MCM5 as a target of BET inhibitors in thyroid cancer cells

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Abstract

Anaplastic thyroid carcinoma (ATC) is an extremely aggressive thyroid cancer subtype, refractory to the current medical treatment. Among various epigenetic anticancer drugs, bromodomain and extra-terminal inhibitors (BETis) are considered to be an appealing novel class of compounds. BETi target the bromodomain and extra-terminal of BET proteins that act as regulators of gene transcription, interacting with histone acetyl groups. The goal of this study is to delineate which pathway underlies the biological effects derived from BET inhibition, in order to find new potential therapeutic targets in ATC. We investigated the effects of BET inhibition on two human anaplastic thyroid cancer-derived cell lines (FRO and SW1736). The treatment with two BETis, JQ1 and I-BET762, decreased cell viability, reduced cell cycle S-phase, and determined cell death. In order to find BETi effectors, FRO and SW1736 were subjected to a global transcriptome analysis after JQ1 treatment. A significant portion of deregulated genes belongs to cell cycle regulators. Among them, MCM5 was decreased at both mRNA and protein levels in both tested cell lines. Chromatin immunoprecipitation (ChIP) experiments indicate that MCM5 is directly bound by the BET protein BRD4. MCM5 silencing reduced cell proliferation, thus underlining its involvement in the block of proliferation induced by BETis. Furthermore, MCM5 immunohistochemical evaluation in human thyroid tumor tissues demonstrated its overexpression in several papillary thyroid carcinomas and in all ATCs. MCM5 was also overexpressed in a murine model of ATC, and JQ1 treatment reduced Mcm5 mRNA expression in two murine ATC cell lines. Thus, MCM5 could represent a new target in the therapeutic approach against ATC.

Introduction

Thyroid cancers are the most widespread malignancies of the endocrine system and represent approximately 1–1.5% of all tumor-related diseases. The overall 5-year survival of these tumors is approximately 85–90%, with the highest mortality rate reported for poorly differentiated histotypes (Durante et al. 2006, Patel & Shaha 2014). Anaplastic thyroid carcinomas (ATCs), which are wholly or partially composed of undifferentiated cells arising from thyroid follicular epithelium, represent the least common but most lethal thyroid cancer subtype.
Indeed, they are characterized by genetic and epigenetic aberrations that make them unresponsive to treatments based on radioiodine administration (Arturi et al. 2000, Schlumberger et al. 2007, Perri et al. 2011, Elisei 2012). Thus, innovative approaches for ATC treatment are required.

The term ‘epigenetics’ is commonly used to describe the study of hereditable changes in gene expression that are independent from alteration in the underlying DNA sequence. As aberrant gene expression is an essential characteristic that distinguishes neoplastic cells from normal ones, compounds able to target epigenetic alterations are becoming increasingly appealing for cancer treatment. Most of these molecules act on epigenetic ‘writers’ or ‘erasers’, that is, enzymes regulating DNA or histone covalent modifications. Notably, compounds targeting enzymes involved in DNA methylation and histone acetylation have been greatly investigated in several experimental models of solid tumors, including thyroid cancer (Puppin et al. 2005, Russo et al. 2013). Several other chromatin regulators, instead, act as ‘readers’: they possess peculiar domains that are able to interact with nucleosome covalent modifications and respond to upstream signal cascades to control gene expression (Dawson et al. 2012). A rising class of compounds that are able to target chromatin readers is called BET inhibitors (BETis). BET is the acronym of bromodomain and extra-terminal family of proteins (Nicodeme et al. 2010). The mammalian BET protein family includes four members: BRD2, BRD3, and BRD4, which are ubiquitously expressed, and BRDT, whose expression is confined to germ cells. These proteins possess a tandem bromodomain at the N-terminal region, by which they interact with histone acetyl-lysines in order to control gene transcription and cell proliferation (Chiang 2009). Due to the relevance of BET proteins, several inhibitors have been recently developed (Filippakopoulos et al. 2010, Nicodeme et al. 2010). BETis hinder proliferation of a wide range of hematopoietic malignant cell lines and nuclear protein in testis (NUT) midline carcinoma-derived cells (Filippakopoulos et al. 2010, Zubert et al. 2011). The use of BETis, as well as the use of other epigenetic compounds, might be a complicate issue in cancer treatment because these drugs act on a large spectrum of genes (Falahi et al. 2015). However, the antiproliferative outcome could be reasonably due to the deregulation of only a small subset of genes. For instance, repression of the c-MYC oncogene seems to play a major role in mediating the effects of BETis in leukemic cells (Ott et al. 2012). Thus, an attractive approach to study epigenetic compounds is to define their critical effector genes. For this purpose, in this study we investigated the effects of BET inhibition in two human ATC cell lines, analyzing the modulation of their global gene expression profiles. By using this approach, we propose MCM5 as a potential new target for ATC treatment.

Materials and methods

Human and murine cell lines

FRO (purchased from the European Collection of Cell Cultures, Salisbury, UK) and SW1736 (obtained from Cell Lines Service GmbH, Eppelheim, Germany) are human cell lines derived from ATC, both harboring a BRAF V600E mutation (Schweppe et al. 2008, Pilli et al. 2009). Both cell lines have been tested for being mycoplasma free and authenticated by short tandem repeats (STR) analysis to be appropriate cell lines of thyroid cancer origin. FRO and SW1736 were grown in DMEM medium (EuroClone, Milan, Italy) and RPMI 1640 medium (EuroClone), respectively, supplemented with 10% fetal bovine serum (Gibco Invitrogen, Milan, Italy), 2 mM l-glutamine (EuroClone), and 50 mg/mL gentamicin (Gibco Invitrogen), in a humidified incubator (5% CO₂ in air at 37°C). Cultured cells were treated with vehicle (DMSO, Sigma-Aldrich, St Louis, MO, USA) or the following agents: JQ1 (50 nM–10 μM in DMSO) (Cayman Chemical, Ann Arbor, MI, USA) and I-BET762 (50 nM–10 μM in DMSO) (Merck Millipore). T4898M and T3531L are ATC cell lines derived from tumors developed by [Pten, Tp53]thyr–/– mice (Antico Arciuch et al. 2011). Cells were grown in DMEM supplemented with 10% FBS.

Animals

[Pten, Tp53]thyr–/– and Ptenthyr–/– mice have been as described previously (Yeager et al. 2007, Antico Arciuch et al. 2010, 2011). The animals were housed in the Albert Einstein College of Medicine animal facility under specific pathogen-free conditions.

Cell viability and apoptosis

To test cell viability, an MTT assay was performed as described previously (Baldan et al. 2014). Briefly, 5000 cells were seeded onto 96-well plates. On the next day, the growth medium was replaced with vehicle-treated medium (NT, untreated cultures) or with medium containing JQ1
or I-BET762; plates were incubated for 0, 24, 48, and 72 h. All experiments were run in quadruplicate and cell viability was expressed as a percentage relative to baseline samples (T0). The percentage of cell viability assessed by MTT assay was used to determine EC50 concentrations from dose–response curves after 72 h treatment.

Caspase 3/7 activity was measured as a marker of apoptosis using the Apo-ONE Homogeneous Caspase-3/7 Assay Kit (Promega, Milan, Italy) according to the manufacturer’s protocol. Five thousand cells were seeded onto 96-well plates. On the next day, the growth medium was replaced with vehicle-treated medium (NT, untreated cultures) or medium containing JQ1 5 μM or I-BET762 5 μM; the plates were incubated for 48 h. All experimental points were run in triplicate, and apoptosis levels were expressed as a percentage relative to baseline samples (NT).

**Cell cycle analysis**

Cell cycle was determined by fluorescence-activated cell-sorting (FACS) analysis of DNA content. Briefly, FRO and SW1736 cells were collected after a 24, 48, and 72 h treatment with either JQ1 or vehicle (NT); the cells were fixed in cold ethanol 70% and stained with propidium iodide solution containing RNase and Triton-X100. Flow cytometric analysis was performed on a FACSCalibur (Becton Dickinson, Franklin Lakes, NY, USA) using ModFit LT analysis software (Verity Software House, Inc., Topsham, ME, USA). A minimum of 20,000 cells was analyzed for each sample. All experiments were run in triplicate.

**High-throughput RNA sequencing and analysis**

About 1.5 μg of good-quality RNA (RNA Integrity Number >7) was used as a starting material for preparation of library using the Illumina mRNASeq Sample Prep Kit v2.0, following the manufacturer’s instructions (Illumina, San Diego, CA, USA). The poly-A mRNA was fragmented for 1.5 min at 94°C, and all purification steps were performed by using 1X Agencourt AMPure XP beads (Beckman Coulter, Brea, CA, USA). Quality and quantity of each library were assessed by using an Agilent 2100 Bioanalyzer High Sensitivity and Qubit DNA High Sensitivity (Invitrogen). The libraries were pooled together and the obtained pool was checked on an Agilent Bioanalyzer 2100 in order to determine the molarity. Single-read sequencing was performed on the HiSeq2000 (Illumina, San Diego, CA, USA) generating 50-base reads.

The CASAVA 1.8.2 version of the Illumina pipeline was used for image analysis, base calling, and demultiplexing. Reads were first trimmed in order to remove lower base quality data with ERNE (extended randomized numerical aligner) (Vezzi et al. 2012), and adapter sequences were removed with cutadapt (https://pypi.python.org/pypi/cutadapt/1.3). TopHat (Trapnell et al. 2012) was used for mapping and annotation on the HG19 reference sequence. Differential gene expression analysis was performed according to the experimental design using Cufflinks (Trapnell et al. 2012).

**Gene expression assays**

Total RNA from human cell lines, treated with either JQ1 5 μM or vehicle, was extracted with the RNeasy Mini Kit (Qiagen) according to manufacturer’s instructions. About 500 ng of total RNA were reverse transcribed to cDNA using random exaprimers and MMLV reverse transcriptase (Life Technologies). Real-time PCRs were performed using Platinum SYBR Green qPCR SuperMix (Life Technologies) with the ABI Prism 7300 Sequence Detection Systems (Applied Biosystems). The ΔΔCT method, by means of the SDS software (Applied Biosystems), was used to calculate mRNA levels. Oligonucleotide primers for MCM5 (forward, 5'-GATCCTGGCATTTCTACAG-3'; reverse, 5'-CCCTGTATTTGAAGGTGAAG-3') and β-actin (forward, 5'-TTGTACAGGAGTCTCCCTGGC-3'; reverse, 5'-ATGCTATCACCTCCCTGTGTG-3') were purchased from Sigma-Aldrich.

Total mouse RNA (treated with either 0.5 μM or vehicle) was extracted with Trizol and reverse transcribed using the Maxima First Strand cDNA Synthesis Kit (Thermo Scientific). qRT-PCR was performed on a StepOnePlus apparatus using SYBR Green PCR Master Mix (Applied Biosystems) and custom-designed primers (forward, 5'-CAGAGGGGATTCAAAGGTG-3'; reverse, 5'-CGATCCAGTATTCCACGTTG-3'). Each sample was run in triplicate and 18S was used to control for input RNA. Data analysis was based on the Ct method, and experiments were repeated at least three times using at least two independent organ and cell pools.

**Chromatin immunoprecipitation**

The chromatin immunoprecipitation (ChIP) assay was performed as described previously (Baldan et al. 2015). For immunoprecipitation, samples were incubated with 10 μg of rabbit polyclonal anti-BRD4 (Active Motif, Carlsbad, CA, USA) or Rabbit IgG (Millipore) as negative control (NC).
After incubation with Dynabeads Protein A (Invitrogen), washing, and elution, the DNA was extracted with phenol/chloroform, and the aqueous phase recovered and precipitated. The extracted DNA was used as a template in qPCR with the following primers: MCM5 Forward Primer (CCTGTTCGTGGCGCTTTTGTTC), MCM5 Reverse Primer (GATCGTCAATCCCGCATG), NCR Forward Primer (TGGCTGATCCTTTTACAGGGAGTT), NCR Reverse Primer (TTTGGCAAGATGTTGAAAAAC). Fold enrichment was then calculated as signal over background (IgG).

Data are representative of three different experiments.

RNA silencing

For transient silencing of endogenous MCM5, the TriFECTa RNAi Kit (Integrated DNA Technologies, Inc., Coralville, IA, USA) was used. A ‘universal’ NC duplex that targets a site that is absent from human, mouse, and rat genomes was used. siRNA oligonucleotides were transfected at a concentration of 5 nM using DharmaFECT Transfection reagent (Thermo Scientific), according to the manufacturer’s instructions. The day before transfection, cells were plated in antibiotics-free medium and harvested 72 h after transfection. The gene-silencing efficiency was evaluated by mRNA and protein level analysis.

Annexin V assay

In order to test the percentage of viable/apoptotic cells after MCM5 silencing, an Annexin V assay was performed. Briefly, cells were washed with cold PBS, transferred to a polystyrene round-bottomed flow tube (Falcon; Becton Dickinson), and resuspended in 195 μL of 1 binding buffer (BB 10 mM; HEPES/NaOH, pH 7.4, 140 mM NaCl, and 2.5 mM CaCl₂). Five microliter of Fluorescein-conjugated Annexin V (Annexin V-FITC; Bender Med Systems, Vienna, Austria) was added and samples were incubated for 10 min at room temperature. After washing, cells were resuspended in 190 μL of BB in which 10 μL of propidium iodide stock solution (final concentration 1 μg/mL) was added. Flow cytometric analysis was done on CyAN, Dako Cytomation using Summit software (flow cytometer; Beckman Coulter). Forward scatter and side scatter were acquired in a linear mode. FITC and PI fluorescent signals derived from 488 nM excitation were detected in a logarithmic mode at FL1/PMT3 and FL2/PMT4, with FITC 530/30 nM filters and PI 585/42 nM filters, respectively, and a FL1/2 560 nM shortpass dichroic filter. Signals for forward and side scatter and fluorescence were collected for 10⁴ cells using the forward light scatter parameter as the master signal. Data are expressed as mean fluorescence intensity (FI) values. All experiments were run in triplicate.

MCM5 overexpression

To overexpress MCM5 protein, we purchased the TrueClone MCM5 cDNA cloned in pCMV6-AC, which contains a full open reading frame of the human MCM5 gene (Origene, Rockville, MD, USA). The overexpression vector was transfected with Turbofect reagent (Fisher Scientific S.A.S., Illkirch, France) at 2 μL per well. pCMV empty vector was used as NC. Once we verified the overexpression by immunoblotting, we performed a rescue experiment, in order to verify the direct involvement of MCM5 in JQ1 mechanism of action. Cells were transfected with pCMV-MCM5 or NC and exposed to 5 μM JQ1 or vehicle for 48 h. Cell viability was evaluated by MTT assay, as described previously.

Protein extraction and Western blot

Total protein extraction was performed as described previously (Passon et al. 2012). Briefly, FRO and SW1736 cells, incubated in vehicle-treated medium (NT, untreated cultures) or with JQ1 5 μM, were harvested by scraping and lysed with total lysis buffer (Tris–HCl 50 mM pH 8.0, NaCl 120 mM, EDTA 5 mM, Triton 1%, NP40 1%, protease inhibitors). For subcellular fractionation, pellets were resuspended in buffer B (10 mM Tris–HCl (pH 7.5), 1.5 mM MgCl₂, and 10 mM KCl supplemented with 1× protease inhibitor cocktail, 0.5 mM PMSE, 1 mM NaF, and 1 mM Na₂VO₄) and centrifuged at 2000 g for 10 min at 4°C. For nuclear extracts collection, pellets were resuspended in buffer B (20 mM Tris–HCl (pH 7.5), 0.42 M KCl, 1.5 mM MgCl₂, 20% (vol/vol) glycerol supplemented with 1× protease inhibitor cocktail, 0.5 mM PMSE, 1 mM NaF, and 1 mM Na₂VO₄) and centrifuged at 15,000 g for 30 min at 4°C. For western blotting analysis, proteins were electrophoresed on 10% SDS–PAGE and then transferred to nitrocellulose membranes, which were saturated with 5% nonfat dry milk in PBS/0.1% Tween 20. The membranes were then incubated overnight with rabbit polyclonal anti-BRD4 antibody 1:1000 (Active Motif), rabbit anti-lamin-β1 antibody 1:1000 (Abcam), rabbit polyclonal anti-MCM5 antibody 1:250 (Sigma-Aldrich), or rabbit anti-β-actin antibody 1:1000 (Abcam). The next day, membranes were incubated for 2 h with anti-rabbit immunoglobulin coupled to peroxidase 1:4000 (Sigma-Aldrich). The blots were developed using
Results

Biological effects of BETis on ATC cells

In a first set of experiments, we evaluated the sensitivity of two human anaplastic thyroid cancer cell lines (FRO and SW1736) to BETis. As shown in Fig. 1, incubation either with JQ1 (Fig. 1A) or I-BET762 (Fig. 1B) significantly reduced FRO and SW1736 cell viability at different time points. Relying on data obtained, for further experiments, we decided to use the median effective dose of 5 μM; i.e., the dose required to achieve 50% of the desired response in 50% of the population. To assess whether the decrease in cell viability was due to an increase of apoptotic cell death, caspase 3/7 levels were measured after 48 h treatment. Administration of JQ1 increased caspase levels in both cell lines in a quite striking manner, whereas I-BET762 treatment showed a less pronounced effect (Fig. 1C). In order to verify whether a 48 h I-BET762 treatment displayed a caspase-independent cell death, we perform a western blotting analysis of PARP. PARP is known to be a substrate for several proteins such as caspases, calpains, and cathepsins; it produces several specific cleavage fragments that resemble signature patterns of different cell death programs (Gobell et al. 2001; Chaitanya et al. 2010). As shown in Supplementary Fig. 2 see section on supplementary data given at the end of this article (Fig. 2A and B), ATC cells showed a marked increase of both apoptotic- and necrotic-specific cleaved-PARP. We, then, corroborated the BET inhibition effects on cell viability and cell death in 8505c cells, a third anaplastic thyroid cell line (Supplementary Fig. 3). Moreover, we used a third BETi: I-BET151. This latter is a pan-BETi having a slight different chemical structure compared to I-BET762. I-BET151 reduced cell viability similarly to I-BET762 (Supplementary Fig. 1A and B, and Supplementary Fig. 3). Hereinafter, we decided to proceed in our investigation with JQ1-treatment only, as it revealed to be the most effective drug, in terms of biological effects, in both cell lines. A FACS analysis of the cell cycle was then performed on both cell lines treated with 5 μM JQ1 for 24, 48, and 72 h. JQ1-treatment significantly increased the proportion of cells arrested in G0/G1 at each analyzed time point (Fig. 1D).

Differential gene expression after BET inhibition

To identify potential molecular targets connected to BET inhibition, we performed a high-throughput RNA sequencing (RNA-seq) analysis on both FRO and SW1736 cells treated with 5 μM JQ1 or vehicle for 24 h. To assess

Human tissues and immunohistochemistry

A series of 12 normal thyroid glands (NTs), 25 follicular adenomas (FAs), 23 follicular thyroid carcinomas (FTCs), 36 papillary thyroid carcinomas (PTCs), and eight ATCs were selected from the files of the Institution of Anatomic Pathology of the University of Udine (Udine, Italy), and the most representative block of each lesion was retrieved from the archive. Representative tumor-bearing areas were selected and a tissue microarray was constructed. All samples were diagnosed by referral pathologists of the institution and then reviewed by a single experienced pathologist, thus including only patients with a confirmed diagnosis. Briefly, 5 μM formalin-fixed paraffin tissue sections mounted on SuperFrost Plus slides (Menzel-Gläser, Braunschweig, Germany) were placed in the PT Link pre-treatment module (DAKO A/S, Glostrup, Denmark), which performs automatically the entire pre-treatment process of de-paraffinization, rehydration, and epitope retrieval using the Low pH Target Retrieval Solution (0.001 M citrate buffer pH 6.0) at 98°C for 40 min from DAKO. Endogenous peroxidase activity was blocked by incubation using the peroxidase block solution (DAKO) for 5 min. Primary rabbit polyclonal antiserum to MCM5 (Sigma-Aldrich) diluted 1:1000 was applied and incubated for 60 min at room temperature. After washing, the slides were incubated with the DAKO EnVision FLEX System (DAKO) according to the manufacturer’s guidelines. For reaction visualization, 3,3′-diaminobenzidine tetrahydrochloride was used as chromogen. The sections were counterstained with Mayer’s hematoxylin. Using light microscopy, the entire section was scanned at high-power magnification (400×) and nuclear immunostaining was evaluated semi-quantitatively as percentage of positive cells. Staining intensity was scored as light, moderate, and strong. A final score was then calculated by multiplying the percent staining by the intensity level.

Statistical analysis

mRNA and protein levels, cell viability, and apoptosis levels were expressed as means±s.d., and significances were determined by a one-way ANOVA followed by Dunnett’s test performed with GraphPAD Software for Science. *P value<0.05 was considered to be statistically significant.
the transcriptional changes induced by BET inhibition, a comparison between cells treated with JQ1 or vehicle was performed. Heat maps representing each analyzed condition are shown in Fig. 2A. Expression of 14,199 and 13,013 genes was detected in FRO and SW1736 cell lines, respectively. Vehicle-treated (NT) FRO and SW1736 cells showed a quite different mRNA profile. After filtering low quantity reads, results showed that approximately 2000 genes were differentially expressed after JQ1 treatment either in FRO or SW1736 cells (at a log2 fold change >2) (Supplementary Tables 1 and 2, see section on supplementary data given at the end of this article). Among them, 768 genes were significantly upregulated and 1229 were downregulated in response to JQ1 treatment (Fig. 2B). Venn diagrams showed that the majority of altered genes was specific for each cell line, which likely underlies the differential mRNA profile observed with vehicle treatment. However, there was a core set of 288 genes commonly affected by JQ1 treatment in both cell lines, most of which decreased their expression (n = 213). RNA-seq data, then, were confirmed by qPCR (Supplementary Fig. 4). The shared top 20 differentially expressed genes (Table 1) were then subjected to Ingenuity Pathway Analysis in order to outline which pathways were mostly affected by JQ1 treatment. Various signaling cascades turned out to be modified by BETis treatment: regulation of DNA and metabolic processes proved out to be the most deregulated one (Supplementary Table 3). The RNA-seq analysis identified a large number of JQ1 putative targets, primarily in cell cycle-linked pathways.
These data are consistent with Da Costa’s findings in leukemia cells (Da Costa et al. 2013). Several studies have proposed a possible role of c-MYC downregulation as a major mechanism of action of BETis (Alderton 2011, Zuber et al. 2011). Our RNA-seq data show that this oncogene is not a critical factor in JQ1-mediated effects in ATC cells, because JQ1 treatment did not significantly affect c-MYC expression in neither of the two cell lines (Supplementary Fig. 4). BETis, however, modified the expression of several cell cycle regulators such as RB proteins, E2F transcription factors, and CDK inhibitors (Supplementary Table 4).

**BETis effects on MCM5**

Considering that cell cycle regulatory genes were the most affected by JQ1 treatment, we focused on MCM5, one of the top 20 downregulated genes, which is known to be involved in G1/S cell cycle transition, Figure 2. Gene expression modifications in ATC JQ1-treated cell lines. (A) Heat maps showing the hierarchical clustering of mRNA targets in FRO and SW1736 cell lines (n=1997). Cells were treated with either 5μM JQ1 or vehicle for 24h. (B) Venn diagrams representing the comparison of both upregulated (left) and downregulated (right) genes between FRO and SW1736 cell lines after RNA-seq data analysis. Purple gene clusters represent shared modified genes between FRO and SW1736 cell lines.

Figure 3. Downregulation of MCM5 in human and murine ATC cell lines after JQ1 treatment. (A) FRO and SW1736 cells were treated either with 5μM JQ1 or vehicle for 24h, and MCM5 mRNA expression was evaluated by qPCR. All samples were run in triplicate. Vehicle-treated (NT) FRO cells were arbitrarily set at 1.0 and mRNA levels are expressed as relative expression values. (B) FRO and SW1736 were treated with either 5μM JQ1 or vehicle for 72h, and MCM5 protein levels were detected by western blotting. (C) Densitometric analysis of MCM5 protein levels in ATC cells after 5μM JQ1 or vehicle treatment. (D) T4888M and T3531L cells were treated with either 0.5μM JQ1 or vehicle for 72h, and Mcm5 mRNA expression was evaluated by qPCR. All samples were run in triplicate. Vehicle-treated (NT) T4888M cells were arbitrarily set at 1.0 and mRNA levels are expressed as relative expression values. Results are shown as mean±s.d. ***P<0.001, ****P<0.0001 by ANOVA test. Data are representative of three independent experiments.
coordinating origin activation and replication fork progression. MCM5 belongs to the minichromosome maintenance complex family, which has been associated with the highly proliferative phenotype of ATC (Guida et al. 2005). Several studies have established that MCM5 protein level is a good diagnostic marker of malignancy for bladder, esophageal, and gastric cancers (Kebebew et al. 2006). After confirming RNA-seq data on mRNA levels (Fig. 3A), we evaluated whether a 5μM JQ1 treatment significantly altered MCM5 protein levels in FRO and SW1736 cells (Fig. 3B). After 72 h incubation with JQ1, a significant reduction in MCM5 protein levels was observed (Fig. 3C). To extend and further validate this finding, we used two murine ATC cell lines derived from tumors developed by a genetically defined mouse model of ATC.

Treatment of both lines with 0.5μM JQ1 for 72 h resulted in a significant downregulation of MCM5 RNA (Fig. 3D). To evaluate whether MCM5 is a direct target of BRD4, which is the BET protein whose activity is inhibited by JQ1, we performed a ChIP-qPCR analysis in FRO and SW1736 cells. We first evaluated if the two cell lines expressed comparable BRD4 protein levels (Fig. 4A). After the ChIP assay, we observed an average five-fold MCM5 enrichment in the BRD4 immunoprecipitate compared with the IgG one. To confirm specificity, an intergenic region (NCR) has been used as an NC for BRD4 binding (Fig. 4B). In order to test the biological relevance of MCM5 gene repression upon BET inhibition, an RNA interference analysis was performed. MCM5 gene expression was silenced both in FRO and in SW1736 cells, and immunoblot analysis
revealed an almost total abolishment of endogenous MCM5 protein level with siRNA#2 treatment (Fig. 4C). Furthermore, MCM5 silencing, especially with siRNA#2, significantly decreased cell viability ($P<0.001$), increasing the percentage of apoptotic cells after Annexin V staining compared with NC (Fig. 4D). Similarly, to what we observed treating cells with JQ1, this outcome was more pronounced in SW1736 than in FRO cells. To further prove that MCM5 is directly involved in the cell viability decrease after 5 μM JQ1 treatment, we performed a rescue experiment. FRO and SW1736 cells were treated with pCMV-MCM5 or NC and exposed to JQ1 for 48 h. As shown in Fig. 4E, MCM5 protein overexpression partially rescued its protein levels in JQ1-treated cells. To validate that this phenomenon is associated with a cell viability rescue, an MTT assay was performed. FRO and SW1736 cells were treated with pCMV-MCM5 or NC and exposed to JQ1 for 48 h. In vivo MCM5-level evaluation

Our data underline the relevance of MCM5 in thyroid cancer cell proliferation. Thus, MCM5 expression was evaluated in a panel of 12 NTs, 25 FAs, 23 FTCs, 36 PTCs, and 8 ATCs. Representative images of MCM5 staining are shown in Fig. 5A. MCM5 was faintly expressed in normal thyroid tissue and FAs. PTCs showed a dual behavior: some samples displayed an intense and high MCM5 nuclear staining, whereas some others a much lesser one, with only a cytoplasmic signal. In FTCs, nuclear staining was less intense and frequent, with a predominant cytoplasmic MCM5 staining. ATCs, instead, showed a marked nuclear MCM5 localization. No significant differences were observed in terms of MCM5 nuclear or cytoplasmic staining intensity among different areas of each tumor. Quantification of the nuclear immunohistochemical signal was obtained by computing the percentage of cell positivity and intensity of signal (Fig. 5B). As cytoplasmic MCM5 protein represents the inactive MCM5 isoform (Abe et al. 2012), the strong nuclear staining observed in all ATCs and several PTCs would indicate its hyperactivation. Finally, we measured the expression of Mcm5 in four primary ATCs developed by [Pten, Tp53]thyr–/– mice (Antico Arciucho et al. 2011). In agreement with the human data, Mcm5 was found significantly overexpressed in mouse ATC, compared with normal thyroids (Fig. 5C), whereas its expression was not altered in FTCs developed by Ptenthyr–/– mice (data not shown).

Discussion

The cross talk between genomic and epigenomic elements in driving generation and progression of both
hematologic malignancies and solid tumors has been extensively highlighted (Martín-Subero et al. 2013). Among epigenetic anticancer drugs, those targeting BET proteins have recently emerged as promising therapeutics in a remarkable range of diseases (Shi & Vakoc 2014, Wee et al. 2014). ATC is the most aggressive thyroid cancer subtype and it does not significantly respond to any therapy. In order to find new strategies to treat this aggressive thyroid cancer subtype, we investigated BETis target genes in ATC cell lines. As expected, human ATC cell lines FRO and SW1736, treated with three different BETis, displayed a decrease in cell viability associated with an increase in cell death phenomena. These data were also been confirmed in a third ATC cell line (Supplementary Fig. 3). In our experimental model, BET inhibition triggers a multitarget cascade resulting in substantial biological effects, as demonstrated by RNA-seq data. The identification of genes mediating BETis antiproliferative effect could provide innovative therapeutic targets. Among hundreds of deregulated genes after BET inhibition, a large fraction belongs to the regulators of cell cycle and apoptosis, that is, E2F family members, CDKs and CDK inhibitors, and other TP53 pathway-related elements (Møller 2003, Smallridge et al. 2009). These data are similar to those previously published for study in medulloblastoma and metastatic melanoma (Henssen et al. 2013, Segura et al. 2013). All these findings corroborate the hypothesis that BETis disrupt cell cycle control, driving G0/G1 arrest and inducing cell death, by acting on multiple targets. Ott and coworkers observed that JQ1 treatment decrease the expression of IL7R in leukemic cell lines, at both mRNA and protein levels (Ott et al. 2012). In anaplastic thyroid cell lines, a 24 h JQ1 treatment exhibited a significant downregulation of IL7R mRNA levels, but this reduction was not confirmed by analyzing protein levels (Supplementary Fig. 5). We would like to stress that endogenous IL7R protein is not so abundant in these two cell lines; therefore, the misregulation of a protein involved during lymphocyte development seems not to perform a substantial role in JQ1 effects in thyroid cancer cells. Our data, instead, show at least a ten-fold downregulation of MCM5 expression in JQ1-treated cells, suggesting a critical role of this protein in the molecular cascade triggered by BETis. As confirmation, MCM5 proved to be a direct target of BRD4, as highlighted in the ChIP-qPCR assay. Moreover, MCM5 overexpression moderately attenuated FRO and SW1736 JQ1 sensitivity, in terms of cell viability, as shown in the rescue experiment (Fig. 4F). This mild effect is reasonably due to the multitarget activity of epigenetic drugs, which act simultaneously on a wide range of pathways, as shown in our RNA-seq analysis. Accordingly to our data, poor rescue effects have been obtained for other targets of BETis (Lockwood et al. 2012, Bandopadhayay et al. 2014).

MCM5 belongs to the minichromosome maintenance complex, which includes elements of the pre-replicative complex that are essential for DNA replication (Noseda & Karsan 2006). MCM proteins are considered as licensing

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**Figure 5**

MCM5 expression in thyroid tumors. (A) Representative images of MCM5 expression in NTs, FAs, PTCs, FTCs, and ATCs. (B) Quantitation of MCM5 expression in normal and neoplastic thyroid tissues. Quantitation was obtained by using the IHC score, calculated as described in Materials and methods section. (C) Mcm5 expression was evaluated by qPCR in control thyroids from wild-type mice and primary ATCs from [Pten, Tp53]−/− mice. All samples were run in triplicate. Wild-type (WT) expression was set at 1.0 and mRNA levels are expressed as relative expression values. Results are shown as mean±s.d. P<0.05, **P<0.01, ****P<0.0001 by ANOVA test.
components for the S-phase initiation. MCMs oscillate between nuclear and cytoplasmic localizations (Madine et al. 2000). Nuclear MCMs associate with DNA during G1 and early S-phase, allowing the control of replication origin firing, in order to restrict the chromosome replication to only one round per cell cycle (Romanowski & Madine 1997). MCMs are restricted, instead, to cytoplasm during G2/M-phase in order to prevent re-replication phenomena (Laskey 2005). Recently, components of the MCM family have been described as aberrantly expressed in many malignancies, including thyroid carcinomas (Giaginis et al. 2010). In agreement with these findings and corroborating our in vitro and in vivo results, MCM5 immunostaining demonstrated that ATCs contained more MCM5-positive cells than did normal thyroid gland (Fig. 5). Several studies have demonstrated that the expression of MCM5 is controlled by different pathways (Guida et al. 2005, Laskey 2005,Noseda & Karsan 2006, Abe et al. 2012). Fig. 6 summarizes major players controlling MCM5 gene expression, as well as their functional interactions. By comparing Fig. 6 and Supplementary Table 4, it is undeniable that many proteins that upregulate MCM5 expression are also downregulated by BETis. Interestingly, JQ1 reduces the expression of some elements involved in the replication fork assembly, thus affecting cell cycle mechanisms, in which, in turn, MCM proteins are involved. Altogether, this cascade could lead to G0/G1 arrest (Fig. 1D) and cell death. In conclusion, our findings support the hypothesis that MCM5 targeting may be regarded as an effective molecular therapy against ATC.

Figure 6
Biological network related to MCM protein function. Network diagram (upper part) showing key molecules controlling MCM protein levels. White symbols represent proteins whose mRNA levels are modulated by BETis in our transcriptome analysis (see Supplementary Table 4); gray symbols indicate proteins whose mRNA levels are not modulated by BETis. Arrows indicate functional relationship between proteins (delineated in Madine et al. 2000, Guida et al. 2005, Laskey 2005, Abe et al. 2012). The lower part indicates factors cooperating with MCM proteins in the replicative fork assembly. All of these factors are downregulated by BETis (see Supplementary Table 4). Arrowheads show the relationship between MCMs and cell cycle phases.
Supplementary data
This is linked to the online version of the paper at http://dx.doi.org/10.1530/ERC-15-0322.

Declaration of interest
The authors declare that there is no conflict of interest that could be perceived as prejudicing the impartiality of the research reported.

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