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Loss of CITED1, an MITF regulator, drives a phenotype switch in vitro and can predict clinical outcome in primary melanoma tumours

CITED1 is a non-DNA binding transcriptional co-regulator whose expression can distinguish the ‘proliferative’ from ‘invasive’ signature in the *phenotype-switching* model of melanoma. We have found that CITED1 expression is repressed by TGFβ in addition to other ‘proliferative’ signature genes while the ‘invasive’ signature genes are upregulated. In agreement, CITED1 positively correlates with MITF expression and can discriminate the MITF-high/pigmentation tumor molecular subtype in a large cohort (120) of melanoma cell lines. Interestingly, CITED1 overexpression significantly suppressed MITF promoter activation, mRNA and protein expression levels while MITF was transiently upregulated following siRNA mediated CITED1 silencing. Conversely, MITF siRNA silencing resulted in CITED1 downregulation indicating a reciprocal relationship. Whole genome expression analysis identified a phenotype shift induced by CITED1 silencing and driven mainly by expression of MITF and a cohort of MITF target genes that were significantly altered. Concomitantly, we found changes in the cell-cycle profile that manifest as transient G1 accumulation, increased expression of CDKN1A and a reduction in cell viability. Additionally, we could predict survival outcome by classifying primary melanoma tumors using our *in vitro* derived ‘CITED1-silenced’ gene expression signature. We hypothesize that CITED1 acts a regulator of MITF, functioning to maintain MITF levels in a range compatible with tumourigenesis.
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INTRODUCTION

CITED1 is the founding member of the CITED (CBP/p300-Interacting transactivator with glutamic acid [E]/aspartic acid [D]-rich C-terminal domain) family of transcriptional co-regulators and was originally cloned from a differential display screen between pigmented mouse B16 melanoma cells and their dedifferentiated non-pigmented derivative, B16F10s. This led to speculation that CITED1 or msg1 (melanocyte specific gene 1) as it was known at that time, was involved in the process of pigmentation (Shioda et al. 1996). Subsequently, Nair et al. reported that stable overexpression of CITED1 increased the levels of tyrosinase, dopachrome tautomerase (Dct) and melanin in B16 cells, reinforcing the idea that it had a role in melanogenesis (Nair et al. 2001). By 2005, as gene expression profiling became relatively commonplace, CITED1 was identified in several new screens of tumors and cell lines: two studies identified CITED1 as a gene whose expression distinguished nevi from primary melanoma, another found CITED1 to be upregulated in advanced stage melanomas in comparison to benign nevi or melanoma in situ, while expression profiling of an in vitro progression model identified CITED1 among a signature of genes lost in aggressive melanoma lines relative to primary melanocytes in culture (Ryu et al. 2007; Haqq et al. 2005; Talantov 2005; Smith et al. 2005).

Based on extensive gene expression profiling of melanoma cell lines in vitro, Hoek et al. proposed the ‘phenotype-switching’ model of melanoma that was independent of the degree of transformation or disease progression and
sought to explain the observation that melanoma cells altered between two states: those with high proliferative potential that are less invasive and those with high metastatic potential that are less proliferative. These separate but alternating states are controlled by different transcriptional programs and can be defined by specific gene signatures (Hoek, Eichhoff, et al. 2008a). MITF expression and many of its known targets (TYR, MLANA) define the ‘proliferative’ group, while the ‘invasive’ signature group is characterized by expression of negative regulators of the Wnt signaling pathway (WNT5A, DKK1, CTGF). CITED1 expression was associated with the proliferative pathway signature and subsequently confirmed in an updated and expanded data set to be significantly correlated with the proliferative phenotype (P<1.00E-05, http://www.jurmo.ch/hopp, accessed 19 March 2013) (Hoek et al. 2006; Widmer et al. 2012).

Studies on CITED1 suggest that it is a non-DNA binding nuclear transcriptional co-regulator capable of influencing TGFβ induced transcription mediated by ligand-induced SMAD hetero-oligomerization; estrogen-dependent transcription mediated by ERα, and Wnt/β-Catenin-dependent transcription. These effects are dependent on CITED1-CBP/P300 binding via the conserved CITED family CR2 domain and while in the case of ERα, CITED1 is thought to act by stabilizing the CBP/P300-ERα interaction, in the case of β-Catenin it acts to repress transcription by competing for binding with CBP/P300 transcriptional co-activators (Shioda et al. 1998; Yahata 2001; Yahata 2000; Plisov 2005).
Microphthalmia-associated transcription factor, MITF, acts as a master-regulator of melanocyte differentiation and as a result has been intensely studied in the field of melanoma research (Widlund & Fisher 2003; Levy et al. 2006). It is a basic helix-loop-helix leucine zipper transcription factor that recognizes E-box and M-box sequences in the promoter regions of its target genes. Highlighting its importance in the disease, amplification of MITF locus has been found in >15% of metastatic melanomas and germline mutations in MITF that predispose carriers to melanoma development have also been found (Garraway et al. 2005; Bertolotto et al. 2011; Yokoyama et al. 2011). In melanoma cells the target genes of MITF include most notably TYR, MCIP, DCT, MLANA involved in the process of pigmentation; cell cycle regulators such as CDK2 and CDKN1A and the more recently identified BRCA1 gene that has, with other target DNA repair genes, defined a role for MITF in the DNA damage response (DDR) (Strub et al. 2011; Beuret et al. 2011; Giuliano et al. 2010).

The regulation of MITF is complex and tightly controlled, exhibiting both transcriptional and post-translational regulation. There are several transcript isoforms, of which MITF-M is the dominant form expressed in melanocytes. Multiple signaling pathways converge on the MITF-M specific promoter that harbors binding sites for PAX3, SOX10, CREB, FOXD3, LEF-1 and BRN2 among other transcription factors (Yokoyama & Fisher n.d.; Levy et al. 2006). Additionally, the MITF target gene CDKN1A/P21 has been shown to act as
reciprocal transcriptional cofactor independently of its CDK inhibitor function, suggesting the existence of at least one positive feedback loop (Sestáková et al. 2010).

MITF post-translational activity can be affected by phosphorylation, sumoylation, ubiquitination and by binding with proteins that block access to the DNA binding domain such as PIAS3 (Yokoyama & Fisher n.d.; Levy 2001). Oncogenic BRAF (but not wildtype BRAF), which is mutated in up to 50% of melanomas, also regulates MITF via simultaneously stimulating MITF activation through ERK phosphorylation, which leads to its subsequent degradation, and by inducing transcription of MITF via BRN2 upregulation (Davies et al. 2002; Wellbrock et al. 2008).

The consensus regarding why the cell invests such effort in maintaining control of MITF levels and why there are so many regulatory mechanisms, is that melanocytes and melanoma are exquisitely sensitive to even small variations in MITF expression. Ultimately its activity must be sustained within the narrow window permissive for continued survival and proliferation. In this study, we characterise the role of CITED1 as a novel regulator of MITF in melanoma.

MATERIALS AND METHODS

Cell lines
Cell lines were obtained from ATCC. HT144 and SKMEL3 cells were cultured in McCoy’s5A supplemented with 10% and 15% fetal bovine serum (FBS), respectively. A2058, WM852 and WM239 were cultured in RPMI 1640 supplemented with 10% FBS; A375 and HMBC cells were cultured in DMEM supplemented with 10% FBS and SKMEL5 cells were cultured in MEM media supplemented with 10% FBS. Cells were grown in the presence of penicillin and streptomycin (50 l.U./mL).

**Gene expression analysis**

RNA was isolated (4 replicates for each treatment) using a Qiagen RNeasy Plus mini-kit and the quality determined using a Bioanalyser (Agilent). Gene expression experiments were performed using the Illumina HT12 array covering more than 47,000 transcripts and known splice variants across the human transcriptome. The raw data was quantile normalized and Illumina control probes were removed from subsequent analysis using BASE (Vallon-Christersson et al. 2009). The data were exported to MeV, log2 transformed and gene and sample centered (Al et al. 2003). SAM (significance of microarray analysis) was performed using a two-group comparison; for the siRNA experiment the groups corresponded to siNEG vs #1 & #3 siCITED1 and for the TGFβ1 experiment the groups corresponded to cells with or without TGFβ1 treatment. In both cases there was a median false discovery risk of 10 false-positive transcripts. Hierarchical clustering was performed to visualize the data. 312 probes were found to be significantly altered in the
siRNA experiment (208 upregulated and 104 downregulated) while 1009 probes were significantly altered by TGFβ1 treatment. DAVID was used to assist in functional annotation of the final gene lists (Huang et al. 2007).

For the publically available data cited, 120 melanoma cell lines from three cohorts (PMID: 17516929, 16827748, 20406975) analyzed by Affymetrix gene expression microarrays were collected, individually MAS5 normalized, and merged into a single cohort. Probe sets were collapsed into single genes and mean-centered across the entire cohort. Data from Harbst et al. were classified using nearest centroid and pearson correlation. Survival analysis and multivariate cox regression methods were performed in R.

**Transient transfections, promoter-reporter assay and TGFβ1-treatment**

Transient transfections were performed using Lipofectamine2000 and Opti-MEM reduced serum media (Life Technologies) according to the manufactures recommendations. siRNA was purchased from Applied Biosystems and the notations in the text: siNEG, #1 siCITED1 and #3 siCITED1 correspond to the catalogue ID numbers #4390843, #s8965 and #s224062 respectively. For the MITF targeting siRNA; N, siM1 and siM3 correspond to the catalogue ID numbers #4390843, #s8790 and #ss8792, respectively. For the luciferase reporter assay, a Dual-Luciferase Reporter assay system #E1910 (Promega) was used to measure relative reporter activity on a FLUOstar Omeaga microplate reader (BMG Labtech). A375 cells were transfected with a luciferase reporter construct harboring 2.3kb of the
MITF-M specific promoter in a PGL2 vector (Wellbrock et al. 2008). A pRL-Renilla Luciferase reporter vector was used as a control for each transfection. CITED1 was overexpressed using a pRc/CMV containing a N-terminal HA-tagged human CITED1 (transcript isoform 1) referred to as pCITED1 in the text. An empty CMV-promoter expression plasmid, pcDNA3.1 (+) was used as a negative control. Recombinant human transforming growth factor-β1 (TGFβ1), #PHG9203 was purchased from Invitrogen. For the A2058 gene expression experiment, cells were exposed to either 5 or 10ng/ml TGFβ1 in serum-free media for 24 hours. In the case of the Luciferase reporter assay, cells were serum starved the day after transfection for 3 hours and exposed to 5ng/ml TGFβ1 in serum free media for 24 hours prior to harvesting.

Antibodies and Immunoblotting
The following antibodies were used: anti-CITED1, #AB15096 from Abcam; anti-MITF (C5 clone), # MA5-14146 from ThermoScientific; anti-MITF (D5 clone) from Dako, #M3621, (used in Fig. 4c); anti-CDKN1A/P21, #2947 and anti-CDKN1C/P57, #2557 were purchased from Cell Signaling Technology and anti-β-Actin (AC-15), #A5441 from Sigma-Aldrich. Cell lysates were resolved by SDS-PAGE and transferred to 0.45 μm PVDF membranes by electroblotting. The membranes were blocked in 5% non-fat milk in TBST prior to incubation with primary antibodies diluted 2.5% non-fat milk. The blots were probed with the appropriate secondary antibodies (Pierce...
Biotechnology) in 5% non-fat milk. The membranes were developed using ECL (GE Healthcare).

**Cell cycle analysis**

Flow cytometry was performed on a FACSCalibur (BD Biosciences) and subsequently analysed using ModFit (Verity House Software). Briefly, following transfection, confluent cells were detached, washed in 1XPBS and fixed in 70% ethanol. Prior to analysis they were stained with a propidium iodide solution and a 20G syringe was used to obtain a homogenous single cell solution. All events were saved (up to 20,000 events per replicate) ungated, using BD Cell Quest and the data exported to ModFit where following selection of the appropriate ploidy status, a standard auto-analyis fit using autolinerarity was performed. We found that a 2-cycle aneuploid-dip/tetraploid was appropriate for HT144 and A2058 while 1-cycle diploid was suitable for A375.

**Alamar Blue Assay**

The Alamar blue assay reagent was purchased from Invitrogen and used according to the manufactures’ instructions. Briefly, following transfection cells were seeded into 96-well plates at 5000 cells/well. In each experiment, for each of the treatments i.e.: siNEG, #1 siCITED1 and #3 siCITED1, 8 wells spread over 3 rows were used. At the indicated time points (4, 72, 96 and 120 hours post-transfection), Alamar blue was added and the cells incubated at 37°C for 2 hours. Fluorescence was measured (544nm) on a FLUOstar
Omega microplate reader (BMG Labtech). The values obtained at the 4-hour time point were used to normalize the fluorescence readings to account for any initial cell counting error. Cells were also seeded in parallel for Western blot analysis (72, 96, 120 hours) to ensure successful CITED1 down regulation.

**Droplet digital PCR**

RNA was isolated from cells using a Qiagen RNeasy Plus mini-kit and quantified using a Nanodrop spectrophotometer (ThermoScientific). cDNA was generated from 50-100ng total RNAs using ‘iScript Advanced cDNA synthesis for RT-qPCR’ (Bio-Rad). Bio-RAD’s ‘ddPCR Supermix for Probes’ was then used with predesigned TaqMan gene expression assays (Applied Biosystems) consisting of specific primers and FAM labelled probes for MITF (#Hs01117294_m1), MITF-M isoform specific transcript (Hs00165165_m1)*, CITED1 (#Hs00918445_g1) and IPO8 (#Hs00183533_m1). (*There appeared to be no advantage in using the MITF-M isoform specific transcript over the MITF probe that could measure multiple isoforms). A manual cut-off for positive/negative droplets was selected using the Biorad QuantaSoft™ data analysis suite to calculate the relative copies/µl of each transcript.

**RESULTS**
TGFβ induces expression of the invasive signature genes while suppressing a cohort of proliferative signature genes including CITED1.

Hoek et al. noted that many of the genes that defined the invasive phenotype were commonly TGFβ-driven while at the same time only the proliferative signature phenotype cells were sensitive to TGFβ growth inhibition in vitro (Hoek et al. 2006). That MITF levels increase and invasiveness is enhanced in response to TGFβ stimulation was also confirmed subsequently (Pierrat et al. 2012; Pinner et al. 2009). In agreement, we showed that the melanoma cell line A2058 upregulates WNT5A in response to TGFβ exposure and that exogenous Wnt-5a in turn increased their invasive potential (Jenei et al. 2009). For the present study, in an effort to examine what other phenotype specifying genes were directly regulated by TGFβ, we performed gene expression analysis and found TGFβ treatment resulted in both upregulation of invasive signature genes and suppression of genes characterizing the proliferative phenotype (Fig. 1a). The effect is most pronounced if only those signature genes that were deemed significantly altered by TGFβ treatment are examined. The original signature set defined by Hoek et al., was redefined as more public datasets became available and has a slightly different but overlapping gene profile based on the top ranked differentially expressed genes (Fig. 1b). Both MITF and CITED1 are in the proliferative cohort and their response to TGFβ treatment was confirmed at protein level in A2058 cells (Fig. 1C).
CITED1 expression positively correlates with the expression of MITF

Examination of publically available gene expression data on 120 melanoma cell lines demonstrated a consistent positive correlation between CITED1 and MITF expression (r=0.6543). Each cell line was assigned as either ‘proliferative’ or ‘invasive’ based on a score derived from the averaged expression values of the approximately 50 genes in each defining signature set that had matching gene symbols in our data (Fig. 2a). We also confirmed the correlation in cell lines derived from our own lab (Fig. S1). This was important as inconsistency in interlaboratory phenotype signatures has previously been reported (Widmer et al. 2012). We could additionally confirm expression at the protein level (Fig. 2b)

Gene expression analysis reveals CITED1 silencing can induce a phenotype-switch

To investigate the function of CITED1 in melanoma, we transiently downregulated its expression using CITED1 targeting siRNA. We choose the HT144 cell line as it had a relatively high level of detectable CITED1 mRNA and protein expression. A scatterplot of the 120 cell lines assigned as either ‘proliferative’ or ‘invasive’ based on the maximum matching gene signature score demonstrates the shift in phenotype that occurs following CITED1 downregulation (Fig. 3a,b). A heatmap of the expression profiles clearly illustrates that the shift is due to a general induction of the ‘proliferative’ and suppression of the ‘invasive’ cohort (Fig. 3c). It was apparent that the #3 siCITED1 siRNA was not as effective at switching the cells as the #1
siCITED1, this was observed consistently throughout our experiments and may be due to the fact that #3 siCITED1 was not as successful at silencing CITED1 (Fig. 3b, inset).

**CITED1 is a reciprocal regulator of MITF and impacts MITF target gene expression**

A heatmap of only the significantly differentially induced transcripts between siNEG and both #1 & #3 siCITED1 highlights the identity of the genes responsible for the expression switch (Fig. 4a). Of most relevance, we found MITF, a known driver of the proliferative phenotype switch and many of its previously known downstream targets, these also included genes categorized by Gene Ontology annotation (GO) as related to pigmentation and UV/DNA damage response (Fig. 4a) (Hoek, Schlegel, et al. 2008b; McGill et al. 2006; Sánchez-Martín et al. 2002; Strub et al. 2011). We could confirm that indeed MITF protein levels were affected by siCITED1 in HT144 cells and that conversely, overexpression of CITED1 in A2058 cells, resulted in downregulation of MITF (Fig. 4b). Strub et al. identified a large number of genomic targets of MITF by ChIP-seq analysis (Strub et al. 2011). A comparison of the genes differentially expressed by siCITED1 compared to siNEG, revealed that there was significant enrichment of these potential targets (Fig. S2a). Notably, genes both up and down regulated by siCITED1 are represented among genes defined as having MITF-occupied promoters (Fig. S2b). We also found that downregulation of MITF using siRNA in HT144 cells (Fig. 4c) and in WM293A, and SKMEL5 cells (Fig. S3a,b) resulted in
decreased protein expression of CITED1 suggesting reciprocity between these factors.

**Induction of MITF by CITED1 silencing transiently restrains cell cycle progression and impacts cell viability**

To investigate the effect of CITED1 silencing on melanoma cells behaviour we analysed the cell cycle distribution following siRNA treatment, by flow cytometry. In siCITED1 treated HT144 cells we saw G1 accumulation as indicated by an increase in the diploid G1 fraction and a concomitant reduction in the total S-phase fraction peaking at 33 hours but also observed at 48 and 72 hours post-transfection in comparison to siRNA control HT144 cells. Again, the effect was apparent but not as pronounced using the #3 siCITED1 (Fig S4a). Similar effects were seen in #1 and #3 siCITED1 treated A2058 and A375 cells (Fig. S4b,c).

Owing to the previously reported dependency of MITF-induced cell cycle arrest on CDKN1A/P21 we investigated the levels of several cyclin-dependant kinase inhibitors following CITED1 silencing(Carreira et al. 2005). We found that CDKN1A/P21 was transiently increased in siCITED1 treated HT144 cells relative to the siNEG treated HT144 cells. In contrast, in A2058 cells, which do not have detectable levels of CDKN1A/P21 (Fig. S5), the levels of CDKN1C/P57 were suppressed in response to CITED1 overexpression (Fig 5b). We hypothesised therefore that melanoma cells can utilise either CDKN1A/P21 or CDKN1C/P57 to mediate cell cycle arrest induced by MITF and
this is reflected in the expression levels of the alternate CDK inhibitors in different melanoma cell lines (Fig. S5).

In agreement with the cell cycle data, an Alamar Blue assay revealed a significant reduction in cell metabolic activity over 5 days in HT144 cells treated with siCITED1 (Fig 5c). The effect was apparent but not as pronounced in the #3 siCITED1 sample.

The effect of CITED1 silencing on MITF is transient and mediated via promoter activation

We observed that the peak upregulation of MITF and CDKN1A/P21 protein following siCITED1 treatment varied from transfection to transfection, being seen between 24-48 hours post-transfection but appearing as unchanged or even downregulated after this time (Fig. 6a, upper panel). In agreement, later timepoints of the cell cycle analysis (=/>72 hours) exhibited little or no change in G1/S-phase distribution or even a reverse pattern (Fig. 5a HT144, and data not shown: A2058, A375). We therefore sought to examine the transcriptional dynamics more closely, map the changes in MITF following CITED1 silencing and see if they corresponded to cell behaviour and changes at the protein level. We used a quantitative droplet digital PCR based assay (Biorad) to measure mRNA in HT144 cells transfected with siCITED1#1 and siNEG as well as A2058 cells transiently overexpressing CITED1 compared to an empty vector control. MITF, CITED1 and IPO8 specific primers and probes were used to measure exact copies/µl of each mRNA from aliquots of the
same cDNA solution. Plots of siCITED1(copies/µl)/siNEG(copies/µl) and EV(copies/µl)/pCITED1(copies/µl) show the directional change in MITF and CITED1 relative to the housekeeper IPO8. CITED1 expression is rapidly suppressed following siCITED1 treatment of HT144 cells, concomitant with an upregulation of MITF that diminishes over time and in fact is suppressed by 100 hours in accordance with observations at the protein level (Fig 6a, lower panel). In contrast, overexpression of CITED1 in A2058 cells results in transient suppression of MITF at both protein and transcript level (Fig. 6b, upper and lower panels).

The rapid MITF transcriptional response to CITED1 manipulation suggested to us that the effect could be directly mediated at the promoter level. To test this hypothesis, we over expressed an MITF-M promoter-reporter construct and CITED1 in A375 cells. We chose A375 cells, as while they had less endogenous CITED1 and MITF than HT144 or A2058 so as not to cause interference with the assay, we also knew that they could respond adequately as they had an identical G1 accumulation/S-phase decrease to both HT144 and A2058 cells following CITED1 silencing (Fig S4c). TGFβ treatment was used as a positive control for repression of the MITF-M promoter. CITED1 transfection led to significant suppression of the MITF-M promoter luciferase activity relative to the empty vector control, as did TGFβ treatment alone or combination with CITED1 overexpression (Fig. 6c). There did not appear to be an additive or synergistic effect using both TGFβ
treatment and CITED1 overexpression suggesting TGFβ may be dependent on CITED1 for MITF suppression.

The CITED1-silenced gene signature predicts outcome in primary melanoma

The ‘proliferative’ and ‘invasive’ signature phenotypes have served to define the gene expression classification of melanoma cell lines. However, primary tumours and metastatic lesions have also been molecularly classified into several distinct groups by gene expression profiling (Harbst et al. 2012; Jonsson et al. 2010). The four-class structure found in tumours consists of the ‘pigmentation’, ‘proliferative’, ‘high-immune’ and ‘normal-like’ subgroups with a subset falling into an unclassifiable cohort (Jonsson et al. 2010). We used the same tumour classification to subtype the 120 cell lines that had publically available expression data and could show that the tumour ‘pigmentation’ subgroup that highly expresses MITF, corresponds to the cell line ‘proliferative’ phenotype described by Hoek et al. Accordingly, the tumour ‘proliferative’ and ‘high-immune’ subgroups comprise the cell line ‘invasive’ phenotype (Fig. 7a). It is worth noting that the names of the tumour subgroups were derived from a description of the differentially expressed genes that comprised each molecular classification while the ‘invasive-proliferative’ switching phenotypes were named to reflect the behaviour exhibited by lines classified by one or other signature. This distinction helps to explain the confusing occurrence that both classifications have a group referred to as ‘proliferative’ although they are not equivalent.
The overlap between the primary tumour classifying and cell line classifying systems allows us to infer that CITED1 expression is most likely restricted to a subset of MITF high ‘pigmentation’ subtype tumours. As the tumour subtype classification was shown to be prognostically significant in primary melanomas we were interested to know if CITED1 expression itself was independently predictive of outcome. Previously we reported on the analysis of 223 primary lesions using the Illumina WG-DASL protocol (Harbst et al. 2012). As the CITED1 probe in this assay did not produce reliable data we instead derived a CITED1-silenced gene signature score based on the differentially expressed genes from the HT144 siCITED1 experiment (Fig. 3). We subsequently interrogated the gene expression data on the primary melanoma lesions using a nearest centroid approach derived from the CITED1-silenced gene signature. This revealed that primary melanomas with a gene expression signature most similar to the CITED1-silenced signature (CITED1low-class) had a significantly better outcome than those with a signature most disparate from the CITED1-silenced signature (CITED1high-class) (Fig. 7b). Importantly, the CITED1 signature classing had independent prognostic information (HR 1.85, CI 0.30-0.98, p=0.044) from the AJCC staging system (HR 5.05, CI 2.42-10.55, p=1.64x10^-5). These data indirectly imply that CITED1 expression itself is a potential prognostic indicator in primary melanomas and the transcriptional program influenced by CITED1 expression determines tumour behaviour in vivo.
DISCUSSION

One seemingly paradoxical observation from our study and previous investigations is that although CITED1 behaves as a negative regulator of MITF, both their expression levels appear positively correlated across cells lines and tumours. We maintain that this observation simply reflects the fact that where there are high levels of MITF, high levels of its negative regulator are also required. The evidence of the tight control exerted over MITF levels in melanocytes and melanoma simply speaks to the necessity of the cell to maintain a level compatible with survival and proliferation, in a type of biological ‘sweet-spot’ facilitating tumour progression. The cellular effects of both extremes i.e: very low or high levels of MITF, have been elegantly described by a rheostat model in order to reconcile the conflicting observations of the effects of manipulating MITF in vitro, and the fact that counter-intuitively, a lineage-specifying differentiation factor can behave as a potent oncogene(Hoek & Goding 2010; Carreira et al. 2006; Cheli, Giuliano, et al. 2011a). The rheostat model (Fig. S6) explains why MITF silencing can block cells in G1 and induce senescence, while it is also possible to induce a G1 arrest by MITF overexpression via CDKN2A/P16 or CDKN1A/P21 and, as we now propose, potentially also via CDKN1C/P57 (Carreira et al. 2006; Loercher et al. 2005; Carreira et al. 2005). At the extreme high end of MITF expression lies differentiated melanocytic cells, while the lowest levels can lead to senescence and irreversible cell death. Between these two extremes however it is thought that melanoma cells can oscillate from a low-MITF ‘invasive’ to a high-MITF ‘proliferative’ state via phenotype-switching.
We hypothesise that the role of CITED1 in melanoma is to maintain levels of MITF compatible with tumour progression and effectively tip the balance in favour of cell cycle progression rather than MITF-induced G1-arrest. This is supported by our findings that downregulation of CITED1 using siRNA results in a phenotype switch to a more pigmented state driven by increased MITF expression and concomitant upregulation of CDKN1A/P21. Conversely, we could observe that downregulation of MITF resulted in suppression of CITED1 in several cell lines suggesting the existence of a classical feedback loop where low MITF levels result in inhibition of its negative regulator. MITF induced cell cycle arrest was previously shown to be dependent on CDKN1A/P21 and it was demonstrated that MITF does not cause a cell cycle arrest in CDKN1A-deficient mouse embryo fibroblasts (MEF) cells (Carreira et al. 2005). However, our data indicate that in melanoma cells deficient in CDKN1A/P21, the alternative CDK inhibitor CDKN1C/P57 is expressed and responsive to MITF.

Interestingly, while we observed upregulation of most MITF targets following CITED1 silencing, we found that BRCA1 and other DNA damage response (DDR) genes were suppressed, suggesting that CITED1 downregulation does not necessarily facilitate transcription of all MITF targets. It is thus tempting to speculate that rather than simply acting to induce MITF and thereby indirectly enhance transcription of its target genes, that CITED1 may also act as co-factor for MITF at various genomic locations differentially modulating
the MITF target gene response at individual promoters. One way that this might be achieved is via MITF-CITED1 competition for CBP/P300 binding as CBP/P300 is a known transcriptional coregulator for MITF, although it is not required for transcription of all MITF targets (Vachtenheim et al. 2007) (Yan et al. 2013).

As suggested by Sáez-Ayala et al., anti–cancer therapy should be ideally independent of dominant or ‘driver’ genetic alterations so that subclonal populations do not gain a subsequent advantage and the same holds true in the case of targeting a specific phenotype. Successful therapy will necessarily need to switch or push the subdominant phenotype into the susceptible state or eradicate the phenotype resistant to treatment. This approach was initially championed by Cheli et al., who proposed the eradication of low-MITF cells as a therapeutic strategy (Cheli, Guiliano, et al. 2011b). Indeed the idea of lineage-specific therapy has been subsequently proved in principle using methotrexate (MTX) to first activate MITF expression, in turn activating the tyrosinase enzyme, and thereby sensitising tumour cells to a tyrosinase-processed anti-folate prodrug (TMECG) (Sáez-Ayala et al. 2013). However, even without drug targeting, induction of MITF, above what is tolerated by even the highly pigmented tumor cell types, i.e: levels reaching that of melanocytes, would seem to be incompatible with melanoma progression as it inhibits cell cycle progression. Our assertion is that CITED1 acts to repress MITF in order to maintain its level in a range compatible with tumorigenesis. This assertion as a consequence naturally
suggests CITED1 as therapeutic target for genetic manipulation. Successful implementation of such a strategy would result in cell specific enhancement of MITF expression and increased susceptibility to the type of chemotherapeutic eradication demonstrated by Sáez-Ayala et al. or potentially induction of CDKN1A/p21 or CDKN1C/p57-dependent cell growth arrest even without further intervention (Fig. S6) (Sáez-Ayala et al. 2013).

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Figure 1

TGFβ induced gene expression in A2058 melanoma cells

(a) Distribution of the proliferative and invasive signature score genes relative to the heat map of gene expression changes induced by TGFβ treatment. (b) Gene expression heatmap of the proliferative and invasive signature within those genes significantly altered by TGFβ treatment (1009 transcripts following SAM, median FDR q-value=1%. ‘2006’ refers to the signature list (motif1 and motif2) while ‘2012’ refers to the updated signature derived from further datasets (Widmer et al. 2012). (c) Western blot of MITF and CITED showing both proteins are suppressed by TGFβ treatment. β-Actin is used as a loading control.
Figure 2

CITED1 expression correlates with MITF expression

(a) The relative MITF and CITED1 expression levels from the publically available gene expression data of 120 melanoma cell lines (Pearson correlation $r=0.6543$, $p < 0.001$). Arrows indicate the cell lines used in this study. The cell lines are further subdivided into one of either ‘invasive’ or ‘proliferative’ phenotype based on expression signature score. (b) A Western blot is shown of the relative protein expression levels of both MITF and CITED1 in our cell lines in good agreement with the transcript levels.
Figure 3

CITED1 silencing induces a phenotype switch

(a) 120 melanoma cell lines are shown distributed on the basis of the phenotype score. The HT144 cells chosen to study the effects of CITED1 downregulation are indicated. (b) Following CITED1 downregulation a phenotype shift is observed indicated by their scatter position change according to the average expression score of genes that distinguish invasive from proliferative phenotype. For the 120 melanoma cell lines (Affymetrix platform) the expression score was derived from expression levels of 50 and 54 proliferative and invasive genes with matching genes symbols, respectively, while for the HT144 experiment (Illumina platform), 51 and 54 proliferative and invasive genes with matching genes symbols were retrieved. A Western blot of the degree of protein downregulation of CITED1 at the time of the expression analysis is also shown. β-Actin is used as a loading control (inset). (c) A heatmap comprising the ‘invasive’ and ‘proliferative’ signature genes illustrating how they are altered by CITED1 silencing; #1 and #3 denote two separate siRNAs targeting CITED1.
figure 3
Figure 4

CITED1 regulates MITF and its targets genes

(a) A heatmap showing the 312 transcripts identified as significantly changed using a SAM 2-way comparison between siNEG and siCITED1 (#1 & #3 were combined), median FDR q-value = 3%). MITF, as well as a cohort of significantly enriched MITF targets, genes associated with pigmentation, and genes involved in the UV/DNA damage response are highlighted on the right. (b) Western blot confirmation of the effect of silencing CITED1, using siRNA (#1, #3) relative to a negative control siRNA (N), on MITF protein expression in HT144 cells at 24 and 48 hours post-transfection, and the effect of overexpressing CITED1 (pCITED1) relative to an empty vector control (EV) in A2058 cells at 24 hours post-transfection. β-Actin is used as a loading control in each case. (c) Western blot showing the effect of silencing MITF using two siRNAs (siM1, siM3) on both MITF and CITED1 levels in HT144 cells at 48 hours post-transfection relative to a negative control siRNA (N). β-Actin is used as a loading control.
figure 4
Figure 5

CITED1 silencing restrains cell cycle progression and reduces cell viability

(a) A bar chart showing the % change in cell cycle distribution in #1 siCITED1 treated HT144 cells relative to siNEG treated HT144 cells. The reduction in the total S-phase is shown at 33 hours, 48 hours and 72 hours post-transfection in addition to the corresponding increase in the diploid G1 fraction. (b) Western blots showing upregulation of CDKN1A/P21 following CITED1 silencing in HT144 cells and suppression of CDKN1C/P57 following CITED1 overexpression in A2058 cells. (c) An Alamar Blue based metabolic assay shows a reduction in cell viability over 5 days in HT144 cells treated with siCITED1 relative to those treated with siNEG. Stars indicate significance for siNEG vs. #1 siCITED1 where ***p<= 0.0005, **p<= 0.005 and *p<= 0.05. In the case of siNEG vs. #1 siCITED1, the difference is significant (*) at 96 and 120 hours.
figure 5
Figure 6

CITED1 silencing transiently upregulates MITF via promoter activation

(a) A Western blot of HT144 cell lysate samples taken at the indicated time points post-transfection and showing the corresponding levels of MITF protein in #1 siCITED1 and siNEG treated cells. Shown underneath are the changes in mRNA levels of MITF-M, CITED1 and a housekeeper gene IPO8, as measured by specific ddPCR assays over a time course of 4-100 hours following transfection of HT144 cells with either siCITED1 or siNEG. (b) A Western blot of A2058 cell lysate samples taken at the indicated time points post-transfection and showing the corresponding levels of MITF protein between CITED1 overexpression (pCITED1) and empty vector (EV) control. Shown underneath are the changes in mRNA levels of MITF and a housekeeper gene IPO8, as measured by specific ddPCR assays over a time course of 33-72 hours following transfection of A2058 with either pCITED1 or an empty vector control. (c) The relative luciferase activity of the MITF-M promoter reporter measured in lysates of A375 cells transfected with the pCITED1 expression plasmid or empty vector (EV) control and treated with or without TGFβ for 24 hours (**p<= 0.0005, **p<= 0.005 and NS= not significant).
Figure 6

(a) Western blot analysis showing expression levels of CITED1, MITF, CDKN1A/P21, and β-ACTIN at 48, 72, 100, and 120 hours.

(b) Graphs showing the expression ratio of CITED1 and MITF over time.

(c) Bar graph illustrating the percentage of relative luciferase activity for different treatments: empty vector, empty vector + TGFβ, pCITED1, and pCITED1 + TGFβ.
Figure 7

The CITED1-silenced gene signature predicts patient outcome

(a) In the leftmost panel a scatter plot of the 120 melanoma cell lines are shown distributed on the basis of their ‘invasive’ or ‘proliferative’ phenotype signature score and coloured according to the tumor molecular subtypes as defined by Jönsson et al. to illustrate the overlap between the two classification systems (Jonsson et al. 2010). In the rightmost panel the data is presented as a heat map where each gene of the proliferative or invasive signature genes is represented by a horizontal line and the 120 individual cell lines are grouped by molecular tumour subtype (coloured blocks) and shown vertically. (b) Recurrence free survival (RFS) of primary melanoma patients grouped by gene expression similarity to the CITED1 (siCITED1) silenced gene signature.
**Figure 7**

(a) Heatmaps illustrating the proliferation and invasion scores for various cell lines.

(b) Kaplan-Meier survival curves showing the difference in relapse-free survival between low and high CITED1 classes.