

# Insights into p53 transcriptional function via genome-wide chromatin occupancy and gene expression analysis

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The tumor-suppressor p53 can induce various biological responses. Yet, it is not clear whether it is p53 *in vivo* promoter selectivity that triggers different transcription programs leading to different outcomes. Our analysis of genome-wide chromatin occupancy by p53 using chromatin immunoprecipitation (ChIP)-seq revealed ‘p53 default program’, that is, the pattern of major p53-bound sites that is similar upon p53 activation by nutlin3a, reactivation of p53 and induction of tumor cell apoptosis (RITA) or 5-fluorouracil in breast cancer cells, despite different biological outcomes. Parallel analysis of gene expression allowed identification of 280 novel p53 target genes, including p53-repressed *AURKA*. We identified Sp1 as one of the p53 modulators, which confer specificity to p53-mediated transcriptional response upon RITA. Further, we found that STAT3 antagonizes p53-mediated repression of a subset of genes, including *AURKA*.

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Given the high tumor-suppressor potency of the p53 transcription factor, the strategy to reestablish p53 function is very attractive.<sup>1</sup> However, there remains much to learn, as the cellular response to p53 can range from cell death to cell survival, and the consequences of p53 reactivation are difficult to predict.<sup>2</sup>

Several classes of molecules reactivating p53, such as nutlin3a, reactivation of p53 and induction of tumor cell apoptosis (RITA), PRIMA-1<sup>MET</sup>/APR-246, MI-219 and tenovins, have been shown to efficiently eliminate tumor cells *in vitro* and in animal models.<sup>3–7</sup> Nutlin3a binds the major p53 inhibitor Mdm2, entailing the induction of p53.<sup>3</sup> In contrast, RITA prevents Mdm2 binding and activates p53 by binding to p53 itself.<sup>4</sup> Notably, p53 activation by nutlin3a and RITA in the same cancer cells triggers different transcriptional programs, resulting in growth arrest or apoptosis.<sup>8</sup>

The control of cell fate by p53 is accomplished by activation or repression of p53-target genes, containing the consensus p53-binding motif.<sup>9</sup> However, distinct transcriptional programs triggered by p53 upon different stresses raise the question whether this is due to a differential pattern of p53 binding to promoters, or whether p53 binds the same set of its targets irrespective of the type of stimuli, while the presence or

absence of other factors will decide whether the expression of a particular gene will be regulated by p53.<sup>10</sup> Genome-wide analyses are required to delve into the mechanisms by which it triggers different transcriptional programs in different settings. Previous genome-wide studies of p53 DNA binding *in vivo* using chromatin immunoprecipitation (ChIP) in conjunction with hybridization (ChIP-chip) or ChIP paired-end (ChIP-PET) sequencing approaches helped to refine the p53 consensus binding site and to predict a number of p53-target genes.<sup>11–13</sup>

In this study, we carried out an unbiased analysis of chromatin occupancy by p53 using p53 ChIP followed by deep sequencing (ChIP-seq) upon its induction by three different molecules, target-specific nutlin3a and RITA, as well as chemotherapeutic drug 5-fluorouracil (5-FU) and combined it with the analysis of a new comprehensive data set of p53-induced gene expression changes. We characterized important features of p53 *in vivo* chromatin occupancy and identified novel target genes and transcription factors, which affect p53 transcriptional function, as exemplified by the modulation of p53 transcriptional response by Sp1 and the antagonistic interaction between p53 and STAT3 at the *AURKA* promoter.

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**Abbreviations:** RITA, reactivation of p53 and induction of tumor cell apoptosis; 5-FU, 5-fluorouracil; qPCR, quantitative real-time reverse transcription PCR; static, STAT three inhibitory compound; ChIP, chromatin immunoprecipitation; ChIP-seq, ChIP combined with deep sequencing; ChIP-PET, ChIP paired-end; ChIP-chip, ChIP in conjunction with chip hybridization; ChIP-PCR, ChIP combined with real-time reverse transcription PCR; TSS, transcription start site; TF, transcription factor; RE, response element

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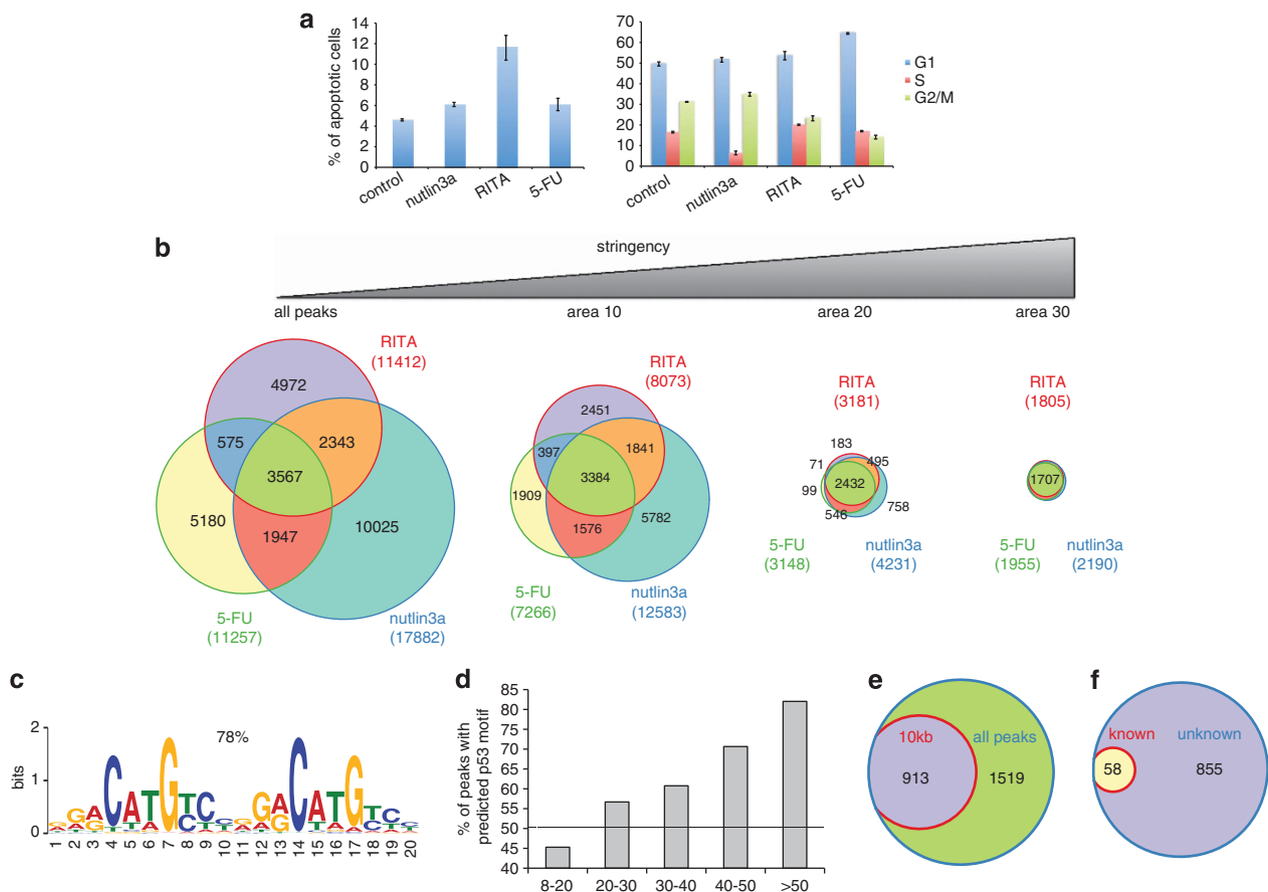
## Results

**Genome-wide map of p53 binding to chromatin upon its induction by nutlin3a, RITA and 5-FU.** We compared the genome-wide chromatin occupancy by p53 upon its activation with three different compounds, non-genotoxic molecules nutlin3a and RITA, as well as chemotherapeutic drug 5-FU, known to induce DNA damage.<sup>14</sup> Despite mechanistic differences, all the three treatments induce p53 protein levels and its transcriptional activity.<sup>3,8,15</sup> Intriguingly, they lead to different biological responses: depletion of S-phase indicating growth arrest upon nutlin3a treatment, mainly G1/S arrest induced by 5-FU and induction of apoptosis as a major response to RITA (Figure 1a).

We performed the p53 ChIP followed by massive parallel DNA sequencing (ChIP-seq) in MCF7 cells treated with nutlin3a, RITA and 5-FU. Applying a threshold of  $P < 0.05$ , we identified 6357 peaks corresponding to p53-bound DNA fragments in the library from mock-treated control cells, and 17882 in nutlin3a-, 11412 in RITA- and 11257 in 5-FU-treated samples (Figure 1b, Supplementary Tables S1–4).

Comparison of the chromatin occupancy by p53 upon different treatments using the web-based tool Galaxy Genome<sup>16</sup> revealed a partial overlap between the data sets (Figure 1b). In order to select the most relevant binding sites, we selected peaks with a minimum of two-fold enrichment over IgG control and large area, which correlated with peak height in a linear fashion (Supplementary Figure S1A). The increase of stringency (area 30) resulted in a significant increase of the proportion of peaks common to all three treatments, for example, for nutlin3a, the proportion of common peaks increased from 20 to 78% (Figure 1b). These results indicate that p53 binds the same set of sites, defined as a ‘default p53 program’ independent from the type of p53 activating stimuli.

Our ChIP combined with real-time reverse transcription PCR (ChIP-PCR) analysis confirmed p53 binding only to a half of randomly selected sites from ‘area 10’ (data not shown), suggesting that these are low confidence peaks. Therefore, for further analysis, we selected all p53-bound loci, which reached the threshold area 20. For these high confidence peaks, the proportion of those common for all



**Figure 1** Characterization of the global chromatin occupancy by p53 upon its activation by nutlin3a, RITA and 5-FU. (a) Induction of apoptosis upon nutlin3a, RITA and 5-FU treatment of MCF7 cells was detected by fluorescence-activated cell sorting (FACS) of annexin V-stained cells (left) and cell-cycle profiles were assessed by FACS of PI-stained cells (right). (b) Venn diagrams were obtained by intersection of all p53-bound DNA fragments (ChIP-Seq peaks,  $P < 0.05$ ) obtained from MCF7 cells treated with nutlin3a-, 5-FU- and RITA and filtering according to the area. (c) The p53 consensus binding motif was identified *de novo* by analyzing the sequences of 500 randomly selected peaks common for all the three treatments using the program MEME (d). The fraction of peaks containing the p53 consensus site increased along with increased stringency of peak selection. (e) A Venn diagram demonstrates the proportion of p53-bound fragments located within  $\pm 10$  kb of the TSS (913 out of 2432). (f) A Venn diagram shows the proportion of peaks located in the vicinity of known and unknown p53-target genes bound within  $\pm 10$  kb of TSS upon all three treatments. See also Supplementary Figure S1, Supplementary Figure S2 and Supplementary Tables S1–S4

the three treatments (a total of 2432 peaks) became predominant, although we detected a certain number of unique peaks (area 20, Figure 1b). However, ChIP-PCR analysis did not confirm their specificity: no unique peaks were found among randomly selected top-score nutlin3a-only peaks, whereas merely 40% of RITA-only peaks were confirmed as being specific (data not shown). Moreover, microarray analysis showed that the expression of the majority of genes, which appear to be selectively bound and significantly changed upon RITA treatment, was also affected by nutlin3a (24 out of 30 genes, Supplemental Table 12). In addition, as indicated in Supplementary Figure S1B, only 10% of peaks detected upon nutlin3a and less than a half of RITA-specific peaks contain p53 consensus motif. In contrast, 82% of peaks detected upon all three conditions (i.e., common peaks) contain p53 consensus.

Thus, we conclude that the majority of high-confidence binding sites are common for all the three treatments and represent the 'p53 default program'.

**In vivo p53-binding sequence preference.** We generated a consensus p53-binding site based on p53-bound fragments common for all the three treatments at threshold area 20 (Figure 1b) by applying the *de novo* motif discovery algorithm MEME for 500 randomly selected common peaks (out of 2432,  $E = 4.4 \times 10^{-682}$ ) (Figure 1c). This motif fits well to the original p53 consensus site as well as the one refined recently.<sup>11,12,17,18</sup> Interestingly, applying a higher threshold for the peak area increased the proportion of peaks containing the p53-binding site from 45 to 82%, as detected by the p53Scan algorithm (Figure 1d), whereas only 3.9% of random sequences contain this motif. Thus, most of the 'default p53 program' sites contained canonical p53-binding sites.

In agreement with previous report,<sup>12</sup> we found that 37% of the peaks are located within 10 kb from the transcription start sites (TSSs, Figure 1e and Supplementary Figure S2).

**Identification of novel p53-target genes.** Analysis of p53 occupancy at promoters showed that, along with a number of known p53 targets, p53 bound to 855 sites corresponding to 688 novel genes (Figure 1f). Comparison of gene-expression profiles with our ChIP-seq data showed that 41% of these (280 genes) were differently expressed upon nutlin3a (Figures 2a and b, see Supplemental Data and Supplementary Tables S5–S9 for a more detailed description). These genes are likely to be novel functional p53-target genes (Supplementary Table S10).

A total of 4484 genes that changed their expression upon nutlin3a, but were not bound by p53 in the vicinity of TSS, most likely represent the secondary effects of p53. In addition, we detected the p53 binding to the promoters of 408 genes, which did not result in expression changes (Figure 2a and Supplementary Tables S5–S9). Thus, our data suggest that the binding of p53 *per se* is not sufficient to regulate gene expression.

**Comparison of p53-bound and induced versus p53-bound and repressed genes.** Among 280 novel p53-target genes, 214 were induced and 66 were repressed (Figure 2b).

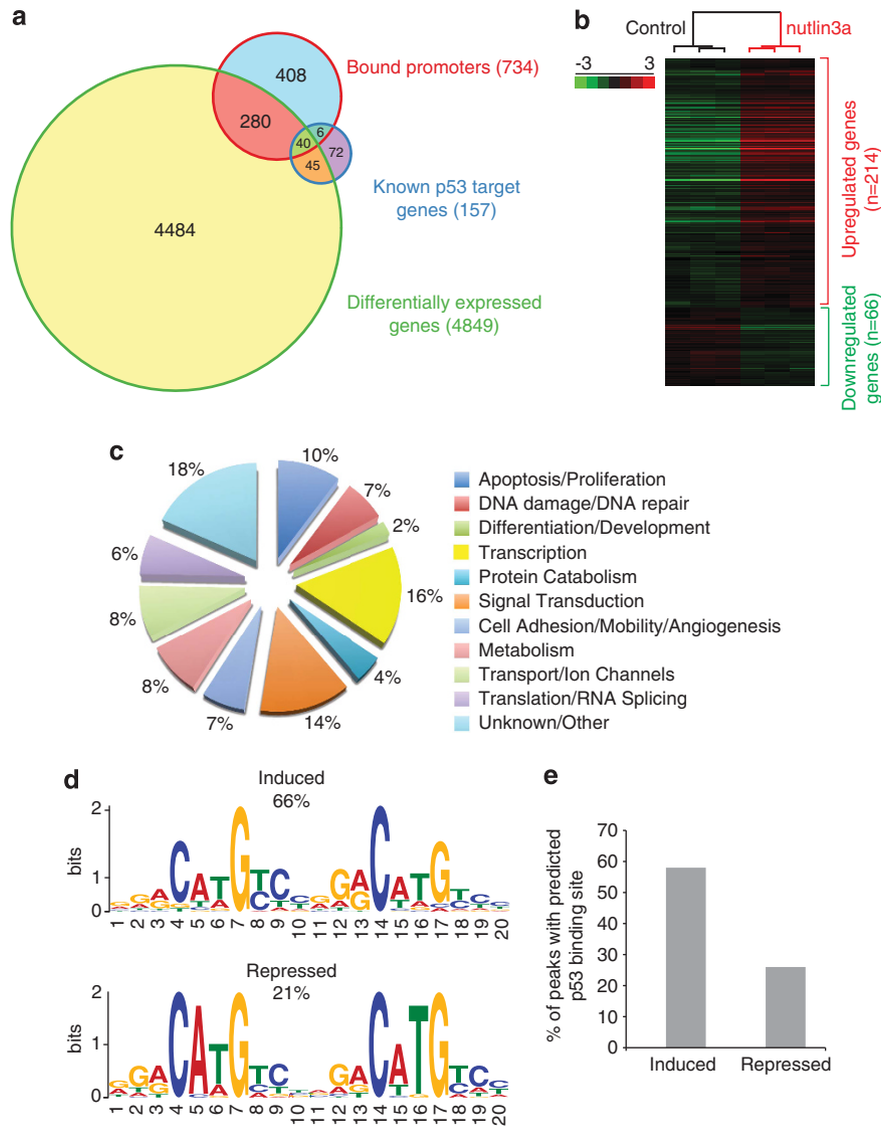
Using gene ontology, we functionally annotated these genes (Figure 2c, Supplementary Table S10). Along with expected categories, such as apoptosis, cell-cycle regulation, DNA repair/damage, signal transduction, metabolism, transport/ion channels, protein catabolism and translation/splicing, we found that one of the most highly represented categories involved genes encoding regulators of transcription, mainly transcription factors (TFs) (e.g., E2F7, HES1, HES2, FOSL1, ZNF219, Sp1). This might help to explain the gross changes in gene expression upon p53 activation, that is, the differential expression of 4484 genes after nutlin3a treatment.

We searched for the p53-binding motif in our set of p53-target genes and applied the *de novo* discovery algorithm MEME to the p53-bound fragments separately for induced and repressed genes, which resulted in two distinct motifs, both of which are similar to the p53 consensus motif. However, although this motif was present in 66% of induced genes (E-value of  $10^{-244}$ ), the proportion of repressed genes, which contained the p53 consensus was much lower, only 21% (E-value reached  $5.3 \times 10^1$ ) (Figure 2d). This conclusion was confirmed using p53Scan and p53MH programs (Figure 2e and Supplementary Figure S5B). Interestingly, the extent of occupation by p53 was significantly higher in induced than in repressed genes (Supplementary Figure S5A).

Taken together, our data implies that the majority of p53-bound genes, which are induced, share the classical consensus motif and have more pronounced p53 binding than the repressed genes. The consensus motif generated from repressed targets (Figure 2d, lower panel) fits with the classical consensus site even better, but is infrequently present. Interestingly, analysis of p53 occupancy in control and treated samples allowed to identify two classes of genes bound by p53 in the absence of activating stimuli (Supplementary Figures S3 and S4).

**Analysis of composite response elements (REs) in the promoters of p53-repressed genes revealed two possible modes of repression by p53.** We performed composite pattern search for TF-binding sites adjacent to sites occupied by p53 (distance < 100bp) in the promoters of repressed genes using geneXplain platform.<sup>19</sup> Top-score TF include E2F, MYB, HNF3, DEAF-1, HSF-2, IRF, STAT and others (Table 1). E2F1 is known to enhance apoptosis induced by p53 and cooperate with p53 in activation of several genes; moreover, E2F and p53 have been shown to repress two anti-apoptotic genes *Bcl-2* and *Mcl-1*.<sup>20</sup> Further, E2F1 is activated by nutlin3a and its transcriptional activity is a crucial determinant of suppressor effect of nutlin3a.<sup>21</sup> Enrichment of E2F RE next to p53-binding sites in repressed genes suggests that the cooperation between these two factors in transcriptional repression via direct binding of both TFs to promoters might occur on a larger scale than previously anticipated.

Apart from acting together in induction of gene expression, IRF and p53 have been shown to cooperate in repression of hTERT gene.<sup>22</sup> Taken together with our data, this provides a support to the notion that p53 and IRF can collaborate in repression of at least several genes.



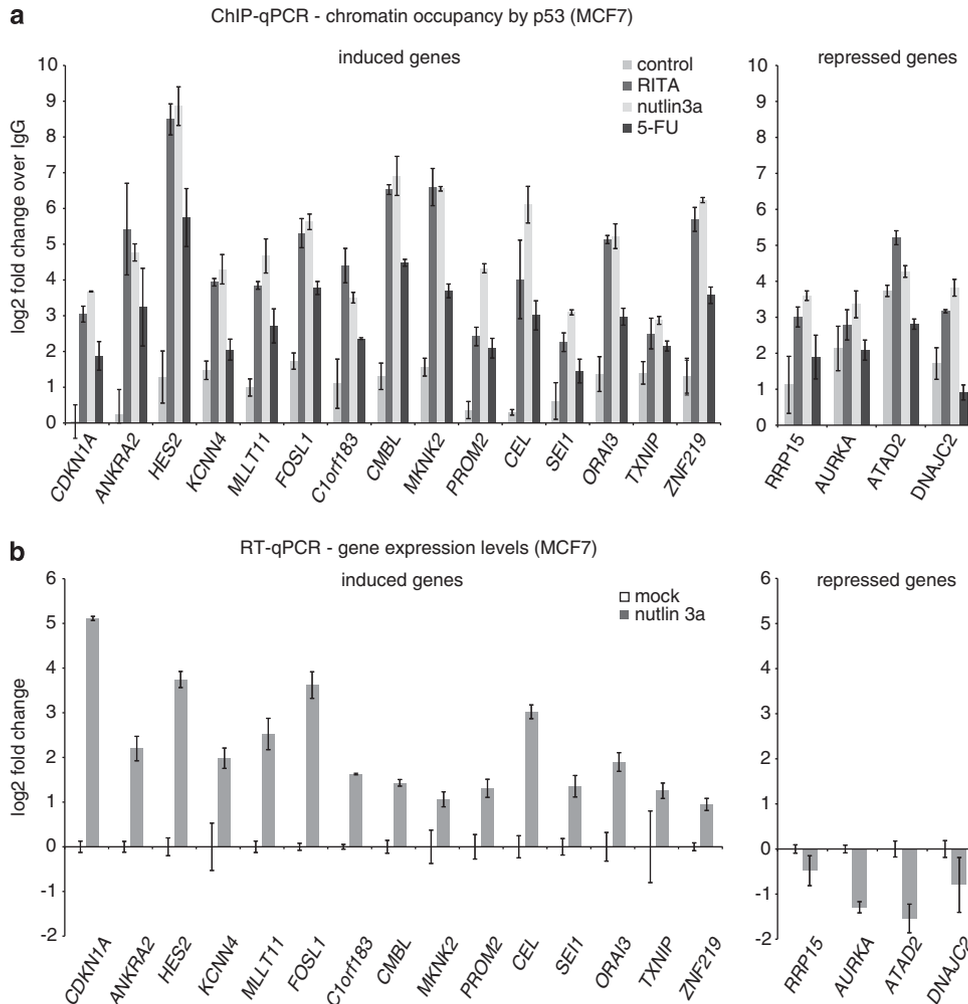
**Figure 2** Novel p53-target genes bound by p53 and differentially expressed upon nutlin3a treatment. (a) Bioinformatic analysis of the genes bound by p53 within 10 kbp from TSS and differentially expressed upon 8 h of nutlin3a treatment ( $P < 0.05$ ) revealed 280 putative novel p53-target genes. (b) A heatmap shows the differential expression of selected 280 p53-bound genes upon nutlin3a treatment. Columns indicate separate arrays and rows indicate genes. The values are row-mean-centered. Green corresponds to lower and red to higher than mean value expression. (c) The pie chart of the functional categories of 280 novel p53-target genes. (d) The p53 consensus binding site was detected much more often in the proximity of TSS of induced, than repressed genes (66% and 21%, respectively), as analyzed using the MEME algorithm. (e) The probability of the presence of a p53 consensus binding motif in the p53-bound fragment is higher in induced than in repressed genes

**Table 1** Composite regulatory elements in p53-repressed promoters

Composite elements	Frequency in peaks (fP)	Frequency in background (fB)	Ratio	P-value
P53-E2F	0.23849	0.029404	8.1108	5.9188E-09
P53-MYB	0.29784	0.046216	6.44452	3.9883E-07
P53-GATA4	0.790771	0.588456	1.34381	2.3550E-02
P53-HNF3	0.304779	0.101922	2.99032	1.5959E-04
P53-DEAF-1	0.48582	0.179621	2.70469	5.5473E-06
P53-HSF-2	0.310844	0.085733	3.62572	1.1939E-05
P53-STAT	0.239011	0.024201	9.87608	7.9175E-09
P53-IRF	0.093085	0.00199	46.7764	3.6570E-06
P53-TBX5	0.123387	0.005495	22.4544	7.5897E-07

Several top-score TF, which have their RE in p53-repressed genes are proto-oncogenes, including MYB, HNF3 (FOXA), DEAF-1, HSF-2 and STAT (Table 1). It is tempting to speculate that p53 might mediate transrepression via inhibition of these TF at their target promoters. Indeed, it has been found that p53 can antagonize MYB by binding to MYB-activated promoter<sup>23</sup> and to compete out HNF3(FOXA) from its promoters.<sup>24</sup> We addressed the question whether there is an antagonistic interaction between p53 and STAT3 at the promoters in a series of experiments presented below.

Thus, analysis of TF RE overrepresented in p53-bound and repressed promoters revealed two possible modes of p53-mediated repression: via cooperation with factors, such



**Figure 3** Validation of chromatin occupancy and mRNA expression of a selected panel of newly identified p53-target genes. (a) ChIP using the anti-p53 antibody DO-1, followed by qPCR, was used to detect relative p53 occupancy in the promoters of 18 selected genes in MCF7 cells treated with nutlin3a, RITA and 5-FU for 8 h. CDKN1A serves as a positive control for p53 binding. (b) The mRNA levels of 18 selected genes were detected by qPCR in MCF7 cells treated with nutlin3a

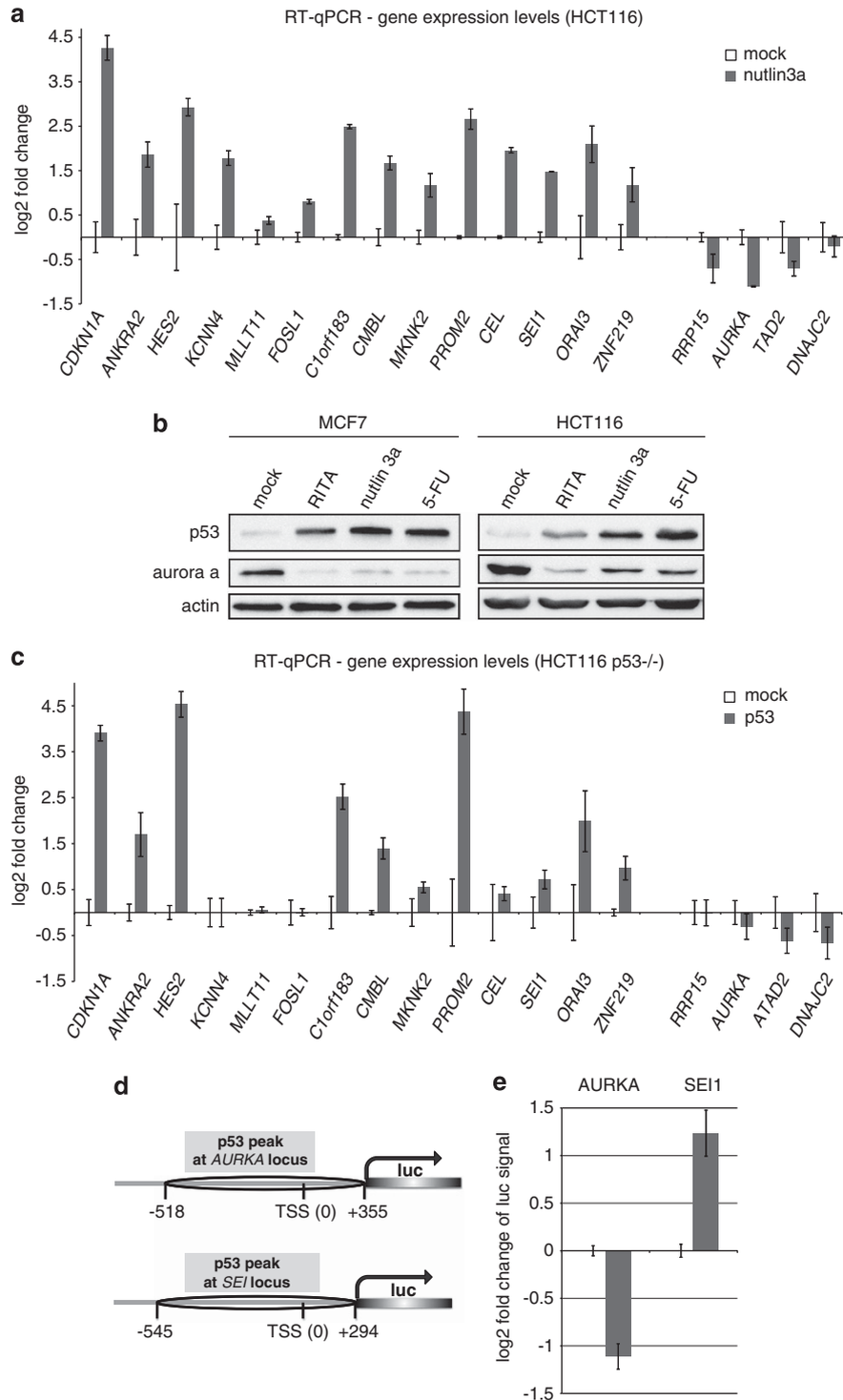
as E2F and IRF, and inhibition of activity of proto-oncogenes, such as MYB, HNF3 and STAT, at their target promoters.

**Validation of a set of novel p53-target genes.** Assessment of p53 binding to a set of 14 activated and 4 repressed novel p53 targets by ChIP-PCR, as well as their differential expression using quantitative PCR (qPCR), both in MCF7 and HCT116 cells, confirmed that the selected genes are novel p53-target genes (Figures 3a and b and Figure 4a, respectively). In addition, ectopic expression of p53 in the p53-null HCT116 *TP53*<sup>-/-</sup> led to the induction of 10 out of 14 genes (Figure 4c).

One of the most interesting novel p53 targets is *AURKA*, encoding aurora kinase A, a promising target for the development of novel anticancer therapies.<sup>25</sup> In line with mRNA decline as detected by qPCR (Figures 3b and 4a), the level of aurora kinase A protein was strongly decreased in MCF7 and HCT116 cells upon p53 induction by nutlin3a, RITA and 5-FU (Figure 4b).

A more detailed investigation of the role of the p53-binding site in the promoters of *SEI1*, as a representative of a p53-induced gene, and *AURKA*, representing a p53-repressed gene, using luciferase reporter assay confirmed the functionality of p53 RE in these genes (Figures 4d and e). Taken together, our results provide a solid evidence that *AURKA* is directly repressed by p53, whereas *SEI1* is induced by p53.

**Transcription-factor-binding motif enrichment in p53-target genes.** Our finding of the p53 default program implies that differential gene expression upon p53 binding to its promoters is highly dependent on the presence of other TFs at p53-target promoters. In order to identify these factors, we searched for TFs, which have their binding motifs in the interval of  $\pm 100$  bp of the ChIP-seq peak maximum in p53-induced and -repressed genes. Using the Match algorithm from geneXplain, we found that RE for a number of TFs, such as Sp1, WT1, ZF5, ETS1 and AP2, as well as p53, were overrepresented in the promoters of both categories of genes



**Figure 4** Further validation of novel p53-target genes. (a) Colon carcinoma HCT116 cells were treated with nutlin3a and the level of expression of 18 novel p53-target genes was assessed by qPCR. (b) The protein level of aurora kinase A was decreased upon p53 activation by nutlin3a, RITA and 5-FU in MCF7, as well as HCT116 cells, as assessed by immunoblotting. (c) Changes of mRNA levels of 18 selected genes were assessed by qPCR upon overexpression of p53 in the p53 null HCT116 p53<sup>-/-</sup> cells. (d) Schemes depicting p53-bound fragments derived from the p53-induced *SEI1* and p53-repressed *AURKA* promoters, which were cloned into a luciferase reporter plasmid. (e) The activity of luciferase reporter expressed under the control of *AURKA* and *SEI1* promoters was assessed upon ectopic expression of p53 in HCT116 p53<sup>-/-</sup> cells

(Table 2). Notably, RE for several TFs were enriched only in induced genes, for example, NF- $\kappa$ B, SMAD3 and MAZ, whereas for others – only in repressed genes, for example,

HIF1 and BCL6 (Table 2). Expression of 18 out of 45 TFs, whose RE were enriched in the p53-regulated promoters, was affected upon p53 activation by nutlin3a treatment,

**Table 2** Transcription factors predicted to have their binding sites in the vicinity ( $\pm 100$  bp) of the p53-binding sites

Common for induced and repressed				Unique for induced				Unique for Repressed			
Factor	N sites per 1 kbp	Enrich	P-value	Factor	N sites per 1 kbp	Enrich	P-value	Factor	N sites per 1 kbp	Enrich	P-value
AHR	0.7117	1.8792	0.001829	BACH2	0.2491	2.587	0.0098112	<u>ARNT</u>	1.7143	2.811	3.59E-05
AP2	5.1246	3.1678	9.43E-40	ETV7	2.9181	1.3733	0.0006481	<u>BCL6</u>	2.0714	1.9207	0.0017116
AP4	1.2278	1.6777	0.0006008	HAND1	4.1815	1.494	7.40E-07	<b>DEAF1</b>	0.2143	33.38	0.0021042
AR	0.3559	2.1322	0.0095111	HNF1A	0.1779	3.9597	0.0049341	<u>HIF1A</u>	1.2143	2.6642	0.0007813
CP2	4.7865	1.6139	7.10E-10	<b>AP1</b>	0.4804	2.5806	0.0004129				
<u>CREB/ATF</u>	1.9217	1.7404	7.09E-06	<u>MAF</u>	0.1779	6.9295	0.0005729				
<b>E2F</b>	1.2278	3.4153	1.15E-11	<b>MAZ</b>	0.7117	2.174	0.0002464				
EBF	0.9431	2.2258	1.73E-05	MYOG	5.8541	1.416	2.89E-07				
<u>EGR1</u>	0.8363	3.7221	3.98E-09	<b>NFKB</b>	1.121	1.7289	0.0005755				
ETS1	1.3701	2.1558	6.09E-07	<u>NR3C1</u>	0.1068	16.631	0.0018787				
HIC1	0.4804	3.7419	7.46E-06	PAX6	8.2206	1.177	0.0020025				
NR5A2	1.1744	1.5373	0.0037999	<u>SMAD3</u>	4.6085	1.2729	0.0008693				
PAX3	5.8363	1.2877	0.0001103	SP3	4.2705	1.974	3.22E-15				
PAX4	1.2811	1.5234	0.0030804	ZBTB33	0.2491	2.985	0.0044078				
PAX5	6.9751	1.6266	5.52E-14								
PPARA	6.4057	1.2319	0.000642								
SP1	1.1566	5.4596	2.06E-16								
<u>STAT</u>	2.2143	1.8347	0.0022804								
<b>TEAD-2</b>	1.2989	3.0658	7.00E-11								
<b>TP53</b>	3.2562	8.0514	4.66E-55								
V-MYB	0.8541	2.2174	4.48E-05								
<b>VDR</b>	4.7687	1.6693	6.62E-11								
WT1	7.0463	2.3304	8.54E-34								
<b>ZBTB7A</b>	10.854	1.6118	3.70E-20								
ZF5	7.8114	3.3521	6.24E-64								
ZFP67	0.7651	11.919	1.58E-16								
ZIC3	7.8114	1.4215	2.58E-09								

Expression level after nutlin3a treatment: underlined – induced; bold – repressed; plain – no changes.

suggesting that changes in their expression might in turn modulate the regulation of p53 targets.

**Sp1 as a key determinant of alternative transcriptional program induced by RITA.** Sp1 is one of the top factors whose RE are enriched in p53-bound sites (Table 2). Taken together with our recent finding that Sp1 contributes to the induction of apoptosis by RITA in MCF7 cells,<sup>26</sup> this prompted us to test whether Sp1 might be one of the factors that confer specificity to RITA-induced transcriptional response.

Therefore, we performed microarray analysis in MCF7 cells stably depleted for Sp1 by short hairpin RNA, treated or not treated with RITA. We studied the expression of a set of 51 genes bound by p53 both upon nutlin3a and RITA treatment, but whose expression was significantly changed only by nutlin3a, but not RITA.

Strikingly, the expression of genes induced by RITA in Sp1-depleted cells was very similar to that in nutlin3a-treated cells; these data sets were clustered together (Figure 5a). Thus, Sp1 depletion converted RITA-induced expression profile of these genes to the nutlin3a-like pattern.

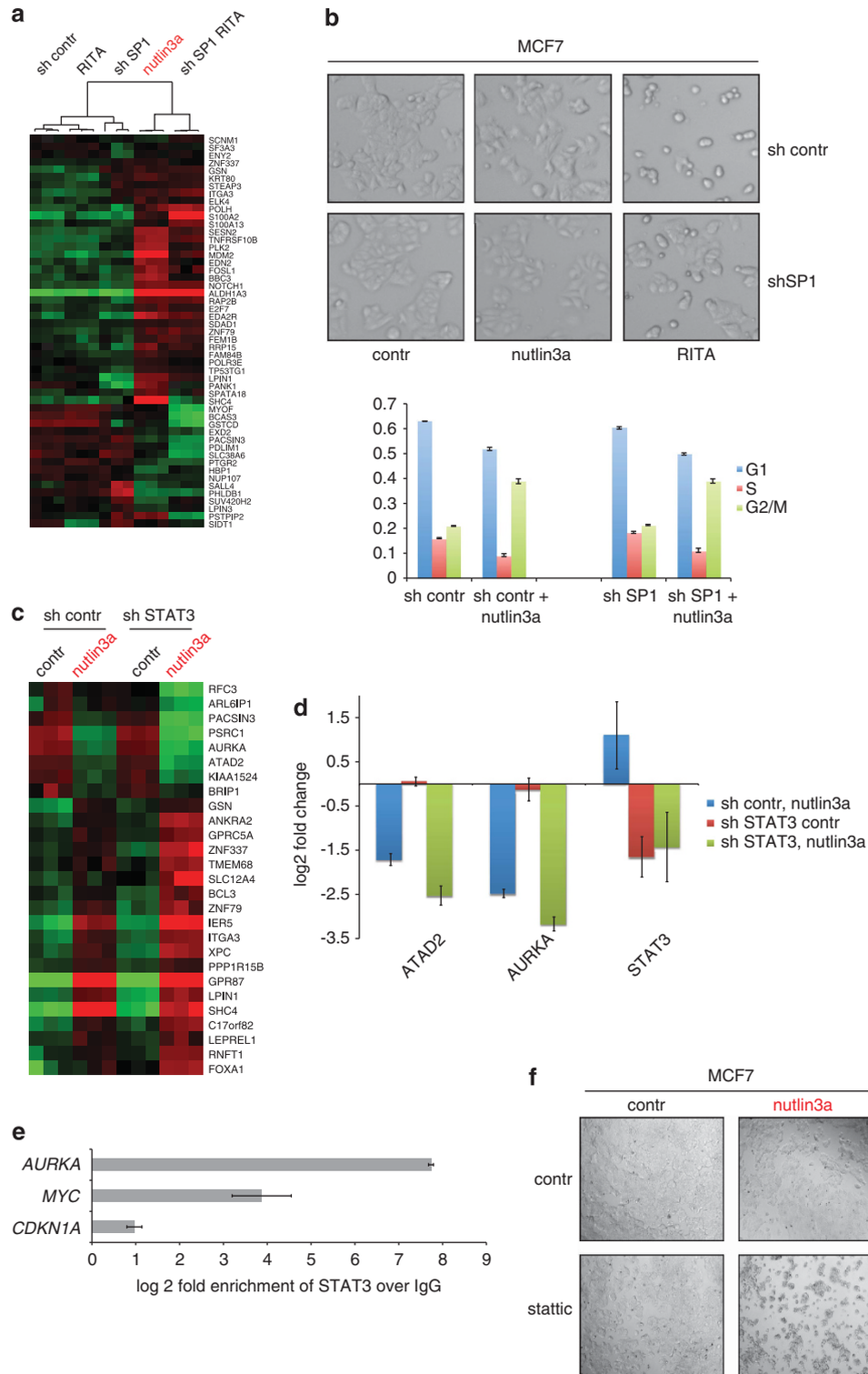
Analysis of genes induced by RITA in Sp1-depleted cells revealed the robust activation of expression of negative p53 regulators *MDM2* and *Notch* (Figure 5a). Induction of these two negative feedback loops can attenuate p53-induced biological response by RITA in Sp1-depleted cells, as it has been shown that enhanced expression of *Mdm2* and *Notch* by nutlin3a restrains the potential therapeutic efficacy of nutlin3a.<sup>27,28</sup> Further, the expression of pro-proliferative

genes *ALDH1A13*, *EDN2*, *Plk2* and *FOSL1* was strongly induced, which might also counteract growth suppression. In addition, two pro-apoptotic genes *BBC3* and *EDA2R* were also induced to the same level as upon nutlin3a treatment. However, these genes are not expected to contribute to nutlin3a-induced outcome, because it is mainly growth arrest/senescence in these cells.

Thus, the induction of two negative feedback loops and several pro-proliferative genes upon p53 activation by RITA in the absence of Sp1 might play a role in attenuation of apoptosis, which we observe in RITA-treated Sp1-depleted cells<sup>26</sup> (Figure 5b, upper panel). Sp1 silencing did not affect nutlin3a-induced suppression of MCF7 cells (Figure 5b, lower panel), suggesting that Sp1 does not play a role in nutlin3a-induced p53 response.

Taken together, our findings implicate Sp1 as one of the key determinants of alternative transcriptional programs elicited by p53 upon RITA and nutlin3a treatment. Because the set of differentially expressed genes, which we analyzed, was bound by p53 upon both treatments, these results imply that Sp1 has a discriminatory impact on some p53-target genes after RITA treatment.

**Antagonistic interaction between p53 and STAT3 at p53-repressed promoters.** As shown in Table 1, p53-STAT composite element is enriched in p53-bound and repressed genes. Because STAT3 is an important oncogene and a promising target for anticancer therapy, we studied its impact on p53 targets.



**Figure 5** Identification of Sp1 and STAT3 as important modulators of p53 transcriptional activity. (a) The heatmap shows the results of gene expression analysis of RITA-treated MCF7 cells in which *Sp1* was stably depleted by short hairpin RNA (shRNA). (b) Sp1 depletion attenuated RITA-induced apoptosis, but did not affect nutlin3a-induced growth arrest, as assessed by microscopy analysis (upper panel) and FACS of PI-stained cells (lower panel). (c) The heatmap shows the results of gene expression analysis of nutlin3a-treated MCF7 cells in which *STAT3* was stably depleted by means of shRNA. (d) Depletion of *STAT3* by shRNA enhances p53-mediated repression of *AURKA* and *ATAD2* as assessed by qPCR in MCF7 cells treated with nutlin3a. (e) *STAT3* ChIP in MCF7 cells demonstrated the binding of *STAT3* to the promoter of *AURKA*. *CDKN1A* served as a negative and *MYC* as a positive control for *STAT3* chromatin binding. (f) Inhibition of *STAT3* by static facilitated the growth-suppression effect of nutlin3a as assessed by light microscopy analysis of the MCF7 after 48 h of treatment. See also Supplementary Figure S5

Analysis of gene expression profiles upon p53 activation by nutlin3a in STAT3-depleted MCF7 cells showed that STAT3 silencing promotes p53 transcriptional activity, as a large set of p53 targets was affected by nutlin3a to a greater extent in STAT3-depleted cells (Figure 5c). Notably, *AURKA* and *ATAD2*, which we identified and validated as direct p53 targets, are also among STAT3-regulated genes (Figure 5c). Indeed, qPCR analysis confirmed that their repression by p53 is stronger upon STAT3 depletion (Figure 5d), suggesting that STAT3 hampers p53-mediated repression of *AURKA* and *ATAD2*.

Furthermore, we found a significant binding of STAT3 to the *AURKA* promoter (Figure 5e), suggesting that STAT3 antagonizes p53-mediated repression of *AURKA* by direct binding to the promoter.

Next, we found that the growth suppression by nutlin3a was facilitated upon chemical inhibition of STAT3 by stattic (STAT3 inhibitor), as well as by knockdown of STAT3 (Figure 5f and Supplementary Figure S6, respectively), suggesting that STAT3 counteracts p53-induced biological response. Taken together, our results suggest that p53 reactivation by small molecules combined with STAT3 inhibition might have a beneficial effect in the clinical setting.

Moreover, we found that the direct p53 target genes *AURKA* and *ATAD2* were among the most significantly differentially expressed genes in mutant compared with wild-type p53 breast tumor samples (Supplementary Figure S7), suggesting their high importance in p53-mediated prevention of tumor progression in patients.

## Discussion

Our data suggest that the most frequently occupied p53 sites were the same upon p53 induction by three different stimuli, nutlin3a, RITA or chemotherapeutic drug 5-FU. Thus, this pattern can be thought of as a 'default p53 program'. This is in line with the conclusions of recent ChIP-chip and ChIP-seq studies that the occupancy of the core p53-binding sites is similar regardless of the mechanism of p53 activation and cell fate.<sup>29,30</sup> Among the 'p53 default program' genes, we identified 61 known p53-target genes that are bound by p53 independently of the cell context and mechanism of p53 activation.

Parallel analysis of p53 genomic occupancy and gene expression allowed to identify 280 novel p53 target genes. Interestingly, the promoters of 63 out of 280 genes have been previously found to be bound by p53 in U2OS cells upon actinomycin D treatment using ChIP-chip analysis<sup>11</sup> (Supplementary Table S10, marked with a star).

Only 41% of genes bound by p53 were differently expressed upon nutlin3a. Thus, the binding of p53 *per se* is not sufficient to regulate gene expression. Recent study suggests that although p53 phosphorylation at S15 and S46 does not significantly affect the pattern of p53 binding to promoters, it appears to affect p53 transcriptional program,<sup>30</sup> probably via recruitment of additional factors.

Our study highlights the key role of TFs bound to p53-regulated promoters in the selective regulation of target genes by p53. We show that Sp1 can redirect p53 towards a specific transcriptional program facilitating the induction of apoptosis.

Sp1 modulates p53 transcriptional program via attenuation of the p53-mediated induction of two negative feedback loops and pro-proliferative genes, as well as via cooperation with p53-mediated repression of metabolic genes, as we previously showed.<sup>26</sup>

Compared with gene activation, less is known about p53-mediated transcriptional repression.<sup>18</sup> Our analysis of novel p53 target genes implies that the transrepression by p53 might be achieved due to the direct binding to its consensus only in certain cases. It is possible that p53-mediated repression might be indirect and/or the binding mode of p53 is different (i.e., via another TF or non-consensus binding site). Another possibility is that p53 regulates these genes via 'looping mechanism'.<sup>31</sup>

The analysis of composite binding motifs revealed the enrichment of E2F-p53 and IRF-p53 composite sites, suggesting the involvement of these two factors in p53-mediated repression of genes on a larger scale than previously anticipated. Moreover, our data suggest that another mode of p53 repression involves inhibition of proto-oncogenic TF, such as STAT3, MYB and others, via direct binding to promoters in the close proximity to their RE.

We found that STAT3, important oncogene and a promising target for anticancer therapy, antagonizes p53 activity at several promoters, including the newly identified p53 target *AURKA*. This implies that the combination of p53 activators with STAT3 inhibitors might be beneficial in a clinic.

Low occupancy of p53 at repressed promoters most probably reflects lower number of cells in which p53 is bound to these promoters at a given moment, that is, more transient binding mode. Transient binding of p53 to repressed promoters might be related to a different mode of transcription activation *versus* repression. Activation of transcription requires the constant presence of an activator to ensure open chromatin structure and accessibility for transcriptional machinery. In contrast, once the repressive chromatin state at the promoter is established, it is self-maintained in the absence of an inducing signal due to initial modification, providing new binding sites for the enzymes that maintain chromatin modifications. Therefore, it does not require the continuous presence of the initial repressor.<sup>32</sup>

In conclusion, our data strongly suggest that p53-mediated target gene expression upon activation by nutlin3a, RITA or 5-FU is determined not only by a selective binding of p53 to its target promoters but also depends on cooperating or antagonizing TFs. These results are expected to help in a future to harness p53 activity by targeting p53 modulators and achieve a desired p53 transcriptional response.

## Materials and Methods

**Cell-based assays.** MCF7 is a human breast carcinoma line and HCT116 and HCT116 p53<sup>-/-</sup> cells, a gift from B Vogelstein, are colon carcinoma. Cell lines were treated with 1  $\mu$ M RITA, 10  $\mu$ M nutlin3a or 100  $\mu$ M 5-FU for 8 or 4 h before analysis. Stattic 5  $\mu$ M was used to inhibit STAT3 activity. The pGL3 luciferase construct, used for the cloning of promoter fragments, was provided by S Mkrtchian of the Karolinska Institutet. The *AURKA* promoter (-518 to +355) and *SEI1* promoter (-545 to +294) were cloned using the primers listed in Supplementary Table S11. Sp1-depleted cell line was established as described in Zawacka-Pankau *et al.*<sup>26</sup> Transfections were performed using Lipofectamine 2000 (Invitrogen, Grand Island, NY, USA). Luciferase activity was measured using the Dual-Luciferase Reporter System (Promega, Madison, WI, USA) and analyzed as described before.<sup>33</sup>

Cell viability and fluorescence-activated cell sorting analysis were performed as previously described.<sup>15</sup> Western blotting was performed according to the standard procedures. The proteins were detected using the following antibodies: anti-p53 DO-1 (Santa Cruz Biotechnology, Santa Cruz, CA, USA), anti-Aurora A ab13824 (Abcam, Cambridge, UK) and anti-beta-actin (Sigma-Aldrich, St. Louis, MO, USA).

**ChIP and ChIP-seq experiment.** ChIP in MCF7 cells, library preparation, massive parallel sequencing and ChIP-Seq primary analysis was performed as previously published.<sup>34</sup> Sc-126 (Santa-Cruz Biotechnology, Santa Cruz, CA, USA), Genespin DO-1 p53 and Sc-2025 mouse IgG antibodies were used for IP.

Approximately 6 million high-quality sequencing reads were mapped to the human genome (NCBI36) and used to calculate the height of the peaks as the number of overlapping reads extended to the fragment length. The number of extended reads in each p53 peak was compared with the number of extended reads in the same region in an IgG-precipitated sample and enrichments were calculated. Data are publicly available from the NCBI sequence Read Archive and in Supplementary data (Supplementary Tables S1–S4). Data are archived at NCBI Sequence Read Archive under Accession SRP007261.

Intersection of the peaks ( $P < 0.05$ ) after different treatments was performed with a web-based tool, Galaxy,<sup>16</sup> and imported into R (<http://www.r-project.org/>). Venn diagrams were plotted with Vennable R package (<http://R-Forge.R-project.org/projects/vennerable/>). The distribution of the peaks around the TSS was calculated using the ChIPpeakAnno package.<sup>35</sup> For *de novo* motif discovery analysis, we used the MEME<sup>36</sup> or p53MH<sup>37</sup> and p53Scan<sup>11</sup> programs. As input data, we used sequences at a distance  $\pm 200$  bp from the position of the peak maximum height.

The presence of TF-binding sites located at  $\pm 100$  bp from p53-binding sites in ChIP-Seq fragments and composite patterns were analyzed with the tool MATCH that uses a library of about 600 positional weight matrices (PWMs) from the TRANSFAC database for vertebrate TFs.<sup>38</sup> Overrepresentation of TF-binding sites and composite elements (TF site pairs) were computed by the F-Match algorithm and composite module analyst program (Composite Module Analysts) that is implemented in the geneXplain platform ([www.genexplain.com](http://www.genexplain.com)).

**Microarray experiment.** Expression profiling of MCF7 cells was performed using Affymetrix (Santa Clara, CA, USA) human genome 219 arrays as previously described.<sup>26</sup> Primer sequences for qPCR are listed in Supplementary Table S13.

### Conflict of Interest

The authors declare no conflict of interest.

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