Genome-wide profiling reveals stimulus-specific functions of p53 during differentiation and DNA damage of human embryonic stem cells

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ABSTRACT

How tumor suppressor p53 selectively responds to specific signals, especially in normal cells, is poorly understood. We performed genome-wide profiling of p53 chromatin interactions and target gene expression in human embryonic stem cells (hESCs) in response to early differentiation, induced by retinoic acid, versus DNA damage, caused by adriamycin. Most p53-binding sites are unique to each state and define stimulus-specific p53 responses in hESCs. Differentiation-activated p53 targets include many developmental transcription factors and, in pluripotent hESCs, are bound by OCT4 and NANOG at chromatin enriched in both H3K27me3 and H3K4me3. Activation of these genes occurs with recruitment of p53 and H3K27me3-specific demethylases, UTX and JMJD3, to chromatin. In contrast, genes associated with cell migration and motility are bound by p53 specifically after DNA damage. Surveillance functions of p53 in cell death and cell cycle regulation are conserved in both DNA damage and differentiation. Comparative genomic analysis of p53-targets in mouse and human ESCs supports an inter-species divergence in p53 regulatory functions during evolution. Our findings expand the registry of p53-regulated genes to define p53-regulated opposition to pluripotency during early differentiation, a process highly distinct from stress-induced p53 response in hESCs.

INTRODUCTION

Tumor suppressor protein p53 is well studied in its roles of cellular surveillance by activation of cell cycle arrest and cell death pathways (1). Recent work expands p53 regulatory functions to cellular metabolism and homeostasis, implantation, aging and quiescence of stem cells [reviewed in detail in (2)]. Tumor suppressor p53 has been implicated in limiting the self-renewal of stem cells in variety of systems (3–5). Reports that p53 loss in breast cancers activates genes with embryonic stem cell (ESC) transcription signatures (6) and that p53 mutation enables aberrant self-renewal of hematopoietic stem cells to promote acute myeloid leukemia (7) support p53 functions in opposition to a stem cell state.

The characteristics shared between specific cancers and stem cells (8), and the dysfunction of p53 in a majority of human cancers (9), led to a proposal that p53 imposes differentiation as a tumor-suppressive mechanism (2,10). Multiple studies show that developmental reprogramming of a fully differentiated somatic cell to an induced pluripotent stem cell state is facilitated by depletion of p53 or malfunction in pathways that activate p53 [reviewed in (11)]. However, whether p53 is a passive barrier to reprogramming and creation of induced pluripotent stem cells, due to its inhibitory effects on cellular proliferation, or if it actively opposes pluripotency remains controversial (12). Recent work from our laboratory revealed that p53 plays a significant role during retinoic acid (RA)-mediated differentiation of human ESCs (hESCs) by regulating cell cycle and activating micro-RNAs that repress stem cell factors (5). Prior studies comparing mouse embryonic
stem cells (mESCs) and hESCs show fundamental differences between the two, including basic mechanisms of transcription factor function (13,14) and the impact cell cycle has on self-renewal (15). These differences may reflect different stages of development: inner cell mass (mESCs) versus epiblast (hESCs) (16). However, further discovery and analysis of the earliest stages of human embryonic development are needed to resolve such controversies.

Genome-wide profiling of protein interactions with chromatin and intersection with global gene expression analysis allow functional annotation of directly regulated biological pathways. These approaches have been instrumental in revealing networks of transcription and epigenetic regulators of pluripotency and self-renewal of ESCs (17,18). Here, we applied these methods and identified genome-wide downstream targets of p53 in hESCs undergoing differentiation or DNA damage. Using a combination of genomic and biochemical approaches, we found that p53 responds to RA signaling by direct regulation of genes significant in developmental processes. These gene targets of p53 are distinct from those induced during DNA damage, which function in cellular migration and motility. Moreover, during differentiation p53 interacts with genomic regions occupied by core pluripotency factors OCT4 and NANOG in pluripotent hESCs. Using conventional ChIP and sequential ChIP, we show that p53 binds OCT4-occupied gene loci and mediates loss of H3K27me3 to activate expression of specific developmental genes. Further, by co-immunoprecipitation analysis of chromatin, we show that p53 interacts with H3K27me3 demethylases, UTX and JMJD3, which alter chromatin, we show that p53 interacts with H3K27me3 to activate expression of specific developmental genes. Collectively, our results broaden the known transcriptional functions of p53 and reveal directly regulated gene signatures unique to either differentiation or damage response, as well as the prime functions of p53 in hESCs conserved in both.

**MATERIALS AND METHODS**

**Cell culture**

Human Embryonic Stem (WA09) Cells were obtained from National Stem Cell Bank (Madison, WI, USA) and cultured according to the protocol from WiCell Research Institute. Briefly, WA09 cells were maintained in hESC cultured according to the protocol from WiCell Research Institute. hESCs were obtained from Cell culture

**ChIP**

hESCs grown on inactivated MEF feeders were cultured in complete hESC media or treated with RA for 2 days in differentiation media were collected, PBS washed and cross-linked in 1% formaldehyde for 10 min at room temperature. After glycine followed by PBS wash, cells were lysed using lysis buffer [5 mM Hepes (pH 8.0), 85 mM KCl, 0.5% NP40] supplemented with protease inhibitor (Calbiochem). After removal of cytoplasmic extract, remaining cell pellet was lysed in Nuclear lysis buffer [50 mM Tris–HCl (pH 7.5), 10 mM EDTA, 1% SDS] and protease inhibitor. Lysates with glass beads (Sigma) were fragmented with a Bioruptor (Diagenode) to obtain DNA fragments of average length under 500 bp. After centrifugation, the supernatant was preabsorbed with 40 µl of Protein A beads (GE Healthcare) and immunoglobulin G (IgG) for 2 h, then incubated with 5 µg of p53 (Santa Cruz Biotechnology), OCT4, Histone H3, (both from Abcam), H3K4me3 or H3K27me3 antibody or control IgG (all from Millipore) overnight at 4°C. The immunocomplex was collected on protein A beads, washed and bound DNA was eluted and reverse cross-linked overnight at 65°C. The DNA region of interest was detected by SYBR real-time quantitative PCR using primers encompassing p53-response elements (p53RE) or OCT4-binding sites on the respective gene (see Supplementary Table S7 for sequence of primers used for ChIP-qPCR).

**ChIP-seq analysis**

Sequencing of p53-bound DNA was performed at the Bioinformatics Core of the Cincinnati Children’s Hospital Medical Center, Cincinnati, OH. p53-bound DNA (~10 ng) was purified by PAGE to obtain 100–300 bp fragments and sequenced on an Illumina Solexa GAII sequencer. Sequence reads derived from Illumina GAII sequencer were aligned to the NCBI Build 36 (UCSC hg18) human genome using ELAND software (Illumina) to produce uniquely matched reads with up to two mismatches per read allowance. We obtained 15012584, 14655088 and 11280919 uniquely mapped reads to the genome in untreated, differentiation and
damage conditions, respectively. Enriched regions for each condition were normalized to input DNA and detected by MACS v1.4.0 (19) with a cut-off \( P < 10^{-5} \) for damage and differentiation, and \( P < 10^{-10} \) for untreated. Wiggle files were generated by the same sequence file and later normalized. Peaks share at least one base under their enriched regions called as overlapped between different conditions. The distance between unique peaks in each condition was measured using a gradually increasing window and determining the unique peaks summits coinciding in the same window. Obtained numbers were plotted, and pie charts were generated by ratios of overlapping versus non-overlapping summits for a certain window length. PhastCons conservation scores for 44 vertebrates were downloaded from UCSC website (http://hgdownload.cse.ucsc.edu/goldenPath/hg18/phastCons44way/vertebrate/), and individual chromosome files were merged into a single wiggle file. Aggregate plots for conservation scores across enriched sites were generated using the Sitepro program under CEAS (20).

Motif analysis was performed using MEME-ChIP (21), which is specifically designed to discover associated motifs in large sets of DNA sequences. Zero or one motif per sequence was searched with the motif lengths between 6 and 30 bp. Analysis of motif enrichment (22) outputs for each data set are shown with a \( P \)-value cut-off \(< 1.00 \times 10^{-10} \). SeqPos motif discovery program in Cistrome analysis pipeline (http://cistrome.org/ap/) was also performed for motif discovery underneath enriched sites (~400 bp of the peak’s center) in each condition.

Human RefSeq gene information was obtained from UCSC table browser (http://genome.ucsc.edu/cgi-bin/hgTables?command=start). Weighted binding coefficient was calculated as previously described (23), to determine the fold enrichment over the randomized binding sites. Genes with a nearby peak 10 kb up/downstream of transcription start sites (TSS) were designated as targets. Gene Ontology (GO) analyses for each set of target genes were performed using DAVID (24). Developmental transcription factors were obtained from the HUGO website (http://www.genenames.org/genefamily.html), previously published study (17) and NCBI’s Gene database (http://www.ncbi.nlm.nih.gov/gene). Each dot shown represents a member of a particular family only if the gene’s ontology terms entail transcription or DNA binding and also development or differentiation.

INTERPRO protein domain analysis was performed using Genomic Regions Enrichment of Annotations Tool (GREAT) (25). Peak files were imported into GREAT by setting a gene association rule as a single gene within 10 kb ranges of binding sites. The top five categories are shown.

ChIP-Seq data sets of OCT4 (GSM518373) and NANOG (GSM518374) were obtained from GEO database (http://www.ncbi.nlm.nih.gov/geo/). Raw sequences were re-analyzed with MACS v1.4.0. Obtained peaks were used for overlap analysis and circular plot. Circos (26) was used to visualize p53, OCT4, NANOG and Histone 3 Lysine 27 trimethylation (H3K27me3) around four Homeodomain-box (HOX) clusters. H3K27me3 data were obtained from UCSC genome browsers’ ENCODE project website (http://hgdownload.cse.ucsc.edu/goldenPath/hg18/encodeDCC/wgEncodeBroads/ChipSeq/) (27). Random locations showing similar distribution in each chromosome were generated 10 000 times and used for determining statistical significance. Wiggle files were used to generate a density plot, using the heatmap tool in the Cistrome analysis pipeline (http://cistrome.org/ap/). K-means clustering was applied to the condition specific regions. Normalized wiggle files were used to generate histone aggregate plots. TSS of p53 target genes was used as the center of the window and each window was divided into 40 bins of 25 bp resolution. Average ratios were plotted for each category.

Microarray analysis

Affymetrix U133 Plus2.0 microarrays were performed for each condition (Pluripotent, +Adr and +RA) in triplicates. Robust multi-array average method was used with default options (with background correction, quantile normalization, and log transformation) to normalize raw data from batches using R/Bioconductor’s affy package (28). For genes, which are represented by multiple probes on the array, maximum expression value was retained for further analyses. A gene is called as differentially expressed if FDR corrected \( P < 0.05 \), which is calculated with empirical Bayes method by eBayes function in Bioconductor’s limma package. GO analysis of differentially expressed gene was performed using DAVID.

RNA knockdown

hESC colonies were grown on Matrigel in 6-well plates. siRNA-targeting human TP53, and non-target (control) were purchased from Dharmacon. In all, 75 nM siRNA oligos were transfected twice into cells using Lipofectamine2000 (Invitrogen) transfection reagent according to manufacturer’s protocol within a period of 5 days. First siRNA transfection was performed 24 h after splitting the cells followed by media change 6 h post-transfection. Thirty-six hours after first siRNA transfection, cells were cultured in differentiation media (−FGF, +RA) for 3 days and harvested. Cells were transfected one more time with siRNA on beginning of day 2 of RA treatment to maintain the knockdown efficiency.

RNA isolation and real-time qPCR analysis

RNA was isolated using TRIzol Reagent (Invitrogen) as per manufacturer’s specifications. For RNA analysis 500 ng total RNA was treated with DNAse and cDNA was synthesized as previously described (29), and amplified with human gene specific primers (Supplementary Table S8) with Power SYBR Green PCR Master Mix (Applied Biosystems). The average threshold (Ct) was determined for each gene and normalized to Actin mRNA level as internal normalization control.
**Chromatin fractionation**

Chromatin fractions were prepared using biochemical fractionation protocol as described previously (30) with some modifications. Briefly, pluripotent or RA treated hESCs were harvested and washed with chilled phosphate-buffered saline (PBS). The cells were then resuspended in buffer A [10 mM HEPES (pH 7.9), 10 mM KCl, 1.5 mM MgCl2, 0.34 M sucrose, 10% glycerol, 1 mM DTT and protease inhibitor cocktail]. Triton X-100 was added to cells at a final concentration of 0.1%, followed by incubation on ice for 8 min, and nuclei were collected by centrifugation (5 min, 1300g, 4°C). The supernatant was clarified by high speed centrifugation (5 min, 20000g, 4°C) to recover the cytoplasmic extract. The nuclei were washed once with buffer A and resuspended for 30 min in buffer B (3 mM EDTA, 0.2 mM EGTA, 1 mM DTT and protease inhibitor cocktail), and insoluble chromatin pellet and soluble fractions were separated by centrifugation (5 min, 17000g, 4°C). The chromatin pellet was resuspended in immunoprecipitation (IP) buffer (50 mM Tris–HCl, 1% Triton X-100, 150 mM NaCl and 15 U of MNase (Worthington) at 37°C for 10 min, centrifuged (5 min, 10000g, 4°C), and the supernatant was used for immunoprecipitation and western blot analyses.

**Co-immunoprecipitation analysis**

hESCs grown on inactivated MEF feeders were cultured in complete hESC media or treated with RA for indicated time points in differentiation media, were subjected to chromatin fractionation as described above. 0.5 mg of chromatin fractions were used for immunoprecipitation with 2.5 μg of antibody overnight at 4°C with shaking. Thirty microliters of washed Protein A Sepharose CL-4B bead suspension (GE Healthcare) was added to fraction-antibody mixture and incubated for additional 1 h at 4°C with shaking. Immune complexes were washed twice with IP buffer without MNase, boiled with 1x SDS SDS–Laemmli sample buffer and resolved on 10% SDS–PAGE followed by immunoblotting.

**Western blotting**

hESCs were lysed in RIPA buffer [50 mM Tris (pH 8.0), 150 mM NaCl, 1% NP-40, 0.5% Deoxycholic acid, 0.1% SDS] supplemented with protease inhibitor cocktail (Calbiochem) and phosphatase inhibitor cocktails I and II (Sigma) to obtain whole-cell lysate. Approximately 50 μg of cell lysate or chromatin fraction was analyzed by immunoblotting with anti-p53 (DO1), anti-OCT4, (Santa Cruz Biotechnologies), anti-UTX (Bethyl Laboratories), anti-MJMD3 (Abcam), anti-H3 (Abcam), anti-LaminB1 (Santa Cruz Biotechnologies) and anti-Actin (GeneTex Biotechnology), followed by corresponding Horseradish peroxidase-tagged secondary antibody (GE Healthcare) as described previously (5).

**Generating stable-knockdown hESCs**

hESCs were cultured in mTeSR1 medium as described above. Lentiviral supernatants were produced by transfecting 293T cells with the pLKO.1 control shRNA vector (Open Biosystems) or pLKO.1-shp53 vector (Addgene plasmid 19119), along with pSbPAX2 and pMD2.G packaging plasmids (Open Biosystems), using FuGene HD transfection reagent (Roche) and collected viral media 48 h post-transfection. Stable knockdown hESCs were generated by incubating the cells with viral media followed by spin infection at 2500 rpm at 32°C for 90 min with polybrene (4 μg/ml) and incubated at 37°C for an additional 2 h before medium replacement with fresh mTeSR1. Three days after infection, cells were selected with increasing doses of Puromycin (0.2–2 μg/ml) and finally maintained in 1 μg/ml Puromycin. p53-knockdown efficiency was confirmed by RNA and protein analysis.

**Comparing p53-binding profiles in human and mouse ESCs**

ChIP-Seq data sets of mouse p53 DNA-damage (GSE26362) and H3K27me3 in mESC (GSM397410) (31) were obtained from GEO database. Raw sequences were re-analyzed with MACS v1.4.0 with a P-value cut-off of $P < 10^{-8}$. UCSC lift over tool (http://genome.ucsc.edu/cgi-bin/hgLiftOver) was used to convert mouse genomic coordinates to human hg18 genome assembly. Obtained peaks were used for overlap analysis and circular plot. Genes with a p53 peak within 10 kb up/downstream of TSS were designated as p53 targets. Based on the interspecies transcription factor target gene classification as described previously (32), mouse p53 bound to aligned region in human genome for homologous genes were designated as ‘Conserved’ target genes. Homologous genes with unaligned p53 binding sites between species were designated as ‘Turnover’ p53-targets. GO analyses for each set of target genes were performed using DAVID annotation tool (24). Circos (26) was used to visualize p53, Phastcons Scores and H3K27me3 around four HOX clusters in human and mouse genomes.

**Data access**

p53, H3K4me3 and H3K27me3 ChIP-seq data can be accessed at NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/) under accession no. GSE39912. Affymetrix microarray expression data are available under accession no. GSE39762.

**RESULTS**

**Genome-wide mapping of p53 in hESCs reveal distinct functional binding sites**

Alkaline phosphatase (AP)-staining of hESCs, as one measure of self-renewing (positive staining) versus differentiated (negative staining) state, was monitored for hESCs treated with siControl oligos or depleted of p53, followed by RA-induced differentiation or DNA damage induced by Adriamycin (ADR) (Supplementary Figure S1A). Adr elicited a robust induction of p53, no major change in OCT4 protein levels and a minor reduction in AP-staining of hESCs, indicating no significant differentiation under DNA damage conditions whether
p53 is depleted or not (Supplementary Figure S1A and B). In contrast, RA treatment induced p53 protein and both p53-dependent loss in AP-staining and p53-dependent reduced OCT4 protein levels (Supplementary Figure S1A and B), as we previously reported using multiple measures of stem cell status (5).

To determine the molecular basis for these signal-specific responses and define a landscape of p53-chromatin interactions in hESCs, we mapped p53 occupancy throughout the genome by deep sequencing of p53-binding chromatin fragments (ChIP-Seq) isolated from hESCs in a pluripotent state (untreated), undergoing differentiation (+RA) or after DNA damage (+Adr). In pluripotent hESCs, p53 is enriched at 4509 genomic sites, compared with 8282 and 4941 in hESCs undergoing differentiation or damage, respectively (Figure 1A). A fraction of p53-bound peaks (26.5%) overlapped between differentiation and damage induction (Figure 1A). Comparison of unique sites in a gradually increasing genomic window revealed that only 44% of unique sites overlapped in a 100 kb window, suggesting highly diverse p53 functions in these two states (Supplementary Figure S1C). PhastCons score analysis, comparing genomic regions within 4 kb of each p53-peak summit in 44 vertebrate species, revealed high evolutionary conservation of p53 binding regions (Figure 1B).

Comparison of p53 ChIP-Seq signals to input suggests robust p53 enrichment during both differentiation and damage (Supplementary Figure S2A), consistent with p53 protein stabilization under these conditions (Supplementary Figure S1B). p53-bound regions were significantly enriched with consensus p53-binding sites (p53-motif) in both differentiation and damage (P < 10^{-3} and P < 10^{-235}, respectively, Figure 1D), a motif that is similar to the p53 consensus obtained from TRANSFAC (Figure 1C, Supplementary Figure S1D and Supplementary Table S1). Interestingly, sequences bound by p53 in pluripotent hESCs (untreated) did not generate a similarly high confidence, p53-consensus motif (P > 10^{-5}, Figure 1D). Our finding of lower significance for p53 interactions with chromatin, in the absence of inductive signaling, is likely the result of unstable interactions by low levels of p53 in pluripotent hESCs (5). These findings are also consistent with a model proposed by Prives and colleagues (33). In this scenario, basal p53, lacking stimulus-dependent post-translational modifications and present at low concentration, scans and interacts with p53 response element (p53RE) sequences with a high on-off rate. These sites, where p53 interaction may be low-affinity/high off-rate but sufficient for cross-linking in ChIP, display similar parameters of relative distance to gene TSS, histone marks and chromatin accessibility (DNaseI) as p53 sites of treated hESCs (data not shown). Basal p53 interactions with chromatin are unlikely productive with regards to transcription regulation; hence, we did not consider these data for further analyses.

Interestingly, high confidence p53-bound regions in hESCs undergoing differentiation were significantly enriched in OCT4- and SOX2-binding motifs (P < 10^{-16} and P < 10^{-12}, respectively) (Figure 1C and D), whereas no OCT4-SOX2 motifs were found in p53-bound genomic regions in hESCs exposed to damage or untreated (Figure 1C and D). Reciprocal analysis of OCT4-SOX2 and NANOG-enriched sites in pluripotent hESCs, using previously published ChIP-Seq data sets (13), revealed overlapping p53 response elements (p53REs) in both data sets (Supplementary Figure S1E). Thus, the presence of consensus binding motifs for OCT4 and SOX2 in p53-bound regions, and vice-versa, suggests a possible interplay between these transcription factors in determination of specific stem cell states.

p53 targets developmental transcription factors during differentiation

Across the genome, a significant portion of p53-binding sites (42% for +RA and 28% for +Adr) are enriched (weighted binding coefficient of 0.68-fold for +RA and 0.61-fold for +Adr compared with randomized binding sites) (23) within 10 kb of the nearest annotated TSS (Supplementary Figure S2B). Therefore, we used a 10 kb window of distance from the p53-peak summit to the nearest gene TSS to define a p53 target gene (Figure 2A). Gene comparison analysis revealed only 22% overlap in identity (717 genes) between damage (1326 genes) and differentiation (3172 genes) (Figure 2A), suggesting distinct roles for p53 dependent on cellular environment. GO analysis revealed a striking distinction between genes regulated by p53 during differentiation versus damage (Figure 2B). During differentiation, a majority of p53-target genes is associated with functions in development and transcription regulation (P < 10^{-4}). In contrast, damage-specific p53-gene targets are primarily linked to cell migration and motility (P < 10^{-4}), as determined by GO analysis of each set of stimulus-specific gene targets (Figure 2B and Supplementary Table S2). Highly studied p53-regulated genes with functions in cell death and cell cycle, e.g. CDKN1A, MDM2, are significantly represented (P < 10^{-5}) in p53-bound genes common to both differentiation and damage (Figure 2B).

Next, we determined enrichment of protein domains encoded by p53-target genes in each condition using InterPro terms of the GREAT functional annotation tool. Homeobox domains were revealed as differentiation targets (P < 10^{-11}), and the broad classification of EGF-like domains was enriched across damage-specific p53 targets (P < 10^{-6}) (Figure 2C). Several transcription factors that regulate specification and development are highly represented as differentiation targets (Figure 2D and Supplementary Table S3). These include members of the HOX gene family, which are activated as an early response to RA (34); LIM homeobox (LHX) genes, which are involved in embryonic development and specifically neuronal differentiation (35); the forkhead box (FOX) family of genes, which are involved in axial patterning and tissue development from all three germ layers (36); the sex-determining region-Y box (SOX) gene family that regulates cell-fate specification (37); and, Zic family members (ZIC) that are important during neuronal development, mutations of which cause a wide variety of
congenital malformations (38) (Figure 2D and Supplementary Table S3). These findings suggest that, during differentiation of hESCs, the regulatory influence of p53 is extensive and amplified by targeting transcription factors that promote a committed cellular state.

p53-binding sites coincide with ESC transcription factors during differentiation

Specific developmental genes, which are inactive in pluripotent ESCs and activated early during differentiation, have been described as ‘poised’, owing to sequence-specific binding of core pluripotency factors (OCT4, SOX2 and NANOG) within chromatin enriched in bivalent histone modifications: both repressive H3K27me3 and activation-associated H3K4me3 (18,39). The bivalent state of chromatin is thought to predispose poised genes, e.g. HOX genes, with major roles in differentiation for rapid response to differentiation signals. A circular plot of human chromosomes 2, 7, 12 and 17, representing a ~100 kb region of each of four HOX gene clusters, was used to illustrate enrichment of OCT4, NANOG and H3K27me3 in pluripotent hESCs (Figure 3A). When we compared sites of p53 binding (induced by RA or Adr) to OCT4, NANOG and H3K27me3 enrichment across the HOX genes, we saw that RA-induced p53 binds (20 binding sites for 11 genes) in and around the poised HOX genes during differentiation; whereas, in contrast, there is only one p53-bound site induced by DNA damage at these loci, and this lies outside regions of OCT4/NANOG binding.

Overlap between RA-induced p53 and pluripotency-associated OCT4 and NANOG-binding sites is widespread across the genome, as ~50% of the 1000 highest confidence, differentiation-bound p53 sites are occupied by OCT4/NANOG or both in pluripotent hESCs; only a small fraction (~12%) overlap with damage-specific p53 sites (Figure 3B and Supplementary Figure S3A). Co-occupancy analysis genome-wide showed a considerably higher ratio of p53:OCT4:NANOG overlap and stronger p53 peaks at differentiation- versus damage-induced binding sites (Supplementary Figure S3B). Heat map analysis revealed that there is a substantial enrichment of OCT4 and NANOG at genomic sites bound by p53 during hESC differentiation versus damage (Figure 3C). Binding profiles and comparison of p53 and NANOG
peaks reveal that OCT4 enrichment at p53 peaks, established during differentiation, is of the same magnitude as at NANOG sites. However, NANOG enrichment is stronger at OCT4-binding sites than at sites bound by p53 (Supplementary Figure S3C). Taken together, this suggests that a specific subset of genes is kept in a repressed but poised state by OCT4/NANOG during pluripotency and, in response to RA, p53 occupies these genes to promote hESC differentiation. The lack of overlap between OCT4/NANOG binding in pluripotent

hESCs and damage-induced p53 sites of binding, versus a high degree of overlap with differentiation-induced p53 sites, suggests that distinct mechanisms are involved in stimulus-specific p53-mediated regulation in hESCs.

Transcription of development genes is dependent on p53

To uncover the functional consequences of p53 interactions with chromatin, we performed microarray-based gene expression analysis of hESCs undergoing

Figure 2. p53 targets developmental transcription factors during differentiation. (A) Numbers of distinct and overlapping p53-target genes in hESCs undergoing differentiation and DNA damage. (B) GO term analysis revealed significant and diverse functions of p53 downstream target genes that are specific or shared in response to each treatment. (C) Enrichment analysis of protein domains encoded by p53 downstream targets. Top categories from each data set are listed. (D) Gene families of developmental transcription factors are targets of p53 during differentiation. p53 (green circle) regulation is linked to individual transcription factors (cyan circles), shown grouped by family.
Figure 3. p53-binding sites coincide with ESC transcription factors during differentiation. (A) Circos plot of four human HOX gene clusters showing differential binding patterns of OCT4 (blue), NANOG (red), p53 (damage:yellow, differentiation:orange) and H3K27me3 (green). Numbers outside the circle is for specific human chromosome. (B) Overlap of p53 occupancy with OCT4 and NANOG in hESCs undergoing differentiation or damage. (C) Heat map of binding signals of p53, OCT4 and NANOG within −500 bp to +500 bp of p53 peak summits.
differentiation and integrated these data with our p53 ChIP-Seq data set (Figure 4A and B). Expression analysis revealed a total of 1220 up- and 1221 downregulated genes during differentiation of hESCs. Intersection with our p53 ChIP-Seq revealed that >25% of genes regulated during differentiation (262 down- and 361 upregulated) are bound by p53. Differentiation-specific p53 targets (198 down- and 271 upregulated) were assigned by eliminating genes also responsive to DNA damage (Figure 4A and B). GO analysis of RA downregulated p53 targets revealed that these genes are enriched for cell motility and mesodermal differentiation (Figure 4C). These genes include FOXO3: an essential activator of mesodermal marker Brachyury (40); KLF6: associated with hematopoiesis (41); chromatin modifiers HDAC5 and HDAC9; class II HDACs with critical functions in heart development (42); and, telomere repeat binding factor TERF1: a telomere maintenance factor associated with pluripotency (43) (highlighted in Figure 4A and Supplementary Table S4). RA upregulated p53 targets revealed significant ($P < 10^{-5}$) representation in neuro-ectodermal development, embryonic morphogenesis and pattern specification categories (Figure 4C). These genes include homeobox domain genes (HOXA1, HOXA3, HHEX and HOXB1), developmental transcription factors (GATA2, LHX8, ZIC1 and TCF7L2) and RA nuclear receptors (RARα and RARB) (highlighted in Figure 4A). Several of these genes are repressed by Polycomb complexes and classified as poised by core pluripotency factors in pluripotent hESCs (17), but a role for p53 in their activation during differentiation has not previously been reported.

We performed quantitative RNA and p53 ChIP-PCR analyses of selected genes to assess the impact and specificity of p53 binding (Figures 4 and 5). RA treatment for 2 days resulted in significant activation of genes belonging to HOX and GATA families (Supplementary Figure S4A). Four days of RA increased expression of these genes, as well as developmental transcription factors: TBX5, homeobox genes MSX2 and GBX2, hedgehog receptor PTCH1, Notch co-repressor TLE3, polycomb protein BMI1 and histone H3K36 demethylase KDM2B (Figure 4D). RA-mediated activation is dependent on p53, as hESCs transfected with siTP53 showed a significant reduction in activation of these genes with RA treatment compared with siControl. In contrast, p53 induction by DNA damage had no significant effect on these genes (Supplementary Figure S4A). Expression of CDKN1A and MDM2 was induced during both differentiation and damage in a p53-dependent manner (Supplementary Figure S4B).

We obtained previously published OCT4 and NANOG ChIP-Seq data sets in hESCs from GEO database (13), see ‘Materials and Methods’ section and compared those with p53-enriched regions of the human genome to map OCT4, NANOG and p53-binding elements at specific developmental genes: HOXAI, PTCH1 and TBX5 (Figure 5A, schematic diagram). ChIP-qPCR analyses revealed significant enrichment of p53 binding, within 2 days of RA exposure, at the p53REs of PTCH1, HOXAI, TBX5 and CDKN1A (Figure 5B). Importantly, p53 enrichment at RA-specific genes, PTCH1, HOXAI and TBX5, is little changed in response to DNA damage. Both differentiation and damage induce enriched p53 binding at CDKN1A and activation of p21 (CDKN1A) transcription (Figure 5B). We performed sequential ChIPs (re-ChIP) to assess whether OCT4 and p53 co-occupy or exclusively bind to their overlapping binding sites at PTCH1 and TBX5, where the distance between p53- and OCT4-binding sites is within the size range of our fragmented chromatin length (<500 bp), as diagrammed in Figure 5A. We assessed binding of p53 on OCT4-enriched chromatin fragments from hESCs treated with RA for 2 days and found that RA significantly induced p53 enrichment and co-occupancy at OCT4-associated regions of PTCH1 and TBX5 (Figure 5C), roughly equivalent to the increase in p53 association induced by RA (Figure 5B). The OCT4-OCT4 re-ChIP indicates equal efficiency of OCT4 binding to chromatin sites in both untreated and 2-day RA-treated hESCs. Our analysis indicates that, early in RA-mediated differentiation, co-occupancy of p53 and OCT4 occurs within the same region of chromatin and is consistent with activation of gene expression. Robust expression of these p53-target genes, as OCT4 protein levels decrease with differentiation, is likely dependent on alteration and activation of chromatin structure during differentiation.

p53 target genes lose repressive histone marks during differentiation

Developmental genes are held silent in pluripotent ESCs by repressive chromatin enriched in specific histone post-translational modifications (H3K27me3), which are lost during differentiation (18). We generated hESCs stably expressing non-target (shControl) or shRNA against p53 (shTP53) to determine whether RA-activated p53 had an impact on levels of H3K27me3 at the promoters and/or p53REs of PTCH1, TBX5 and CDKN1A (Figure 5D and Supplementary Figure S5). Stable integration of shTP53 resulted in substantial knockdown of p53 protein and failure to elicit an RA-response, as no reduction in AP staining and OCT4 protein was observed in shTP53-hESCs as compared with control (Supplementary Figure S5A and B). In response to RA, H3K27me3 levels are significantly reduced at PTCH1, TBX5 and CDKN1A loci (shControl), whereas no change in H3K27me3 levels were observed in hESCs stably depleted of p53 (shTP53) (Figure 5D and Supplementary Figure S5C). Together, these results suggest that RA-induced signals of differentiation mobilize p53 to bind and activate chromosomal locations, which are repressed but relatively accessible due to binding by OCT4/NANOG in pluripotent hESCs, and this p53 binding alters chromatin status during differentiation.

We tested this hypothesis by categorizing differentially expressed, p53-dependent gene targets as those with overlapping p53, OCT4 and/or NANOG-binding sites versus those bound by p53 only (Figure 6). Gene expression profiling revealed that although the average expression of the two sets are comparable, p53 gene targets that
Figure 4. Transcription of development genes is dependent on p53. (A) Volcano plot of microarray gene-expression data. Each point corresponds to RefSeq gene; in RA-treated samples with average log2-fold change compared with pluripotent hESCs and negative log10 P-value scores. Colored points correspond to genes bound by p53: significantly up- (red) or downregulated (green) p53 targets are highlighted. Target genes overlapping with damage data sets are discarded. (B) Heat map, generated for differentiation-specific p53-target genes, reveals up- or downregulated targets during differentiation compared with pluripotent hESCs. (C) The GO term analysis of differentiation-specific up- or downregulated p53-target genes is shown. (D) RT-qPCR analyses of selected genes in hESCs after 4 days of RA treatment with TP53 or control non-targeting siRNA. Error bars represent SEM from three independent experiments (*P < 0.05, **P < 0.001). For determining statistical significance, data for untreated siTP53 are compared against untreated siControl; siControl + RA 4D treatment is compared with siControl; siTP53 + RA 4D treatment is compared with siControl + RA 4D.
are shared with those bound by OCT4 and/or NANOG before differentiation are the most robustly changed (up- or downregulated) genes (Figure 6A and Supplementary Figure S6A). Consistent with the biological functions of all differentiation-specific p53 targets (Figure 4C), GO analysis for upregulated p53 targets with overlapping OCT4 and/or NANOG sites revealed genes responsible for pattern specification, embryonic morphogenesis and development (Figure 6B, and Supplementary Table S5). Downregulated p53 targets with overlapping OCT4 and/or NANOG sites are involved in mesodermal differentiation, metabolism and cell motion (Supplementary Figure S6B and Supplementary Table S5). On the other hand, p53-bound differentially expressed genes did not result in significant functional categories after GO analysis.

We performed ChIP-Seq for active (H3K27me3) or repressive (H3K4me3) histone marks in hESCs after 2 days of RA treatment, compared with pluripotent hESCs, to assess global changes in active versus repressed chromatin structure during differentiation of hESCs (Figure 6C and D). Profiling of averaged genome-wide histone modifications, 2 kb upstream and downstream of the TSS, confirmed that upregulated p53 targets, bound by OCT4 and/or NANOG in pluripotent hESCs, are highly enriched for bivalent histone marks (H3K4me3 and H3K27me3) before RA treatment. During differentiation, these p53/OCT4/NANOG-regulated genes exhibit marked loss of H3K27me3 status on gene promoter or p53RE of PTCH1 and TBX5 in hESCs treated with RA for 2 days. Error bars represent s.d. from three independent experiments (*P < 0.05, **P < 0.001).
with OCT4 and/or NANOG bound in a pluripotent state and are likely activated by p53-mediated recruitment of chromatin modifying complexes that reduce repressive histone marks.

**p53 interacts with H3K27me3 demethylases to induce expression of poised genes**

To uncover p53-mediated reduction of H3K27me3 at the promoters of developmentally poised genes, we determined whether p53 interacts with H3K27me3-specific demethylases in hESCs (Figure 7). UTX and JMJD3 specifically demethylate H3K27me3/me2 and are members of the JmjC-domain-containing family (44,45). Expression analysis during hESC differentiation revealed <2-fold, but statistically significant, induction of UTX after 5 days of RA treatment, and no obvious effect of RA on EZH2 (an H3K27 methyl transferase) expression (Supplementary Figure S7A). During the same time course, expression of JMJD3 was significantly induced as early as 2 days of RA treatment (Figure 7A and Supplementary Figure S7A). JMJD3 expression during differentiation of hESCs is partly dependent on p53, as JMJD3 induction was impaired in hESCs depleted of TP53 (Figure 7A). Analysis of proteins stably bound to fractionated chromatin, prepared from hESCs undergoing RA-induced differentiation during a time course of 5 days, revealed a marked enrichment of p53, UTX and JMJD3 that coincides with concomitant reduction of OCT4 (Figure 7B). Co-immunoprecipitation analyses of these chromatin fractions show that p53 interactions with UTX and JMJD3 are increased after RA-treatment (Figure 7C), suggesting that p53 binding to chromatin may enrich H3K37me3 demethylases at specific gene sites to alter chromatin structure.

Recent reports suggest that UTX is required for proper differentiation (46,47), and both JMJD3 and UTX are actively involved in removing H3K27me3 marks for activation of the HOXA gene cluster (44,48). Consistent with these reports, we observed UTX enrichment on HOXA1 and HOXA2 promoters, associated with a significant reduction in H3K27me3 mark during RA-induced differentiation of hESCs (Supplementary Figure S7B and C). In response to RA, UTX is significantly enriched at PTCH1,
TBX5 and HOXA1 loci (shControl), whereas no change in UTX enrichment was observed in hESCs stably depleted of p53 (shTP53) (Figure 7D and Supplementary Figure S7D). These results, combined with those summarized in Figures 5 and 6, suggest that p53 recruits H3K27me3 specific demethylases to modify repressive chromatin structure of poised genes and favor gene activation during hESC differentiation.

Divergence in p53 transcription functions between mouse and human ESCs

We determined whether our findings on p53’s role in differentiation of hESCs are evolutionarily conserved by comparison with p53 functions in mESCs at a molecular level. Recent studies support evolutionary divergence in the functions of transcription factors by similar comparisons of mESCs and hESCs (13,14). In mESCs, p53 was reported as a direct repressor of Nanog expression in response to DNA damage, suggesting a potential role for p53 during mESC differentiation (3). Recently, by comparing global p53 chromatin-interaction maps and gene expression analyses, Huang and colleagues showed that p53 both activates differentiation-associated genes and represses stem cell-specific genes in response to DNA damage in mESCs (49). This is in contrast to the highly distinct, DNA damage- and differentiation-induced profiles of p53 interactions with chromatin and regulation of gene expression in hESCs, reported here. To determine whether this is an example of interspecies disparity in p53 transcriptional functions, we compared genome wide p53–chromatin interaction data sets, generated by analysis of mESCs undergoing DNA damage (49), to our p53-chromatin binding profiles in hESCs, both in response to DNA damage and RA-mediated differentiation (Figure 8). We analyzed the mESCs p53 ChIP-Seq data by the same method as used for p53 peak calling in hESCs and found 39150 significantly enriched genomic regions, 28032 of which have homologous sequences in the human genome (Supplementary Figure S8A). We used a 10 kb window of distance from the p53-peak summit to the nearest mouse gene TSS to call a p53 target gene. The p53 binding sites that are homologous within the human
Figure 8. Comparison of p53-mediated transcriptional regulation of developmental genes in mouse and human ESCs. (A) Target genes are classified as 'Conserved' or 'Turnover' based on p53-binding sites to corresponding aligned regions in human and mouse genome. p53 binds to align regions in two species at Conserved targets; however, when p53 binding sites on orthologous genes do not align, those targets are categorized as Turnover genes. (B and C) Numbers of distinct and overlapping p53-target genes in mESCs during DNA damage (green) (49) with hESCs undergoing differentiation (blue) or DNA damage (red) and enriched GO-terms for each subset (C). (D) Circos plot of four human (hs) and mouse (ms) HOX gene clusters (HOXA, B, D and C) showing species-specific differential binding patterns of p53 (damage:Red, differentiation:Blue). Ribbons show syntenic genomic locations between mouse and human (orange:Conserved p53-targets, Yellow:Turnover p53-targets). Green tracks represent repressive H3K27me3 marks around the displayed regions in mouse and human ESCs. Tiles (black, orange or yellow) show RefSeq HOX gene annotations. Purple heat map depicts the Phastcons scores around the displayed regions. Numbers outside the circle is for specific human (hs) and mouse (mm) chromosomes. (E) Tracks represent normalized p53 sequence tag enrichments in differentiating human (top) and DNA-damaged mouse (bottom) ES cells at PTCH1, HOXAI and TBX5 genomic locations (numbers indicate distance from TSS).
that complement its surveillance functions; fewer still address roles for p53 in early differentiation (1). Genome-wide profiling and expression analyses show that RA-mediated p53-response is highly distinct from stress-responsive events occurring downstream of DNA damage in hESCs. We find that p53 promotes differentiation of hESCs by activating expression of developmental transcription factor genes involved in patterning, morphogenesis and organ development. This activated cascade of transcription factors amplifies the functional effects downstream of p53 induction beyond the transient time period when p53 protein is elevated (5). Differentiation-specific p53 gene targets in hESCs include several members of the homeodomain family (HOX, LHX, DLX, PAX) (34,35), the forkhead family of FOX genes (36), the SOX gene family (37) and members of the TBX family (50) of genes, all of which regulate a wide variety of developmental processes. In addition, p53 targets members of the CBX family, specifically CBX2 and CBX4, which are part of the Polycomb complex and are crucial for cell fate determination (51). Our integration analysis reveals several p53 gene targets are downregulated during RA-mediated differentiation, including genes that direct mesodermal differentiation (FOXO3, KLF6, HDAC5, HDAC6) and telomere repeat binding factor TERF1, associated with pluripotency (43). RA signaling directs differentiation of ESCs along a neuro-ectodermal lineage (34). Taken together, p53 may play a role in lineage determination by RA-induced p53-mediated repression and activation of specific genes in hESCs.

Remarkably, more than half of the differentiation-induced p53-bound genomic sites are coincident with binding by either OCT4 or NANOG, or both, in pluripotent hESCs, suggesting that core pluripotency factors may ‘poise’ these genes for p53-mediated activation during the earliest stages of differentiation (18). Our expression analyses confirmed that specific developmental genes, e.g. HOXA1, PTCH1 and TBX5, previously reported to be in a poised state in pluripotent hESCs and essential in differentiation (52–54), are rapidly and highly upregulated during hESC differentiation in a p53-dependent manner. During differentiation, p53 is robustly enriched at the promoters of these genes, whereas OCT4 and NANOG remain bound. Sequential-ChIP analyses confirmed that p53 localizes to and co-occupies regions bound by OCT4. Whether OCT4 is temporarily displaced by p53 at these sites to de-repress gene expression, or OCT4 plays a gene-specific role in initial activation of these genes, requires further study. A similar phenomenon has previously been reported for Sox17, which competes with Nanog for DNA binding to promote expression of development genes during differentiation of mESCs (55). Analysis of histone status around these specific genomic loci (PTCH1 and TBX5) revealed that these genes reside in repressed chromatin enriched in H3K27me3, which is decreased during differentiation of hESCs in a p53-dependent manner.

A comprehensive genome-wide analysis of H3K4me3 and H3K27me3 status indicated that a significant portion of p53 targets might be held in a poised state in pluripotent ESCs by bivalent histone modifications, which
are altered during differentiation with a gain in H3K4me3 and loss of H3K27me3. Removal of H3K27me3 marks is a key step for developmental gene activation during embryonic development (44). Processes responsible for deposition of trimethylation marks on H3K27me3 mediated by polycomb repressive complex 2 (PRC2) have been extensively studied (17,56,57). However, the enzymes that remove methyl marks from H3K27me3 were identified relatively recently. Three members of JmjC-domain containing protein family, UTX, UTY and JMJD3, were reported to specifically demethylate H3K27me3/me2; however, UTY is not active in vitro (44,45,58). UTX and JMJD3 counteract Polycomb-mediated repression to induce gene expression required for proper cellular differentiation and development (44,46,47,59–62). In addition to catalyzing loss of repressive chromatin structure, JMJD3 has been implicated in transcriptional elongation of bivalent genes (62). Depletion of UTX impairs reprogramming, whereas expression of JMJD3 acts as a barrier to generation of iPSC cells, suggesting that precise maintenance of the H3K27me3 marks is crucial for cellular differentiation (63,64). Analysis of p53 interactions with H3K27me3 demethylases, enriched in fractionated chromatin of hESCs, revealed that p53 binding to UTX and JMJD3 is enhanced during differentiation. In parallel, UTX is enriched at development gene promoters (PTCH1, TBX5 and HOXA loci) in a p53-dependent manner, which is associated with depletion of H3K27me3 marks and subsequent transcriptional activation. This suggests that p53 plays a role in modifying chromatin structure at OCT4/NANOG-poised genes, in partnership with specific H3K27me3 demethylases, to activate a developmental transcription program.

Target genes conserved in both differentiation and DNA-damage response are active in the biological pathways of cell cycle regulation, cell motion and cell death response. This finding underscores the role that cell cycle control plays in hESCs (5) and complements a large body of work that shows the importance of cell death pathways in development and in blocking tumor progression (2). Ranked by significance, the next category of downstream p53 targets in both damage and differentiation of mESCs, although binding of p53 at Nanog was not found (49). We find that p53 targets a diverse set of genes during differentiation versus DNA damage, resulting in induction-specific outcomes that largely differ. The absence of p53 binding at genes encoding major stem cell factors in hESCs suggests that p53 does not directly repress pluripotency, supporting species-specific differences in p53 functions especially at the level of cell cycle and differentiation (5,15).

We compared our genome-wide p53–chromatin interaction profiles in hESCs undergoing differentiation or DNA damage to a p53 ChIP-Seq data set in mESC undergoing DNA damage, obtained from Li et al. (49). Interestingly, we observed that p53-targets during hESC differentiation substantially overlap with those activated in mESC during DNA damage. To determine whether a p53-binding event detected in one species occurred at a corresponding aligned region in a second species, the homologous p53-gene targets in human and mouse ESCs were categorized as ‘conserved’ (aligned) or ‘turnover’ (unaligned), as described previously (32). The overlapping p53-targets during differentiation in hESC and DNA damage in mESC are enriched for development-specific functional categories, such as gene expression, embryonic morphogenesis and differentiation. Our comparative location analyses reveal that p53 activates a specific transcription program to induce developmental gene expression in hESC undergoing differentiation or mESC exposed to DNA damage. This is suggestive of inter-species disparity in p53 transcription functions as demonstrated previously for tissue-specific transcription factors (FOXA2, HNF1A, HNF4A, HNF6A and CEBPA) (32,67). Species-specific divergence in gene expression signature (14) and OCT4 and NANOG-binding sites in human versus mouse ESCs (13,68) suggests possible differences in transcription factor functions in these cell types, which may reflect the developmental stage-specific programs in mESCs and hESCs (16). Overall, these observations support the notion that p53 is central in promoting cellular differentiation in both human and mouse ESCs. Based on these results and other studies that describe pro-differentiation effects of p53 (3,5), one might expect that deletion of p53 would have considerable impact on embryonic development. However, p53-null mice appear to develop normally, although with some problems, and p53-null hESCs contribute to all three germ layers during teratoma formation in Severe Combined Immunodeficiency (SCID) mice (69). Compensation by structurally related protein family members, p63 and p73, likely occurs (70), but it is incomplete, as detailed examination of p53-null mice reveals developmental anomalies and implantation defects (71,72) [reviewed in detail in (2)]. Whether p63 or p73 isoforms act in hESCs to regulate any or all of the downstream targets of p53 is unknown.

The p53-regulated gene signatures uncovered in pluripotent, damaged and differentiating hESCs showed that p53 has specific functions and target genes in opposing self-renewal versus maintenance of cellular surveillance. The most significant category of DNA damage-specific p53 gene targets in hESCs encompasses regulation and functions of cell migration and motility, characteristics associated with metastatic carcinomas (6). Further comparisons with other data sets reveal that DNA damage-induced targets of p53 in hESCs, colon cancer-derived cells (73) and normal fibroblasts (74) are strikingly different, suggesting that p53 interactions with chromatin are dictated by stress-specific upstream signaling and cellular
microenvironment. How p53 functions are parsed among the genes that are important in development and those involved in guarding cellular integrity is not known, but likely involves the regulatory lexicon imposed by post-translational modifications of p53 and interactions with co-regulatory protein partners. The characteristics shared between specific cancers and stem cells (8), and the dysfunction of p53 in a majority of human cancers (9), support the need for greater understanding of mechanisms and pathways regulated by p53 in multiple types of stem cells.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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