Lymphoma

The novel lncRNA BlackMamba controls the neoplastic phenotype of ALK− anaplastic large cell lymphoma by regulating the DNA helicase HELLS

Valentina Fragliasso1 · Akanksha Verma2,3 · Gloria Manzotti1 · Annalisa Tameni1,4 · Rohan Bareja2 · Tayla B. Heavican5 · Javeed Iqbal5 · Rui Wang6 · Danilo Fiore6 · Valentina Mularoni1 · Wing C. Chan7 · Priscilla Lhoumaud8 · Jane Skok8 · Eleonora Zanetti9 · Francesco Merli10 · Alessia Ciarrocchi1 · Oliver Elemento2 · Giorgio Inghirami6

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Abstract

The molecular mechanisms leading to the transformation of anaplastic lymphoma kinase negative (ALK−) anaplastic large cell lymphoma (ALCL) have been only in part elucidated. To identify new culprits which promote and drive ALCL, we performed a total transcriptome sequencing and discovered 1208 previously unknown intergenic long noncoding RNAs (lncRNAs), including 18 lncRNAs preferentially expressed in ALCL. We selected an unknown lncRNA, BlackMamba, with an ALK− ALCL preferential expression, for molecular and functional studies. BlackMamba is a chromatin-associated lncRNA regulated by STAT3 via a canonical transcriptional signaling pathway. Knockdown experiments demonstrated that BlackMamba contributes to the pathogenesis of ALCL regulating cell growth and cell morphology. Mechanistically, BlackMamba interacts with the DNA helicase HELLS controlling its recruitment to the promoter regions of cell-architecture-related genes, fostering their expression. Collectively, these findings provide evidence of a previously unknown tumorigenic role of STAT3 via a lncRNA-DNA helicase axis and reveal an undiscovered role for lncRNA in the maintenance of the neoplastic phenotype of ALK−ALCL.

Introduction

Peripheral T-cell lymphomas (PTCLs) are a heterogeneous group of tumors, which include 29 different subtypes according to the World Health Organization (WHO). Among them, systemic CD30-positive anaplastic large cell
lymphomas (ALCL) are characterized by distinct histopathological and clinical features. ALCL are stratified based on the presence of anaplastic lymphoma kinase (ALK) translocations [1–4]. Clinically, systemic ALK−ALCL displays a poor response to therapy and inferior survival compared with ALK+ALCL [4–9]. Within ALK−ALCL,
Fig. 1 ALCL samples expressed a restricted set of aberrantly activated previously unknown lncRNAs. a Schematic representation of human samples used to perform directional RNA sequencing. b Bioinformatic pipeline for the discovery of previously unknown lncRNAs. c Density plot for transcript length shows shared pattern between previously unknown and known lncRNAs compared with protein-coding genes, which are much longer. d Coding Potential Score obtained from GENEID shows that previously unknown lncRNAs and known lncRNAs have comparable and lower average coding potential than do protein-coding genes. e Comparing previously unknown lncRNAs against all transcripts with at least two or more exons show a greater number of exons for the protein-coding genes. f Unsupervised analysis of previously unknown lncRNAs profile across normal T-cell lymphocytes and ALCL primary samples. g Flowchart for the discovery of BlackMamba in ALK− ALCL samples. h Schematic representation of locus, structure, and aligned reads of BlackMamba. Numbers represent the length (bp) of exons and introns (gray). Sashimi plots of representative ALK− and AL+ ALCL samples were generated by Integrative Genomics Viewer software. The genomic coordinates are measured along the horizontal axis and the RPKM (Reads Per Kilobase per Million mapped reads) values up the vertical axis. qRT-PCR analysis of BlackMamba in a validation set of ALCLs samples (i) and in a panel of cell lines (j).

Distinct subsets have been described [7]. ALK− ALCL share unique transcriptional signatures [6, 10] and can be further classified using recurrent genomic abnormalities including JAK1/STAT3 mutations, DUSP22 or TP63 translocations, loss of TP53 and PRDM1/BLIMP1, and aberrant expression of ERBB4 [3, 4, 11]. However, the molecular mechanisms leading to ALK− ALCL transformation, maintenance, and immune evasion remain quite elusive. Hyperactivation of STAT3 is documented in about 40% of ALK− ALCL [3], and contributes to the tumorigenesis and maintenance of STAT3+ ALCL via canonical gene regulation [3, 12] a paradigm shared by many human cancers. However, new findings suggest that STAT3 can also contribute to cancer tumorigenesis by regulating long noncoding RNA (lncRNA) [13, 14].

LncRNAs are transcripts, longer than 200 nucleotides, often display an intron–exon organization, and share close similarities to protein-coding genes. They have pleotropic properties, controlling gene expression, protein stability, localization and function, and cell identity [15]. Unbiased genome-wide analyses discovered thousands of lncRNAs, whose number outweighs those of protein-coding RNAs. More than 8000 lncRNAs are aberrantly expressed in cancer, making these genes ideal tumor-specific biomarkers and putative targets for therapeutic interventions [16].

Remarkably, lncRNAs can shape the T-cell compartment and modulate the adaptive immune system [17]. Detectable lncRNAs are often located within the neighborhood of lineage-specific mRNA [17], and their dynamic expression is cell/stage-specific regulated [18] and linked to cell differentiation and identity [19]. Indeed, IFNG-AS1, linc-Ccr2–59AS, Th2 LCRR, and GATA3-AS1 are associated with distinct T-helper 1 (Th1) or Th2 phenotypes [20–22] while lincRNA-MAF-4 promotes T-lymphocyte differentiation by interacting with chromatin modifiers LSD1 and EZH2 via MAF [19]. Conversely, their oncogenic contributions in T-cell lymphoma remain poorly elucidated and their contribution to the ALCL phenotype remains unexplored [23–25].

Here, by performing deep expression profiling in conjunction with de novo transcriptome assembly, we discovered a panel of previously unknown lncRNAs of ALCL. We focused on a chromatin-associated lncRNA, selectively expressed by ALK− ALCL lymphoma, named BlackMamba. Mechanistically, BlackMamba is regulated via STAT3 and its expression is required to sustain proliferation and clonogenicity of ALK− ALCL through the transcriptional regulation and the functional control of the lymphoid helicase HELLS. These findings provide new evidence on the mechanisms leading to STAT3-mediated ALCL transformation and foster the implementation of STAT3 target therapies for these lymphomas.

Materials and methods

Tissue samples

Fresh and viable cryopreserved cells were isolated from diagnostic/relapsed primary lymphoma biopsies. Diagnoses were assigned according to the WHO classification. Tissues used for NGS analyses were selected for their high tumor cell content (>50%). All studies were approved through institutional human ethics review boards, and patients provided written informed consent in accordance with the Declaration of Helsinki.

Cell growth, colony formation assays, and cell division

For cell growth assays, cells were washed with phosphate-buffered saline seeded at 2.5 × 10^5 cells/ml and treated with drugs. Viable cells were counted by trypan blue exclusion.

Colony formation assays were performed as previously described [26]. Cell division was evaluated using 5 μM of carboxyfluorescein succinimidyl ester (CFSE) fluorescent dye following the manufacturer’s instructions (ab113853, Abcam). The fluorescence was read by FACSCanto II instrument after 6 or 9 days and data were analyzed using BD FACSDiva Software (BD).

Analysis of mRNA stability

Actinomycin D was used to inhibit nascent RNA synthesis. MAC2A, TLBR-1, and TLBR-2 cells (5 × 10^5 cells/ml)
Fig. 2 *BlackMamba* is a previously unknown long noncoding RNA in ALK− ALCL subset. a Schematic representation of *BlackMamba* genomic locus showing the position and the level of histone modifications of the putative promoter and the representation of TSS region. Luciferase activity of overlapping fragments into pGL3 vector (b) and luciferase activity of P4 fragment into pGL3-promoter vector (c). Data are represented as normalized ratio of firefly-Renilla luciferase activities and are expressed as mean values ± SD (n = 3). **p ≤ 0.01. ChIP quantitative qRT-PCR detection of H3K27ac (d), H3K4me3 (e), and phospho- S5-RNA-polymerase II (f) on P4 fragment in a panel of cell lines. GAPDH promoter was used as CTR whereas, BlackMamba promoter +33,459 bp (*BlackMamba*) and a non-coding intergenic region (CTR−) served as negative controls. The values represent mean ± SEM (n = 3). **p ≤ 0.01.

were treated with actinomycin D or DMSO (5 μg/ml, Sigma-Aldrich) for 2, 4, 8, 16, and 24 h and collected for RNA isolation.

**siRNA transfection**

FEPD, MAC2A, TLBR-1, and TLBR-2 cells (1 × 10⁶) were transfected with 5 or 10 pmol siRNA using the Cell Line Nucleofector Kit SF and Amanax 4D Nucleofector (program FI115 for FEPD and MAC2A, DS-130 for TLBR-2 and for TLBR-1). Twenty-four hours after transfection, cells were harvested and plated 2.5 × 10⁵ cells/ml. For STAT3, we used a Silencer Select Validated siRNA ID:s743 (Ambion, Life Technologies); for *BlackMamba* we used three different Silencer Selected siRNAs (Ambion, Life Technologies) (Supplementary Table 1).

**Luciferase assay**

MAC2A and TLBR-2 cells (1 × 10⁶) were transfected with 0.4 μg of reporter pGL3-luciferase plasmids using the Cell Line Nucleofector Kit SF and 4D Amanax Nucleofector (program FI115 for MAC2A, DS-130 for TLBR-2).

Twenty-four hours after transfection, cells were harvested and luciferase activity was measured using the Dual-Luciferase Reporter Assay System (Promega) in a GloMax Discover Luminometer (Promega) according to the manufacturer’s instructions. For each sample, firefly luciferase activity was normalized on Renilla luciferase activity and transactivation of the various reporter constructs was expressed as fold induction on empty vector (pGL3-basic or pGL3-promoter) activity.

**Chromatin immunoprecipitation (ChIP)**

ChIP experiments were performed as previously described [27]. Chromatin was precipitated with antibodies against H3K4me3 (Abcam-ab8580), H3K4me1 (Abcam-ab8895), H3K27Ac (Abcam-ab4729), RNA polymerase II STD repeat YSPTSPS ((phospho S5) (Abcam – ab5408), STAT3 (124H6 Mouse mAb#9139), HELLS (H-4, sc-46665 Santa Cruz), or IgG (as negative control). For each experiment, a chromatin amount corresponding to 0.5% of chromatin used for immunoprecipitation was kept as input control. Each qRT-PCR value was normalized over the appropriate input control and reported in graphs as % of input (qRT-PCR value/input value × 100).

**RNA immunoprecipitation (RIP)**

RIP was performed as described by Abcam RIP protocol. The precleared lysate was incubated for 2 h with 6 μg of antibodies specific for HELLS (sc-46665, Santa Cruz Biotechnology, Inc.) or with IgG as negative control. All experiments were repeated at least three times.

**Quantitative PCR (qRT-PCR)**

One microgram of total RNA was reverse transcribed using RT (iScript, Biorad). The amplified transcript level of each specific gene was normalized on CHMP2A housekeeping. ΔΔCt quantification method was used for RT-qPCR analyses.

The list of primers used is provided in Supplementary Table 2.

**Western blot**

Western blot analysis was performed using standard techniques [3]. The primary antibodies were: STAT3 (124H6 Mouse mAb#9139, 1:1000 Cell signaling Technology), pSTAT3 (Tyr705, D3A7 Rabbit mAb#9145, 1:1000 Cell signaling Technology), STAT5 (D206Y Rabbit mAb#94205, 1:1000 Cell signaling Technology) pSTAT5 (Tyr694, Rabbit mAb#9351, 1:1000 Cell signaling Technology), ALK (D5F3 Rabbit mAb#83633, 1:1000 Cell signaling Technology), pALK (Tyr1604, Rabbit mAb#3341, 1:1000 Cell signaling Technology), β-Actin (sc-47778, 1:1000 Santa Cruz Biotechnology, Inc.), and HELLS (sc-46665, 1:1000 Santa Cruz Biotechnology, Inc.). All secondary antibodies (rabbit and mouse) were HRP-conjugated (GE Healthcare) and diluted 1:5000.

**Immunofluorescence**

Immunofluorescences were performed as previously described [28].

**Statistical analyses**

Statistical analyses for identification and the analysis of lncRNAs are described in Supplementary methods.
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conserved spliced transcripts (Fig. 1c, d), and they binding, whereas GAPDH promoter was used as positive control for control. ATF3 promoter was used as positive control for STAT3

Fig. 3 STAT3 regulates BlackMamba expression. a STAT3 ChIP-sequencing analysis (top panel) and ATAC-sequencing analysis (bottom panel) along BlackMamba locus in ALK+ ALCL cell lines and PDTX (Belli ALK− ALCL and IL89 BIA-ALCL). b qRT-PCR analysis of BlackMamba expression after 24 h of treatment with 1 μM of ruxolitinib or 2 μM of INCB039110. The values represent mean ± SD (n = 3). *p ≤ 0.05; **p ≤ 0.01. ChIP qRT-PCR detection of: (c) STAT3 antibody on P1 fragment, (d) Phospho-5 s RNA Polymerase II, and (e) anti-H3K27ac antibodies on P4 fragment of BlackMamba in ALK− ALCL cell lines treated for 24 h with 1 μM of ruxolitinib. BlackMamba region +33,459 bp (BlackMamba) was used as a negative control. ATF3 promoter was used as positive control for STAT3 binding, whereas GAPDH promoter was used as positive control for d, e. The values represent mean ± SEM (n = 3). **p ≤ 0.01; *p ≤ 0.05. f qRT-PCR analysis of BlackMamba in two different regions (+438 bp and +33,459 bp) in ALK+ ALCL cells after transfection with specific STAT3-siRNA (96 hours). CD30 was used as positive control. The values represent mean ± SD (n = 3); *p ≤ 0.05; **p ≤ 0.01.

Statistical analyses were performed using GraphPad Prism Software (GraphPad). Statistical significance was determined using the Student’s t test. Each experiment was replicated multiple times (>3 up to 6).

Results

ALKL express a large pool of previously unknown IncRNAs

To define IncRNAs preferentially associated with ALCL, we performed high coverage and directional RNA sequencing (RNA-Seq) of 21 ALK+ ALCL and 16 ALK− ALCL primary samples. We included normal T-lymphocytes, corresponding to different stages of differentiation and 10 ALCL cell lines (Fig. 1a and Supplementary Table 3). Firstly, to confirm the appropriateness of pathological samples within our discovery set, we used a 3-gene model classifier (TNFRSF8, BATF3, TMOD1), proven to accurately define ALK+ and ALK− ALCL (Supplementary Fig. 1A) [6]. A principal component analysis based on canonical coding gene expression further resolved the discovery cohort into distinct clusters corresponding to normal T-cell and ALCLs (Supplementary Fig. 1B); the latter group was further stratified into two distinct subgroups, largely represented by ALK+ and ALK− ALCL samples (Supplementary Fig. 1C).

Next, we executed a de novo transcriptome analysis of the aligned primary tumor samples [29], which identified 106,014 new transcripts. Applying filtering cutoffs based on transcript length, exon count, and coding potential (based on cross-species comparisons), we discovered 1208 previously unknown ALCL-specific IncRNAs (Fig. 1b). These IncRNAs showed transcript length >200 bp (a canonical IncRNAs feature), a stringent number of exons n = 2 for conserved spliced transcripts (Fig. 1c, d), and they displayed low coding potential compared to canonical coding RNAs (Fig. 1e).

Next, we identified a set of IncRNAs significantly overexpressed in ALCL samples (18/352) by unsupervised analysis (Supplementary Table 4). Even if this set was not able to fully discriminate ALK+ and ALK− ALCL, these novel IncRNAs were found to be significantly overexpressed in ALCL compared with normal T-lymphocytes (Fig. 1f and Supplementary Fig. 1D, E). Then we restricted the analysis to IncRNAs expressed in at least 20% of ALCL samples (five or more samples), this analysis led to the discovery of a pool of 14 IncRNAs (FDR < 0.05) (Fig. 1g). Since the deregulated STAT3 signaling is oncogenic in ALCL [3, 12], we then searched for canonical STAT3 binding sites within putative promoter regions of these previously unknown IncRNAs (Fig. 1g). We identified the IncRNA XLOC_043524 only, which we named BlackMamba.

BlackMamba is a nonannotated IncRNA located on q26.3 of chromosome 10, predicted to be transcribed from the minus strand, with an estimated transcript length of 70,292 bp (Fig. 1i) and with half-life of ~4 h (Supplementary Fig. 2a-d). BlackMamba is composed of three exons, lacking alternative isoforms. Within the discovery cohort, 9/16 ALK− ALCL (56%) and 1/21 ALK+ ALCL (4.7%) expressed detectable levels of BlackMamba. The preferential expression of ALK− ALCL was further confirmed in an independent set of 15 ALCLs and 9 PTCLs, and no detectable transcripts were seen in a cross-validation cohort of healthy donor resting and activated PBMCs (Fig. 1i and Supplementary Table 5). When we extended this analysis to a panel of T-cell lines, we confirmed that BlackMamba expression was consistently detectable in ALK− ALCL and breast implanted associated (BIA)-ALCL cell lines, albeit with variable levels of expression, with TLBR-1 and TLBR-2 expressing the highest levels. Very low transcripts were detected in ALK+ ALCL (L82 and Karpas299), systemic and cutaneous ALK-ALCL (FPED, Mac1), and in cutaneous T-cell lymphoma MJ, while T-ALL lines (CUTLL1, KOPT-K1, and Jurkat), and mycosis fungoides HUTL-78 cells were negative (Fig. 1j).

BlackMamba is a promoter-associated IncRNA transcriptionally regulated by STAT3

To explore the mechanism(s) which regulates the expression of BlackMamba, we investigated the genomic elements responsible for its transcription. H3K4me3 profile by ChIP-seq in ALK− ALCL cell lines and patient-derived tumor xenograft (PDTX) (Belli) lines showed a high-density profile within a 2000 bp region, spanning the putative transcription start site (TSS) (Fig. 2a). These data are in agreement with a relatively high level of H3K4Me3 (but not
of H3K4Me1) within the same region in ENCODE (in several cell lines), suggesting that BlackMamba is likely to be a promoter-associated gene rather than an enhancer-associated RNA. To elucidate the promoter region of BlackMamba, we then cloned multiple DNA fragments upstream of a luciferase reporter cassette corresponding to a
region of 2000 bp (P1-P5), spanning from −811 to +1076 bp of the BlackMamba TSS (Fig. 2a). High luciferase signals were observed with the segment spanning from −459 bp to −90 bp (P4) in both MAC2A and TLBR-2 (Fig. 2b). By cloning P4 in an “enhancer-like” position downstream to the reporter gene, we demonstrated that this segment did not act as an enhancer (Fig. 2c). The histograms show the fluorescent intensity of CFSE-labeled TLBR-2 and MAC2A expressing pLKO-shRNA BlackMamba (9 days of treatment). Each data point represents the mean ± S.D. (n = 3). One-tailed t-test. *p ≤ 0.05; **p ≤ 0.01. d The histograms show the fluorescent intensity of CFSE-labeled TLBR-2 and MAC2A expressing pLKO-shRNA BlackMamba (9 days of treatment). Each data point represents the mean ± S.D. (n = 3). One-tailed t-test. *p ≤ 0.05; **p ≤ 0.01. e Histogram shows methylcellulose colony formation of pLKO-shRNAs BlackMamba expressing cells pretreated with doxycycline for 6 days. Colonies were counted on day 10, after plating. Each data point represents the mean ± S.D. (n = 3). f The upper panel shows the percentage of polynucleated cells in at least 100 cells stained with May–Grunwald Giemsa (6 days after doxycycline induction). Each data point represents the mean ± S.D. (n = 3). *p ≤ 0.05; **p ≤ 0.01. The lower panel shows the polynucleated cells in TLBR-2. The scale bar represents 100 μm. g FISH analysis of chromosome 17 centromere in pLKO-shRNAs BlackMamba expressing TLBR-2 and MAC2A cells (6 days of doxycycline induction). The scale bar represents 10 μm.

BlackMamba is required for active proliferation and clonogenicity of ALK− ALCL cells

To test the biological properties of BlackMamba, we silenced its expression using two different approaches: a transient siRNA transfection and a doxycycline inducible shRNA. Because BlackMamba is a large gene (~70 kb), we targeted different regions (Supplementary Fig. 4A and Supplementary Table 2). We found two independent shRNAs that effectively reduced its expression after doxycycline induction in MAC2A (~50% for each shRNAs) without affecting top-scoring off-targets significantly (Supplementary Fig. 4B). Both shRNA#2 and #6 could effectively knockdown (KD) the lncRNA in TLBR-2, although with different potency (~30%, shRNA#2 and ~60%, shRNA#6) (Fig. 4a). Next we studied the cellular localization of BlackMamba at steady state, demonstrating it was enriched in the nucleus and strongly associated to the chromatin fraction, suggesting a putative role in chromatin organization and gene expression regulation (Fig. 4b and Supplementary Fig. 4C).

Functionally, the KD of BlackMamba resulted in a dose-dependent cell growth inhibition (Fig. 4c and Supplementary Fig. 4D, E), in the absence of an increased rate of apoptosis (Supplementary Fig. 4F). This phenotype was reproducibly detected in shRNA#6-treated ALCL cells, although significant shRNA#2 mediated changes were observed only in MAC2A cells, which expressed lower levels of mRNA (Supplementary Fig. 4G). Next we demonstrated a delayed cell division in BlackMamba KD cells (Fig. 4d and Supplementary Fig. 4H, I), and an impaired ALCL lymphoma colony formation (Fig. 4e). Conversely, the growth and survival in control K562 (chronic myeloid leukemia) and CUTLL1 KD cell lines...
were not affected (Supplementary Fig. 4J). Interestingly, the cytological inspection of May–Grunwald Giemsa stained ALK−ALCL cells upon BlackMamba KD showed an increased number of polynucleated cells (Fig. 4f and Supplementary Fig. 4K), with a polyploid DNA content by partial FISH-based karyotyping (Fig. 4g).
Lastly, since cytokinesis requires an appropriate cytoskeleton organization, we investigated the cytoskeleton architecture of BlackMamba silenced cells. As actin filaments are major components of the contractile structure that guide cytokinesis in eukaryotes [30], we examined the cellular cytoskeleton using phalloidin staining of actin filaments. We found that after BlackMamba KD, cells lose actin polarization and have a displacement of filaments from their membrane localization, supporting its role in cytoskeleton reorganization and cytokinesis (Fig. 5a). This hypothesis is supported by the transcriptional changes observed after RNA sequencing in BlackMamba KD (59 downregulated genes and 61 significantly upregulated genes, ≥2-fold and FDR < 0.05, Fig. 5b and Supplementary Table 6) where top-scored genes modulated (i.e., RGS1, CCL22, CCL17, PAK2, RHOU) epitomize by the modulation of actin cytoskeleton, integrin-mediated cell adhesion, and focal adhesion genes (Fig. 5c, d and Supplementary Fig. 5A).

Overall these data suggest that BlackMamba is involved in the maintenance of appropriate completion of cytokinesis.

**BlackMamba regulates the transcription of the lymphoid-specific DNA helicase HELLs**

Chromatin-enriched lncRNAs are spatially correlated with transcription factors [31], can act as cell type-specific activators of proximal gene transcription [32] and chromatin-associated lncRNAs (such as XIST or KCNQ1ot1) can influence local chromatin organization, leading to in cis transcriptional repression of genes within large genomic regions [33].

To test whether BlackMamba could operate according to this model, we correlated its expression with transcriptional factors and chromatin-remodeling genes (16 ALK− ALCL and 21 ALK+ ALCL samples). The lymphoid-specific helicase (LSH) HELLs, the SET domain containing protein PRDM13, and the polycomb repressive complex 2 histone-lysine N-methyltransferase EZH2 were found to be positively correlated with BlackMamba. Conversely, the homeobox protein HHEX showed a negative correlation (Fig. 5e). To test whether this association was directly linked to BlackMamba expression, we quantified the mRNA levels of HELLs and HHEX in inducible shRNA BlackMamba ALCL cells. HELLS expression was consistently downregulated upon BlackMamba silencing, while HHEX was upregulated (Fig. 5f and Supplementary Fig. 5B). No consistent changes were observed for EZH2 (Supplementary Fig. 5C) while PRDM13 mRNA was undetectable (data not shown). In line with the gene expression changes, H3K4me3 and H3K27me3 underwent chromatin reorganization, with a significant reduction of H3K4me3 binding on HELLS promoter and a parallel loss of H3K27Me3 on HHEX promoter after silencing (Fig. 5g).

Being also HHEX and HELLS the only TFs located on the same chromosome of BlackMamba, we hypothesized that BlackMamba could regulate gene expression in cis.

To enforce this concept, we quantified the mRNA levels of several neighboring genes spanning 1 Mb from BlackMamba locus. After BlackMamba KD, 5/6 genes were concordantly downregulated in MAC2A and TLBR-2 cell lines (Supplementary Fig. 5D).

These data support the model which predicts that BlackMamba regulates in cis the expression of genes located on the same chromosome.

**BlackMamba interacts with HELLS to control the BlackMamba-dependent transcriptional program**

HELLS, also known as LSH or proliferation-associated SNF2-like (PASG), belongs to a large family of SNF2 chromatin-remodeling ATPases. HELLs is critical to the normal development and survival of lymphoid cells [34] and regulates chromatin organization and gene expression [35, 36]. HELLS mutations or misregulated expression were seen in several cancers and some cases of ICF syndrome [37].

To test whether HELLS enforces BlackMamba-mediated transcription, we silenced HELLS by specific shRNAs. qRT-PCR and WB analyses confirmed an effective silencing (Fig. 6a, b and Supplementary Fig. 6A), demonstrating the concomitant downregulation of known target BMI-1 and MLL1 genes [38, 39] (Supplementary Fig. 6B). Interestingly, the loss of HELLS was linked to expression changes of RGS1, CCL17, CCL22, PAK2, and KCNMA1, phenocopying the BlackMamba KD phenotype (Fig. 6c).
Conversely, the ectopic expression of HELLS effectively restored the baseline expression of these genes in Black-Mamba KD cells (Fig. 6d, e).

It is known that lncRNAs control gene expression by recruiting chromatin-remodeling complexes to target promoters or enhancers, thereby influencing histone
The loss of HELLS led to an impaired cell growth (Fig. 7a). Gating the cell growth capacity of HELLS KD cells. Indeed, RIP precipitation [3] and showed that HELLS binds to two distinct regions of BlackMamba at the 3′′-end of the lncRNA. Since the chromatin-remodeling properties of HELLS can be mediated by the lncRNA HOTAIR [41], we reasoned that HELLS might interact with HOTAIR to mediate its recruitment to target genes. To prove a direct association between HELLS and BlackMamba, we used RNA immunoprecipitation [3] and showed that HELLS binds to two distinct regions of BlackMamba at the 3′′-end of the lncRNA (Fig. 6f). No readout was seen in HELLS KD cells. Next, we proved that HELLS was preferentially bound to target gene promoters only in BlackMamba-positive ALK− ALCLs independently from HELLS basal expression level (Fig. 6g and Supplementary Fig. 6C).

Collectively, these data demonstrate that BlackMamba-HELLS could be a part of the regulatory complex that occupied loci to coordinate ALK− ALCL transcriptional program.

**HELLS is required for the maintenance of the neoplastic phenotype of ALK− ALCL**

To determine the contribution of HELLS and BlackMamba in the maintenance of ALCL phenotype, we first investigated the cell growth capacity of HELLS KD cells. Indeed, the loss of HELLS led to an impaired cell growth (Fig. 7a and Supplementary Fig. 6D), reduced duplication rate (Fig. 7b), and clonogenicity (Fig. 7c and Supplementary Fig. 6E), a phenotype associated with an increased number of polynucleated cells, phenocopying the BlackMamba KD cells (Fig. 7d and Supplementary Fig. 6F).

Next we proved that the overexpression of HELLS could counteract the phenotype associated with the KD of BlackMamba, as the growth impairment of TLBR-2 cells expressing inducible shRNA against BlackMamba was effectively rescued (Fig. 7e) a finding associated with the mitigation of the polynucleated phenotype of the BlackMamba KD (Fig. 7f).

Overall, these findings demonstrate that HELLS is an essential downstream mediator of BlackMamba and that BlackMamba-HELLS axis represents a vulnerability of ALCL cells.

**Discussion**

Although a more complete genomic annotation of ALK− ALCL is emerging [42, 43], the mechanistic modalities of action remain elusive even for known recurrent defects [4, 42].

Here we describe a new chromatin-associated lncRNA, named BlackMamba, preferentially expressed in ALK− ALCL. Mechanistically, STAT3 regulates the expression of BlackMamba and its expression is required for ALK− ALCL neoplastic phenotype. This is achieved mainly through the action of the LSH, HELLS (Fig. 7g) and the transcriptional regulation of genes controlling G-protein and cytoskeletal organization, cell migration, tissue recruitment, and inflammation.

While emerging evidence have shown that lncRNAs are pathogenetic in human B-cell hematological neoplasms [44], little is known in mature T-cell neoplasms [23–25, 45]. In this study, we identified 1208 previously unknown lncRNAs linked to normal or neoplastic T-cells and among them, 18 lncRNAs were largely restricted to ALCL. Accordingly, a third of these new lncRNAs were co-shared by both ALCL subtypes, supporting the notion that both ALK+ and ALK− ALCL can share some communalties [46].

Deregulated activation of STAT3 is a hallmark of many human cancers [47]. ALK+ ALCL and a subset of ALK− ALCL have been proven to be addicted to the JAK/STAT signaling pathway [3, 48] and BIA-ALCL display a constitutive JAK/STAT deregulation [49]. Here, BIA-ALCL cell lines, as well as PDTX, were found to express high levels of BlackMamba. In ALCL, the deregulation/activation of JAK/STAT pathway is mediated by gene fusions, somatic mutations [50], and loss of negative regulators [7, 51]. These lead to a distinct transcriptional program associated with defined pathological entities [6, 46]. LncRNAs have been also shown to be linked to STAT signaling by modulating metabolic pathways [52] and conversely, STAT3, controlling the expression of lncRNAs, can regulate cell differentiation [17].

BlackMamba is a target of STAT3. Both siRNA-mediated and pharmacological inhibitions of JAK/STAT3 pathway profoundly repressed BlackMamba expression demonstrating its contribution in cell growth and clonal expansion of ALCL. Interestingly, loss of the BlackMamba
expression was linked to a unique phenotype characterized by an increased number of polynucleated cells with a balanced chromosomal enumeration possibly linked to key genes regulating cytoskeleton and cell motility. As STAT3 can regulate cell migration via RAC1 and Rho [53, 54], our data provide a new layer of complexity demonstrating a...
new role for STAT3 in cell shape and cytoplasmic partition via the axis of BlackMamba-HELLS. This model is supported by phenotype seen in HELLS KD cells, arguing that HELLS represents a critical downstream effector of BlackMamba.

Mechanistically, BlackMamba also contributes to the neoplastic phenotype of ALCL via multiple mediators, modulating a large group of genes encoding chemokines and chemokine receptors (CCL17–22 and CXCL10), G-protein regulators (RGS1) and multiple proteins involved in cell growth and motility (PAK2, ERBB2, RHOU, KIF21b).

Among BlackMamba targets, we found HELLS (16,550,000 bp from the locus of BlackMamba). HELLS controls T-cell growth [34], regulates DNA methylation [55], and modulates the epigenetic states at specific enhancers of key cell cycle regulators [56]. In cancer cells, HELLS sustains glioma stemness and through the interaction with E2F3 controls cell proliferation of prostate cancer cells [38, 57]. By interacting with the epigenetic silencer factor G9a, HELLS represses gene transcription [58]. Remarkably, a recent genome-scale CRISPR-Cas9 screen has shown cancer dependencies to DNA helicase and identified ATP-DNA helicases as promising new synthetic lethal targets in tumors [59]. Our work extends these findings providing a new lncRNA-dependent mechanism controlling the recruitment of HELLS on chromatin sites and its expression in lymphomas.

Our data provide novel insights into the transformation of ALCL via the untapped role of a lncRNA. Collectively, the findings further support the design of target therapeutic strategies to pharmacologically ablate/inhibit the expression of STAT3 and encourages novel discovery programs for the selection of compounds which could impair STAT3 downstream effector elements like HELLS. Lastly, since HELLS is expressed in many human tumors and plays a relevant role in DNA repair and genomic stability of cancers, its pharmacological inhibition represents a viable therapeutic strategy in many human cancers.

**Data availability**

BlackMamba sequence has been deposited in GenBank database with the accession number MN902222.

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**Author contributions** All authors contributed as a team to the experimental design and the interpretation of data. VF, AT, GM, and VM performed experiments and data analysis. AV and RB performed bioinformatics analysis. OE supervised bioinformatics analysis. EZ performed FISH staining. GI, JI, TBH, FM, and VF diagnosed and stratified pathological samples. RW and DF performed the sequencing. VF and GI created the pathological and clinical database. PL, WC, and VM performed experiments and data analysis. AV and RB performed the sequencing. VF and GI created the pathological and clinical database. PL, WC, and JK contributed to the data analyses. GI, VF, and AC wrote the manuscript.

**Compliance with ethical standards**

**Conflict of interest** The authors declare that they have no conflict of interest.

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