET or EOMES overexpression cultures at day 3 and 7 of culture, showed the clear presence of cytotoxic granules, resembling an NK cell morphology (Figure 2D).

Because T-BET and EOMES overexpression in HPC increased the speed of NK cell development, the possibility existed that T-BET or EOMES overexpression overruled the need for IL-15, a cytokine driving NK cell differentiation, maturation and survival through IL2Rβ signaling (3). However, cultures in the absence of exogenous IL-15 showed no NK cell generation in T-BET or EOMES overexpression cultures (Figure 2E).

To further phenotype these early arising NK cells, a more in-depth kinetic analysis of the expression of activating and inhibitory NK receptors was performed on gated CD56"CD94" cells from days 3, 7 and 21 overexpression cultures and were compared to day 21 control NK cells (Supplementary Figure 2C). The earliest receptor expressed on NK cells generated from T-BET- or EOMES-overexpressing HPC was NKG2D, with even higher NKG2D expression on days 3 and 7 compared to control NK cells. Expression of the activating receptors NKp46 and NKp30, and the inhibitory receptor NKG2A gradually increased to control levels in the T-BET cultures, whereas expression in EOMES cultures was already high from day 3 (NKp46 and NKp30) or day 7 (NKG2A). DNAM-1 and NKp44 expression was delayed and control levels were reached from day 21 in both overexpression conditions. Finally, the cytotoxic effector proteins, perforin and granzyme B, were early expressed in both overexpression cultures, whereby the expression of these proteins was even higher on days 3 and 7 compared to control NK cells (Figure 2F).

In conclusion, T-BET and EOMES overexpression in HPC drastically accelerates NK cell generation, whereby these NK cells have a granular morphology, display early or gradual expression of NK receptors and early expression of perforin and granzyme B. In this model, EOMES is a stronger driver of the early NK cell phenotype and a better enhancer of CD16 expression than T-BET. Nevertheless, NK cell differentiation in T-BET or EOMES overexpression conditions remains IL-15 dependent.

**Differentiating NK Cells Upon T-BET or EOMES Overexpression Undergo Terminal NK Cell Maturation, Whereby T-BET Plays a Predominant Role**

To obtain an in-depth analysis of the effects of T-BET or EOMES overexpression in mature NK cells, RNA and Fast-ATAC sequencing was performed on isolated stage 4 (CD16+) and stage 5 (CD16+) NK cells from day 21 control, T-BET and EOMES overexpression cultures (Supplementary Figure 3A). Transcriptome analysis of stage 4 NK cells upon T-BET or EOMES overexpression showed upregulation of IKZF2, IKZF3, RORA, PRDM1 and ZEB2, that are transcription factor genes known to be involved in terminal NK cell maturation (3, 42–45). In addition, CX3CR1, S1PR5 and KLRG1, characteristic for terminal NK cell maturation (45, 46), were exclusively upregulated in stage 4 and/or stage 5 NK cells from T-BET overexpression cultures (Figure 3A and Supplementary Figure 3B). Furthermore, IL7R, which is known to be downregulated in CD56dim versus CD56bright NK cells (47), was also downregulated in stage 4 NK cells from both T-BET and EOMES overexpression cultures (Figure 3A). Comparable transcriptional differences between T-BET- or EOMES-overexpression conditions versus control cells were found in stage 5 NK cells (Supplementary Figure 3B). Importantly, when comparing the transcriptome of stage 5 versus stage 4 NK cells, there were 671 differentially expressed genes in the control cultures, whereas there were only 54 and 102 differentially expressed genes in the T-BET- and EOMES-overexpression cultures, respectively (Figure 3B, right panel; Supplementary Table 3). These findings suggest that stage 4 NK cells upon T-BET or EOMES overexpression share many similarities with stage 5 control NK cells, further confirming terminal NK cell maturation. Similar to the results in HPC, more genes were differentially expressed in NK cells from EOMES compared to T-BET cultures, in both stage 4 and stage 5 NK cells (2629 vs. 1564 and 2763 vs. 1238 genes, respectively) (Figure 3B, left panel; Supplementary Table 4). However, the opposite was true for the number of differential ATAC sites, which was significantly lower in NK cells upon EOMES overexpression compared to T-BET overexpression (4876 vs. 10 278 and 4601 vs. 10 013 sites in stage 4 and stage 5, respectively) (Figure 3B, left panel). As indicated above, EOMES-overexpressing HPC had more differential ATAC sites than T-BET-overexpressing HPC (Figure 1B), which raised the possibility that EOMES overexpression already regulates the chromatin accessibility of genes in HPC that only become regulated by T-BET overexpression from stage 4 NK cells. However, as overlap analysis showed that most ATAC sites that are differential in T-BET-overexpressing vs. control stage 4 NK cells are not differential in EOMES-overexpressing vs. control HPC (Figure 3C), this is not the case. This implies, as expected, that chromatin accessibility continues to be altered during HPC differentiation towards stage 4 NK cells. Analysis of the overlap between the transcriptome and chromatin accessibility profile of stage 4 NK cells showed differences between the T-BET- and EOMES-overexpression cultures. While in cells from T-BET-overexpression cultures a considerable part of the genes with up- or downregulated transcription contained up- or downregulated ATAC sites, respectively, this was considerably less evident upon EOMES overexpression. Here, more than 70% of the ATAC sites was not differentially regulated compared to control NK cells (Figure 3D), indicating that transcription is considerably less
regulated by epigenetic changes on the chromatin level. Finally, in contrast to the findings in HPC (Figure 1E), few differentially expressed genes contained both up- and downregulated ATAC sites (Figure 3D). Similar results were obtained with stage 5 NK cells (Supplementary Figure 3C). Mainly ETS motifs, followed by T-BOX and RUNX motifs were present in upregulated ATAC regions of stage 4 and stage 5 NK cells generated upon T-BET or EOMES overexpression in HPC, while predominantly RUNX motifs followed by ETS and T-BOX motifs were prevalent in downregulated ATAC sites (Figures 3E, F and Supplementary Figures 3D, E and Supplementary Table 5).

Detailed inspection of the chromatin landscape of IKZF2, IKZF3, RORA and ZEB2, that encode transcription factors linked to mature NK cell differentiation (3, 42–45) and that showed increased mRNA expression in NK cells upon T-BET or EOMES overexpression, revealed that promoter and/or putative enhancer sites became more accessible (Figure 3G; upper panel). In sharp contrast, the accessibility of promotor and/or

FIGURE 3 | NK cells developing from T-BET- or EOMES-overexpressing HPC undergo terminal NK cell maturation. eGFP+CD45+CD56+CD94+ stage 4 (CD16-) and stage 5 (CD16+) NK cells were sorted from day 21 cultures and RNA-seq (n=4) and Fast-ATAC-seq (n=2) was performed. (A) Fold change plots illustrating differential mRNA expression of T-BET or EOMES versus control stage 4 NK cells. Blue and red dots indicate T-BET- and EOMES-specific differential genes, respectively. Orange dots indicate genes differentially regulated by both overexpression constructs. Selected differential genes are highlighted. (B) Charts show the number of differential events of RNA- and ATAC-seq for T-BET or EOMES stage 4 and stage 5 NK cells compared to control NK cells (left panel) and for stage 4 compared to stage 5 NK cells (right panel). (C) Overlap analysis of differential ATAC sites of T-BET vs. control stage 4 NK cells and of EOMES-overexpressing vs. control HPC. Charts demonstrate the absolute number of ATAC sites. (D) Overlap analysis of up- or downregulated mRNA and the indicated subgroups of ATAC sites for T-BET or EOMES versus control stage 4 NK cells. Relative numbers of ATAC sites are shown in the bar charts. (C, D) Up = upregulated sites; down = downregulated sites; both = up- and downregulated sites; no diff = detectable sites without differential expression; none = non-detectable sites. (E, F) Motif enrichment analysis of up- or downregulated, or non-differential ATAC sites of T-BET or EOMES compared to control stage 4 NK cells. (E) The top 5 best matched motifs are indicated. (F) The relative presence of the indicated motif families is shown. (G) Representative Genome Browser views of selected gene loci, showing tracks of ATAC-seq for the indicated stage 4 and stage 5 NK cells, and of H3K27ac and H3K4me3 ChIP-seq of mature NK cells. Differentially regulated ATAC regions are indicated in grey or orange, whereby the orange colored regions contain a T-BOX motif. See also Supplementary Figures 3 and 4.
putative enhancer sites of CX3CR1, S1PR5 and KLRG1 exclusively increased upon overexpression of T-BET (Figure 3G; lower panel) and this correlated with the increased mRNA expression in T-BET, but not in EOMES overexpression cultures. Conversely, IL7R displayed downregulated ATAC sites, corresponding to decreased mRNA expression (Figure 3G; lower panel). Corresponding to the motif enrichment analysis (Figures 3E, F), T-BOX DNA binding motifs were less prevalent in the upregulated ATAC regions (Figure 3G), in contrast to HPC accessible ATAC regions whereby T-BOX motifs were predominant (Figure 1H).

Further analysis of the transcriptome showed that there was no detectable KIR mRNA expression in control HPC, whereas there was a gradual increase in control stage 4 and stage 5 NK cells, as expected. Neither T-BET nor EOMES overexpression induced KIR mRNA in HPC. Whereas EOMES overexpression did not affect (KIR2DS4 and KIR2DL3) or slightly increased (KIR2DL1 and KIR3DL1) KIR mRNA expression in stage 4 and stage 5 NK cells, T-BET overexpression resulted in a drastic increase of all these KIR mRNAs (Figure 4A). Strikingly, the chromatin landscape revealed an upregulated ATAC region in KIR2DL1 and KIR2DS4 genes in HPC overexpressing T-BET or EOMES, although no mRNA was expressed. These ATAC regions were also present in control stage 4 and stage 5 NK cells. In stage 4 and stage 5 NK cells upon T-BET or EOMES overexpression, promotor regions of KIR2DL1, KIR2DL3, KIR2DS4 and KIR3DL1 were more accessible compared to control NK cells, whereby these ATAC regions were

![Figure 4](image-url)
NK cells and CD56 bright and CD56 dim cells cannot be activating effect of IL-15, CD56 expression is increased on all BET or EOMES overexpression cultures (data not shown).

In the in vitro NK cell co-cultures, probably due to the activating effect of IL-15, CD56 expression is increased on all NK cells and CD56bright and CD56dim cells cannot be distinguished. We determined the mean fluorescence intensity (MFI) of CD56 in gated NK cells (CD56^bright^CD94^-^) at different time points of culture. At day 3 and 7 of culture, NK cells from T-BET or EOMES overexpression cultures had a significantly lower CD56 MFI as compared to NK cells from day 21 control cultures. In contrast to NK cells from day 21 control cultures, where as expected stage 4 cells expressed lower KIR levels than stage 5 cells, there was no clear difference in KIR expression in stage 4 vs stage 5 NK cells from T-BET or EOMES overexpression cultures (data not shown).

In conclusion, the transcriptome, chromatin accessibility and protein expression profile gave insights on how T-BET and EOMES epigenetically regulate NK cell maturation and subsequently activate the terminal NK cell transcriptome, wherein T-BET plays a more potent role.

**T-BET and EOMES NK Cells Have Increased Functionality and EOMES NK Cells Display Enhanced ADCC**

An important function of human NK cells is cytotoxic killing of malignant or virus-infected cells. NK cells from day 21 overexpression and control cultures were challenged with several target cell lines. K562 cells, the prototypic NK cell susceptible target cell line (48), was similarly killed by NK cells from all conditions. Additionally, the human B cell lymphoma cell lines Daudi, RL, and Raji, were only killed to a low extent by NK cells from control cultures, whereas killing by NK cells from T-BET or EOMES overexpression cultures was significantly increased. In contrast, Nalm-6, another human B cell line, displayed resistance to efficient killing by NK cells from all culture conditions (Figure 5A).

In addition to direct recognition of tumor targets, NK cells also lyse malignant cells upon binding of tumor cell-bound antibody by the CD16 receptor expressed on stage 5 NK cells, a process known as ADCC (49). Substantially more CD16^+^ NK cells were obtained with NK cells from T-BET and EOMES overexpression cultures (Figures 2B, C), whereby both the percentage as well as the level of CD16 expression was higher in EOMES overexpression cultures (Figure 2B and Supplementary Figure 5A). Therefore, the ADCC capacity was tested using the CD20-expressing human Burkitt’s lymphoma cell line Raji in the absence or presence of Rituximab, a humanized monoclonal anti-CD20 antibody (50). The results show that inclusion of Rituximab increased killing, as expected, but importantly, NK cells from EOMES overexpression cultures displayed significantly higher ADCC functionality as compared to the control or T-BET overexpression conditions (Figure 5B). The chromatin landscape of FCGR3A, the gene encoding CD16 in NK cells (49), was highly similar in NK cells from all culture conditions (Supplementary Figure 5B), indicating that increased CD16 expression upon T-BET or EOMES overexpression is not epigenetically regulated.

Another crucial function of NK cells is production of pro-inflammatory cytokines, including IFN-γ, by which NK cells regulate other immune cells. IFN-γ production upon PMA/Ionomycin stimulation of day 21 NK cells was comparable between conditions. However, IL-12/IL-18 or IL-12/IL-15/IL-18 stimulation resulted in significantly higher IFN-γ production of NK cells upon T-BET or EOMES overexpression compared to control NK cells (Figure 5C). The transcriptome profile showed that NK cells from T-BET- or EOMES-overexpression cultures had upregulated mRNA expression of ICOS, LBH, IL12RB2, IL18R1 and IL18RAP (Figure 3A). These NK cells also displayed increased accessibility of promoter and enhancer regions in ICOS and IL12RB2, whereas enhancer regions in LBH and IL18RAP were upregulated mainly by NK cells from T-BET overexpression cultures (Figure 5D).

These findings show that NK cells generated from T-BET- or EOMES-overexpression HPC have increased cytotoxic capacity against certain tumor targets and NK cells from EOMES overexpression cultures have enhanced ADCC capacity. The NK cells generated upon T-BET or EOMES overexpression display increased IFN-γ production upon cytokine stimulation and this increase is potentially epigenetically regulated.

**DISCUSSION**

Overexpression of T-BET or EOMES in UCB-derived HPC alters the chromatin landscape, correlating with loss of an HPC-related transcriptome and induction of an NK cell transcriptome. Subsequent NK cell differentiation is accelerated, with increased ADCC potential in EOMES cultures and increased KIR expression in T-BET cultures.
T-BET and EOMES are the signature transcription factors defining murine and human mature NK cells, but information on their role in human NK cell differentiation is limited. Murine studies have shown that Eomes and T-bet play a crucial role in early and late NK cell differentiation, respectively. Mouse NK cells mature in a 4-stage developmental program, whereby CD27lowCD11blow cells develop successively via CD27+CD11blow and CD27+CD11b+ intermediate stages into CD27lowCD11b+ terminally mature NK cells (51). Eomes- or T-bet-deficient mice display reduced NK cell numbers, wherein Eomes-deficient mice show less differentiation past the CD27lowCD11blow stage, whereas in T-bet -/- mice NK cell development is halted at the CD27+CD11b+ stage. T-bet/Eomes double-deficient mice have a complete block in NK cell development, indicating that these highly homologous T-box transcription factors also have non-redundant functions (16, 17). During murine NK cell maturation, the levels of Eomes and T-bet inversely correlate and this is consistent with Eomes acting upstream of T-bet in the NK cell maturation cascade (17, 52). Chromatin accessibility and transcriptome profiling following T-BET or EOMES overexpression in HPC provide novel insights on the regulatory role of these transcription factors in humans. Both T-BET and EOMES downregulated HPC-linked genes and genes involved in erythropoiesis, platelet formation or myeloid differentiation, whereas they upregulated genes of NK cell-linked transcription factors, membrane receptors and effector proteins. Overall, more differentially expressed genes were identified in EOMES-overexpressing compared to T-BET-overexpressing HPC. Interestingly, ETS-1, which has a proven role in both murine and human NK cell development (32, 53, 54), was only upregulated upon EOMES overexpression. The effects of T-BET and, more pronounced, of EOMES on the transcriptome were reflected in the chromatin landscape. It was striking that a large part of the more accessible ATAC regions contained a T-BOX motif, suggesting transcriptional regulation by T-BET or EOMES, whereas in the majority of downregulated ATAC regions ETS, GATA and RUNX motifs were prevalent. In this context, there is ample evidence that ETS, GATA and RUNX transcription factors bind to T-BET and/or EOMES, thereby inhibiting or inducing DNA binding of these co-factors (55–59).

T-BET or EOMES overexpression both accelerated differentiation of HPC into NK cells, but our kinetic NK cell receptor analysis showed that Nkp46, Nkp30 and NKG2A were readily expressed upon EOMES overexpression, while expression was delayed upon T-BET overexpression. Furthermore, EOMES was a better enhancer of CD16 expression in NK cells, whereby
both the percentage of CD16-expressing cells as well as the level of CD16 expression was higher as compared to T-BET overexpression. Altogether, this indicates that also in human predominantly EOMES has a role during development of HPC into CD16-expressing NK cells. This is compatible with studies showing that a T-BET/EOMES gradient is present during human NK cell maturation, wherein EOMES is expressed prior to T-BET. T-BET upregulation during further NK cell differentiation is mirrored by EOMES downregulation, whereby EOMES\textsuperscript{low}-T-BET\textsuperscript{high} NK cells are terminally mature (18, 19, 60).

In human NK cells, KIR upregulation correlates with decreased EOMES and increased T-BET levels, with highest KIR expression in EOMES\textsuperscript{low}-T-BET\textsuperscript{high} terminally mature NK cells (14, 19). Our data show that, although NK cells from EOMES overexpression cultures had increased KIR expression compared to control NK cells, KIR expression was highest in T-BET overexpression cultures. This was reflected in increased chromatin accessibility and in higher mRNA and protein levels. The functional homologs of KIRs are Ly49 receptors in mice. Eomes-deficient mice exhibit a limited repertoire of Ly49 receptors, with decreased expression of Ly49A/D/G2/H, whereas T-bet-deficient NK cells express a normal repertoire of activating and inhibitory receptors, with a higher proportion of Ly49D and Ly49G2 (16, 17). Our data show that although EOMES can establish KIR expression in human NK cells, T-BET is superior in this regard. This might implicate that during physiological terminal human NK cell differentiation T-BET further increases KIR expression that is initiated by EOMES. Our data further show that both chromatin accessibility and transcription of genes known to be involved in terminal maturation, including ZEB2, CX3CR1, S1PR5 and KLRG1 (43, 45, 46), were upregulated upon T-BET overexpression. As shown in mice and suggested in human (16–19, 52), our data are in agreement with a crucial role of T-BET in terminal NK cell maturation.

Proper NK cell functionality is mandatory for an effective immune response against malignant cells. NK cells generated upon T-BET or EOMES overexpression in HPC showed efficient killing of NK-sensitive K562 cells and increased killing of Daudi, RL and Raji tumor cell lines. The latter cell lines are partially resistant to NK cell-mediated lysis as they express HLA-class I molecules and/or lack ligands for activating NK receptors, like NKGD2 (48, 61, 62). Intrinsic potential of discrete NK cell subsets is established during NK cell differentiation and it has been shown that terminally mature subsets of NK cells display superior ability to kill target cells in comparison to less mature subsets (63).

NK cells from EOMES cultures additionally showed enhanced ADCC potential, that correlated with increased CD16 expression. This was a surprise, as ex vivo stage 5 NK cells express CD16 and display ADCC functionality and these cells are mainly EOMES\textsuperscript{low}-T-BET\textsuperscript{high}. However, there is more complexity in our culture model as NK cells from EOMES overexpression cultures also express endogenous T-BET, and the reverse is true for NK cells from T-BET overexpression cultures. This is clear from the kinetic analysis of EOMES and T-BET expression, as shown in Supplementary Figure 2A. This kinetic analysis also shows that, whereas ectopic EOMES expression is very high from day 0 to day 7, it equals the endogenous EOMES expression of control cells at day 14 and it even becomes lower than endogenous EOMES expression of the control cells at day 21. Our hypothesis is that the high ectopic expression of EOMES early in the culture programs the cells for increased ADCC capacity once the ectopic EOMES expression decreases later on.

CD16 expression is crucial in NK cell activation against tumor cells via ADCC. In vitro generated NK cells from induced pluripotent or UCB-derived stem cells, commonly used as a source of NK cells for adoptive cell transfer therapy, display low levels of CD16 expression (64–67). Attempts to increase CD16 expression of these NK cell products proves its importance in adoptive NK cell therapy (67). The importance of CD16 is further illustrated by therapeutic approaches focusing on monoclonal antibodies against tumor targets, in which the Fc-receptor regions are modified to increase CD16-binding and subsequently improve ADCC activity of the transferred NK cells (68, 69). The synergistic effects of tumor-targeting monoclonal antibodies and transfusion of NK cells have been shown and are further investigated in ongoing clinical trials (70, 71). Moreover, CD16 expression facilitates not only the combination of transferred NK cells with tumor-targeting monoclonal antibodies to exploit ADCC, but it also plays a crucial role in functionality of Bi- or Tri-specific killer engagers containing a single-chain scFv against the tumor antigen and CD16 (72). Therefore, strategies to improve the ADCC potential of NK cells provide a valuable tool to optimize adoptive NK cell therapies.

NK cells are potent producers of pro-inflammatory cytokines, including IFN-γ (73). The early burst of IFN-γ secretion is not impaired in murine T-bet-deficient NK cells, but after 24 h of stimulation significantly less IFN-γ is produced. This suggests that the presence of T-bet is important for a positive feedback loop on IFN-γ expression (16, 17). A possible role for Eomes in inducing IFN-γ in the absence of T-bet is seen in murine CD8 T cells (74). IFN-γ is a direct target of T-bet in NK cells, whereas no evidence exists for Eomes-binding to the IFN-γ locus (16). In human, CD56\textsuperscript{bright} NK cells efficiently produce IFN-γ after IL-12/IL-18 stimulation, which correlates with increased IL12RB2 and IL18RA mRNA expression (60). Our results show that NK cells generated upon T-BET or EOMES overexpression in HPC produced significantly more IFN-γ after IL-12/IL-15/IL-18 stimulation in comparison to control NK cells. Besides, these NK cells displayed upregulated mRNA expression of ICOS, LBH, IL12RB2 and IL18RAP, with increased accessibility of promoter and/or enhancer regions of these gene loci. During NK cell maturation, epigenetic alterations in signaling components occur (3). T-bet is known to recruit JMJD3 and SET7 in Th1 cells which generate chromatin remodeling of the IL12RB2 gene, leading to gene expression (55). Also the IL18R1/IL18RAP gene locus is a T-bet target in T cells (75). The role of EOMES in regulating signaling components during NK cell activation is less clear. The transcriptional activator LBH activates activator protein-1 (AP-1) and is thus involved in the MAPK-pathway employed in IL-18 signaling (76), although a role for LBH in NK cells is currently unknown. A recent study revealed that ICOS plays and important role in NK cell functionality, whereby ICOS-deficient mice display lower IFN-γ levels after in vivo
stimulation (77). Both transcriptional regulators possibly attribute to the increased IFN-γ production in our T-BET or EOMES cultures, due to enhanced downstream signaling after cytokine stimulation.

Altogether, our findings give molecular insights in the regulatory role of T-BET and EOMES upon overexpression in human HPC and subsequent NK cell differentiation, and can be used to optimize adoptive NK cell therapies.

DATA AVAILABILITY STATEMENT

The datasets presented in the study are accessible on GEO with accession number GSE166439 or can be found on https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE166439.

ETHICS STATEMENT

The usage of human umbilical cord blood in this study was reviewed and approved by the Ethics Committee of the Faculty of Medicine and Health Sciences, Ghent University, Ghent, Belgium. The patients/participants provided their written informed consent to participate in this study.

AUTHOR CONTRIBUTIONS

Conceptualization: LK and GL. Methodology: LK, ST, SW, EP, and GL. Investigation: LK, ST, SW, EP, EA, and ZV. Formal Analysis: LK. Software: WL. Visualization: LK and WL. Supervision: GL. Writing – original draft: LK and GL. Writing – review and editing: LK, ST, SW, EP, ZV, PM, FN, TT, PV, BV, and GL. All authors contributed to the article and approved the submitted version.

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REFERENCES

1. Abel AM, Yang C, Thakar MS, Malarkannan S. Natural Killer Cells: Development, Maturation, and Clinical Utilization. Front Immunol (2018) 9:1869. doi: 10.3389/fimmu.2018.01869
2. Caligiuri MA. Human Natural Killer Cells. Blood (2008) 112:461–9. doi: 10.1182/blood-2007-09-077438
3. Vivier E, Di Santo J, Moretta A. Natural Killer Cells. In: Cooper MD, editor. Curr Top Microbiol Immunol. Switzerland: Springer International Publishing (2016). 247. doi: 10.1007/978-3-319-23916-3
4. Freud AG, Yokohama A, Becknell B, Lee MT, Mao HC, Ferketich AK, et al. Evidence for Discrete Stages of Human Natural Killer Cell Differentiation In Vivo. J Exp Med (2006) 203:1033–43. doi: 10.1084/jem.20052507
5. Scoville SD, Freud AG, Caligiuri MA. Modeling Human Natural Killer Cell Development in the Era of Inmate Lymphoid Cells. Front Immunol (2017) 8:360. doi: 10.3389/fimmu.2017.00360
6. Yu J, Freud AG, Caligiuri MA. Location and Cellular Stages of Natural Killer Cell Development. Trends Immunol (2013) 34:573–82. doi: 10.1016/j.it.2013.07.005
7. Eissens DN, Spanholz J, van der Meer A, van Cranenbroek B, Dolstra H, Kwekkeboom J, et al. Defining Early Human NK Cell Developmental Stages in Primary and Secondary Lymphoid Tissues. PLoS One (2012) 7:e30930. doi: 10.1371/journal.pone.0030930
8. Goh W, Huntington ND. Regulation of Murine Natural Killer Cell Development. Front Immunol (2017) 8:130. doi: 10.3389/fimmu.2017.00130
9. Foley B, Cooley S, Verneri MR, Curtissinger J, Luo X, Waller EK, et al. NK Cell Education After Allogeneic Transplantation: Dissociation Between Recovery of Cytokine-Producing and Cytotoxic Functions. Blood (2011) 118:2784–92. doi: 10.1182/blood-2011-04-347070
10. Simonetta F, Pradier A, Bosshard C, Masouridi-Levrat S, Chalandon Y, Roosnek E. NK Cell Functional Impairment After Allogeneic Hematopoietic Stem Cell Transplantation Is Associated With Reduced Levels of T-Bet and Eomesodermin. J Immunol (2015) 195:4712–20. doi: 10.4049/jimmunol.1501522
11. Gill S, Vasey AE, De Souza A, Baker J, Smith AT, Kohrt HE, et al. Rapid Development of Exhaustion and Down-Regulation of Eomesodermin Limit the Antitumor Activity of Adoptively Transferred Murine Natural Killer Cells. Blood (2012) 119:5758–68. doi: 10.1182/blood-2012-03-415364
12. Papaoannou VE. The T-Box Gene Family: Emerging Roles in Development, Stem Cells and Cancer. Development (2014) 141:3819–33. doi: 10.1242/dev.104471
13. Szabo SJ, Kim ST, Costa GL, Zhang X, Fathman CG, Glinccher LH. A Novel Transcription Factor, T-Bet, Directs Th1 Lineage Commitment. Dev.104471
14. Simonetta F, Pradier A, Roosnek E. T-Bet and Eomesodermin in NK Cell Development, Maturation, and Function. Front Immunol (2016) 7:241. doi: 10.3389/fimmu.2016.00241
15. Zhang J, Marotel M, Fauteux-Daniel S, Mathieu AL, Viel S, Marcais A, et al. T-Bet and Eomesoderm Limit the Antitumor Activity of Adoptively Transferred Murine Natural Killer Cells. Eur J Immunol (2018) 48:738–50. doi: 10.1002/eji.201747299
16. Townsend MJ, Weinmann AS, Matsuda JL, Salomon R, Farnham PJ, Biron CA, et al. T-Bet Regulates the Terminal Maturation and Homeostasis of NK and Valpha14i NKT Cells. Immunity (2004) 20:477–94. doi: 10.1016/S1074-7613(04)00076-7
17. Gordon SM, Chaix J, Rupp LJ, Wu J, Madera S, Sun JC, et al. The Transcription Factors T-Bet and Eomes Control Key Checkpoints of Natural Killer Cell Maturation. Immunity (2012) 36:55–67. doi: 10.1016/j.immuni.2011.11.016
18. Knox JJ, Cosma GL, Betts MR, McLane LM. Characterization of T-Bet and Eomes in Peripheral Human Immune Cells. Front Immunol (2014) 5:217. doi: 10.3389/fimmu.2014.00217

19. Collins A, Rothman N, Liu K, Reiner SL. Eomesodermin and T-Bet Mark Developmentally Distinct Natural Killer Cells. JCI Insight (2017) 2: e90063. doi: 10.1172/jcisiht.e90063

20. Kinsella TM, Nolan GP. Episomal Vectors Rapidly and Stably Produce High-Titer Recombinant Retrovirus. Hum Gene Ther (2000) 11:1468–75. doi: 10.1089/106652400372063

21. Lee KN, Kang HS, Jeon JH, Kim EM, Yoon SR, Song H, et al. VDUP1 Is Required for the Development of Natural Killer Cells. Immunity (2005) 22:195–208. doi: 10.1016/j.immuni.2004.12.012

22. Zhang X, Jeong M, Huang X, Wang XQ, Wang X, Zhou W, et al. Large DNA Methylation Nadirs Anchor Chromatin Loops Maintaining Hematopoietic Stem Cell Identity. Mol Cell (2020) 78:506–21.e506. doi: 10.1016/j.molcel.2020.04.018

23. Kousis OI, Collins PL, Cella M, Robinette ML, Porter SJ, Pyfrom SC, et al. Distinct Gene Regulatory Pathways for Human Innate Versus Adaptive Lymphoid Cells. Cell (2016) 165:1134–46. doi: 10.1016/j.cell.2016.04.014

24. Narni-Mancinelli E, Jaeger BN, Bernat C, Fenici A, Kung S, De Gassart A, et al. Tuning of Natural Killer Cell Reactivity by NKp46 and Helios Calibrates T Cell Responses. Science (2012) 335:344–8. doi: 10.1126/science.1215621

25. R Core Team. R: A language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing (2018). [updated 2019/03/11. version 3.5.3.]

26. Andrews S. FastQC: A Quality Control Tool for High Throughput Sequence Data. Babraham Bioinformatics (2010) [updated 2019/01/08. version 0.11.9. Available from: https://www.bioinformatics.babraham.ac.uk/projects/fastqc/]

27. Mootha VK, Lindgren CM, Eriksson KF, Subramanian A, Sihag S, Lehar J, et al. Distinct and Temporary-Restricted Epigenetic Mechanisms Regulate Human Innate Versus Adaptive Lymphoid Cells Defined by Single-Cell Transcriptome. Nat Commun (2019) 10:3931. doi: 10.1038/s41467-019-11947-7

28. Wu Y, Tian Z, Wei H. Developmental and Functional Control of Natural Killer Cells by Cytokines. Front Immunol (2017) 8:930. doi: 10.3389/fimmu.2017.00930

29. Corces MR, Buenrostro JD, Wu B, Greenside PG, Chan SM, Koenig JL, et al. Lineage-Specific and Single-Cell Chromatin Accessibility Charts Human Hematopoiesis and Leukemia Evolution. Nat Genet (2016) 48:1193–203. doi: 10.1038/ng.3646

30. Roels J, Kuchmiy A, De Decker M, Strubbe S, Lavaert M, Liang KL, et al. Tumor- and Cytokine-Primed Human Natural Killer Cells Exhibit Distinct Phenotypic and Transcriptional Signatures. PloS One (2019) 14: e0218674. doi: 10.1371/journal.pone.0218674

31. Wang W, Erbe AK, Hank JA, Morris ZS, Sondel PM. NK Cell-Mediated Antibody-Dependent Cellular Cytotoxicity in Cancer Immunotherapy. Front Immunol (2015) 6:368. doi: 10.3389/fimmu.2015.00368

32. Gluck WL, Hurst D, Yuan A, Levine AM, Dayton MA, Gockerman JP, et al. The Ets-1 Transcription Factor Is Required for the Development of Natural Killer Cells by Cytokines. JEM (2004) 209:1511–24. doi: 10.1101/jem.2003233

33. Lineage-Specific and Single-Cell Chromatin Accessibility Charts Human Hematopoiesis and Leukemia Evolution. Nat Genet (2016) 48:1193–203. doi: 10.1038/ng.3646

34. Ets-1 Transcription Factor Is Required for the Development of Natural Killer Cells by Cytokines. JEM (2004) 209:1511–24. doi: 10.1101/jem.2003233

35. Barton K, Muthusamy N, Fischer C, Ting CN, Walunas TL, Lanier LL, et al. The Ets-1 Transcription Factor Is Required for the Development of Natural Killer Cells in Mice. Immunity (1998) 9:555–63. doi: 10.1016/S1074-7613(00)80638-X

36. Raymire K, Chandler KJ, Spaulding C, Zandi S, Sigvardsson M, Graves BJ, et al. Gene Deregelation and Chronic Activation in Natural Killer Cells Deficient in the Transcription Factor ETS1. Immunity (2012) 36:921–32. doi: 10.1016/j.immuni.2012.04.006

37. Lazarvic V, Gilchrist LH, Lord GM. T-Bet: A Bridge Between Innate and Adaptive Immunity. Nat Rev Immunol (2013) 13:77–89. doi: 10.1038/nri3536

38. Greeningloh R, Kang BY, Ho IC. Ets-1: A Functional Cofactor of T-Bet, Is Essential for Th1 Inflammatory Responses. J Exp Med (2005) 201:615–26. doi: 10.1084/jem.20041330

39. Caret HA, Hildreth BE 3rd, Samuvel DJ, Thies KA, Rosol TJ, Toribio RE, et al. Eomes Partners With PU.1 and MTTF to Regulate Transcription Factors
Kiekens et al.

Critical for Osteoclast Differentiation. *iScience* (2019) 11:238–45. doi: 10.1016/j.isci.2018.12.018

58. Israel A, Bridger J, Liberman M, Lima Silva V, Nguyen M, Thomas S, Le A, et al. EOMES Interacts With RUNX3 and BRG1 to Promote Innate Memory Cell Formation Through Epigenetic Reprogramming. *Nat Commun* (2019) 10:3306. doi: 10.1038/s41467-019-11233-6

59. Cruz-Guilloty F, Pipkin ME, Djuretic IM, Levanon D, Lotem J, Lichtenheld MG, et al. Runx3 and T-Box Proteins Cooperate to Establish the Transcriptional Program of Effector CTLs. *J Exp Med* (2009) 206:51–9. doi: 10.1084/jem.20081242

60. Luette-Eversloh M, Cicek BB, Siracusa F, Thom JT, Hamann A, et al. Luetke-Eversloh M, Cicek BB, Siracusa F, Thom JT, Hamann A, et al. Control of Effector CD8+ T Cell Function by the Transcription Factor Eomesodermin. *Mol Biol Rep* (2003) 30:2929–36. doi: 10.1007/s11033-003-1164-x

61. Hasenkamp J, Borgerding A, Wufl G, Uhrberg M, Jung W, Dingeldein S, et al. Resistance Against Natural Killer Cell Cytotoxicity: Analysis of Mechanisms. *Scand J Immunol* (2016) 64:444–9. doi: 10.1111/1365-3083.2016.01803.x

62. Jardine L, Hambleton S, Bigley V, Pagan S, Wang XN, Collin M. Sensitizing Primary Acute Lymphoblastic Leukemia to Natural Killer Cell Recognition by Induction of NKG2D Ligands. *Leukemia Lymphoma* (2013) 54:167–73. doi: 10.3109/10428194.2012.708026

63. Oei VYS, Siernicka M, Graczyk-Jarzynka A, Hoel HJ, Yang W, Palacios D, et al. Ex Vivo Generated Natural Killer Cells Acquire Typical Natural Killer Receptors and Display a Cytotoxic Gene Expression Profile Similar to Peripheral Blood Natural Killer Cells. *Stem Cells Dev* (2012) 21:2926–34. doi: 10.1089/scd.2011.0659

64. Dolstra H, Boeven MWH, Spanholz J, Hangalapura BN, Tordoir M, Maas F, et al. Successful Transfer of Umbilical Cord Blood CD34+ Cells for Adoptive Cancer Immunotherapy. *PloS One* (2010) 5:e9221. doi: 10.1371/journal.pone.0009221

65. Lehmann D, Spanholz J, Osl M, Tordoir M, Lipnik K, Bilban M, et al. Ex Vivo Generated Natural Killer Cells Acquire Typical Natural Killer Receptors and Display a Cytotoxic Gene Expression Profile Similar to Peripheral Blood Natural Killer Cells. *Stem Cells Dev* (2012) 21:2926–34. doi: 10.1089/scd.2011.0659

66. Dolstra H, Boeven MWH, Spanholz J, Hangalapura BN, Tordoir M, Maas F, et al. Successful Transfer of Umbilical Cord Blood CD34+ Cells for Adoptive Cancer Immunotherapy. *PloS One* (2010) 5:e9221. doi: 10.1371/journal.pone.0009221

67. Carlsten M, Levy E, Karambelkar A, Li L, Rege R, Berg M, et al. Efficient mRNA-Based Genetic Engineering of Human NK Cells With High-Affinity CD16 and CCRe Augments Rituximab-Induced ADCC Against Lymphoma and Targets NK Cell Migration Toward the Lymph Node-Associated Chemokine Ccl19. *Front Immunol* (2016) 7:105. doi: 10.3389/fimmu.2016.00105

68. Ganjoo KN, de Vos S, Pohlman BL, Flinn IW, Forero-Torres A, Enas NH, et al. Phase 1/2 Study of Ocaratuzumab, an Fc-Engineered Humanized Antibody Directed Against CD20 Monoclonal Antibody, in Low-Affinity FcγRIIIa Patients With Previously Treated Follicular Lymphoma. *Leukemia Lymphoma* (2015) 56:42–8. doi: 10.3109/10428194.2014.911859

69. Cartron G, de Guibert S, Dilhudy MS, Morschhauser F, Leblond V, Dupuis J, et al. Obinutuzumab (GA101) in Relapsed/Refractory Chronic Lymphocytic Leukemia: Final Data From the Phase 1/2 GAUGUIN Study. *Blood* (2014) 124:2196–202. doi: 10.1182/blood-2014-07-586610

70. Deng X, Terumuma H, Nieda M, Xiao W, Nicol A. Synergistic Cytotoxicity of Ex Vivo Expanded Natural Killer Cells in Combination With Monoclonal Antibody Drugs Against Cancer Cells. *Int Immunopharmacol* (2012) 12:4593–605. doi: 10.1016/j.intimp.2012.09.014

71. Suen WC, Lee WY, Leung KT, Pan XH, Li G. Natural Killer Cell-Based Cancer Immunotherapy: A Review on 10 Years Completed Clinical Trials. *Cancer Invest* (2018) 36:431–57. doi: 10.1080/07357907.2018.1515315

72. Vallera DA, Felices M, McElmurry R, McCullar V, Zhou X, Schmohl JU, et al. IL15 Trispecific Killer Engagers (TriKE) Make Natural Killer Cells Specific to CD3+ Targets While Also Inducing Persistence. *In Vivo* Expansion, and Enhanced Function. *Clin Cancer Res an Off J Am Assoc Cancer Res* (2016) 22:3440–50. doi: 10.1158/1078-0432.CCR-15-2710

73. Vivier E, Tomasello E, Baratin M, Walzer T, Ugolini S. Functions of Natural Killer Cells. *Nat Immunol* (2008) 9:503–10. doi: 10.1038/nri2582

74. Pearce EL, Mullen AC, Martins GA, Kraszyck CM, Hutchins AS, Zediak VP, et al. Control of Effector CD8+ T Cell Function by the Transcription Factor Eomesoderm. *Science* (2003) 302:1041–3. doi: 10.1126/science.1090148

75. Soderquest K, Hertweck A, Giambartolomei C, Henderson S, Mohamed R, Goldberg R, et al. Genetic Variants Alter T-Bet Binding and Gene Expression in Mucosal Inflammatory Disease. *PloS Genet* (2017) 13:e1006587. doi: 10.1371/journal.pgen.1006587

76. Ai J, Wang Y, Tan K, Deng Y, Luo N, Yuan W, et al. A Human Homolog of Mouse Lhb Gene, hLHB, Expresses in Heart and Activates SRE and AP-1 Mediated MAPK Signaling Pathway. *Mol Biol Rep* (2008) 35:179–87. doi: 10.1007/s11033-007-9068-4

77. Montes-Casado M, Ojeda G, Aragoneses-Fenoll I, Lopez D, de Andres B, Gaspar ML, et al. ICOS Deficiency Hampers the Homeostasis, Development and Function of NK Cells. *PloS One* (2019) 14(7):e0219449. doi: 10.1371/journal.pone.0219449

Conflict of Interest: An international patent application WO2020070070 was filed by Ghent University (Ghent, Belgium) on 30/09/2019, with title ‘Accelerated human hematopoietic stem cell differentiation towards mature natural killer cells with enhanced antibody-dependent cytotoxic activity’, with LK and GL as inventors.

The remaining authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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