RESEARCH ARTICLE

Evolution of p53 Transactivation Specificity through the Lens of a Yeast-Based Functional Assay

Mattia Lion1‡, Ivan Raimondi1‡, Stefano Donati1, Olivier Jousson2, Yari Ciribilli1, Alberto Inga1 *

1 Laboratory of Transcriptional Networks, Centre for Integrative Biology (CIBIO), University of Trento, Mattarello, Trento, Italy, 2 Laboratory of Microbial Genomics, Centre for Integrative Biology (CIBIO), University of Trento, Mattarello, Trento, Italy
‡ These authors contributed equally to this work.
* inga@science.unitn.it

Abstract

Co-evolution of transcription factors (TFs) with their respective cis-regulatory network enhances functional diversity in the course of evolution. We present a new approach to investigate transactivation capacity of sequence-specific TFs in evolutionary studies. Saccharomyces cerevisiae was used as an in vivo test tube and p53 proteins derived from human and five commonly used animal models were chosen as proof of concept. p53 is a highly conserved master regulator of environmental stress responses. Previous reports indicated conserved p53 DNA binding specificity in vitro, even for evolutionary distant species. We used isogenic yeast strains where p53-dependent transactivation was measured towards chromosomally integrated p53 response elements (REs). Ten REs were chosen to sample a wide range of DNA binding affinity and transactivation capacity for human p53 and proteins were expressed at two levels using an inducible expression system. We showed that the assay is amenable to study thermo-sensitivity of frog p53, and that chimeric constructs containing an ectopic transactivation domain could be rapidly developed to enhance the activity of proteins, such as fruit fly p53, that are poorly effective in engaging the yeast transcriptional machinery. Changes in the profile of relative transactivation towards the ten REs were measured for each p53 protein and compared to the profile obtained with human p53. These results, which are largely independent from relative p53 protein levels, revealed widespread evolutionary divergence of p53 transactivation specificity, even between human and mouse p53. Fruit fly and human p53 exhibited the largest discrimination among REs while zebrafish p53 was the least selective.
Introduction

cis-regulatory elements (CREs) are defined as regions of DNA containing a set of transcription factor binding sites (or response elements, REs) recognized by sequence-specific transcription factors (TFs). Latest evidences are highlighting that divergence in cis-regulatory sequences might underlie phenotypic diversity in evolution more commonly than divergence in the sequences of the TFs themselves. Gain, loss or changes in the affinity of TFs at the level of REs can indeed shape gene regulatory networks [1] [2]. In addition, co-evolution of cis-regulatory elements and of TF proteins is emerging as an extremely important evolutionary process, defined as cis-trans co-evolution [3].

Many important TFs, especially those involved in development, or acting as master regulators, appeared to be highly conserved at the functional level during evolution, as revealed by ectopic expression of orthologous genes and complementation assays [2] [3] [4].

Among those TFs, the tumor suppressor protein p53 emerges as one clear example of this functional conservation. p53 is one of the most studied sequence-specific TF and its inactivation frequently occurs in cancer through the expression of mutant p53 proteins [5] [6] [7]. p53 plays a crucial role in the control of genome stability, apoptosis, cell cycle, cellular senescence and angiogenesis; for these reasons it is known as “the guardian of the genome” [6]. Being a master regulator, p53 can also influence many other biological processes including autophagy, mTOR signaling, cell motility and migration, energy and amino acid metabolism, immune response, to name just a few [8] [9] [6] [7].

To modulate so many different biological responses and to control the expression of a myriad of genes, p53 has to be tightly regulated [8] [10]. p53 specifically recognizes a consensus sequence composed of two decamers that can be separated by a short spacer (5'-RRRCWWGYGG-N-

RRRCWWGYGG-3'; R = purine, Y = pyrimidine and W = adenine or thymine; N = spacer) [11] [12]. p53 binds the RE as a tetramer (dimer of dimers) and differences in RE sequences can strongly impact its transactivation capacity [13]. Many studies have investigated the role of RE sequence in transactivation potential and specificity of human p53 protein [14] [13] [12], revealing the crucial role of the core CWWG in determining the level of transactivation, with CATG leading to highest activity followed by CAAG and CTAG sequences and by CTAG associated with the weakest responsiveness. The flanking purines and pyrimidines are also very important, and, in particular, the dinucleotide motifs flanking the core lead to highest transactivation when their sequence is GG or CC and lowest when the sequence is AG or CT [13]. Interestingly, most p53 REs deviates from the optimal consensus and low-affinity REs appear to have been selected to fine-tune the activation of specific p53-induced pathways, such as apoptosis [14] [15] [13] [16]. Non-canonical p53-REs have also been annotated, consisting of 1/2- (a decamer) and 3/4-sites (a decamer + 1/4-site) [17] [18] [16].

These findings have implications when animal models are employed for the study of p53 or of transcription factors in general, as RE differences occur even between closely related species. For instance, while human and mouse p53 proteins share high sequence identity [19] and exhibit highly overlapping structure [20] as well as similar biochemical properties and transactivation potential, at least towards high affinity REs [21], evolutionary divergence is more evident at the level of p53 REs. Indeed, several genes involved in DNA metabolism and repair that are p53 targets in humans are not responsive to p53 in rodents, which could explain the differences observed in response to environmental stress, cellular damage, and cancer development [19] [22] [23]. Sources of evolutionary diversity in p53 regulatory networks between mammalian species have been identified in intronic fuzzy tandem repeats in mouse [24] and endogenous retroviruses (ERVs) in humans [25] containing p53 REs.

Here we investigated changes in transactivation specificity for p53 proteins derived from Homo sapiens and the animal models Mus musculus, Xenopus laevis, Danio rerio, Drosophila
melanogaster, and Caenorhabditis elegans. We focused on the impact of RE sequence, p53 protein level and temperature on transactivation using a yeast-based functional assay that minimizes the impact of variables such as chromatin state, promoter landscape and the influence of cofactors.

**Materials and Methods**

**Yeast strains and culture media**

Ten isogenic yeast strains (yLFM), auxotrophic for tryptophan and containing different human p53 response element (RE) sequences cloned upstream the reporter Firefly luciferase gene were tested [26][27]. The panel of reporter strains comprises five human REs from downstream target genes of p53 (p21, MDM2 P2, MMP2, BAI1, and PUMA) and five ad hoc generated variations of the p53 consensus sequences (canonical CON1, CON2, CON3 and 3/4-sites, CON-J and CON-K [16]). Three additional reporter strains were constructed starting from the previously described yLFM-ICORE strain [23] and a targeting oligonucleotide containing the sequence of the potential p53 RE from the C. elegans ced-13 target gene [28] or the two putative p53 REs from the X. laevis p21 and mdm2 target gene (S1 File). Strain construction was performed and verified as previously described [23].

Rich medium {YPDA; 1% yeast extract, 2% peptone, 2% dextrose (D-glucose), 200 mg/L adenine} was used to expand yLFM strains. Yeast cells were transformed with the appropriate pTSG-based plasmid using the lithium acetate transformation protocol [29] and selected in medium containing dextrose (2%) as carbon source and adenine (200 mg/L) but lacking tryptophan—SDtA- {0.67% Yeast nitrogen base w/o amino acids (Difco, BD Bioscience, Milan, Italy), 50ml/L of non-essential amino acids (αα), 200 mg/L adenine, 1% leucine, 1% histidine, 1% uracil, 1% lysine} [30]. For bacterial cells Luria-Bertani broth (LB, Sigma, Milan, Italy) medium supplemented with ampicillin antibiotic was used for plasmid selection.

Development of Xl_p53 and Dm_p53 expression plasmids in yeast.

pTSG plasmid [30] is based on the centromeric pRS314 plasmid [31] and contains the TRP1 selection marker, the finely-tuned inducible GAL1 promoter and the ampicillin resistance (ampR) gene for selection in bacterial cells [27].

pTSG-Hs_p53 ([Homo sapiens] [27], pTSG-Mm_p53 ([Mus musculus] [23]), pTSG-Dr_p53 ([Danio rerio]), pTSG-Cep1 ([Caenorhabditis elegans]), were already available [32]. Xenopus laevis cDNA sources were kindly provided by Simona Casarosa’s group in Trento, Italy (CIBIO).

pCS2-Dm_p53 plasmid containing the Drosophila melanogaster p53 open reading frame (ORF) was kindly provided by Lucio Collavin’s group in Trieste, Italy (L.N.CIB).

The ORF of both Xenopus laevis (Xl_p53) and Drosophila melanogaster p53 (Dm_p53) were amplified using the Q5 Hot Start High-Fidelity DNA Polymerase (New England Biolabs, Euroclone, Milan, Italy). Primers were designed to amplify the entire p53-ORF of interest and to harbor flanking regions that recombine with the homologous sequences in the multi-cloning site of the pTSG vector (Eurofins MWG Operon, Ebersberg, Germany; sequences available upon request). The p53 ORFs of interest were specifically inserted in the multiple cloning site (MCS) downstream the GAL1 promoter and upstream of transcription terminator derived from the CYC1 gene. The recombination process occurs in vivo in yeast cells through a gap repair transformation assay [33] using pTSG plasmids digested with BamHI HF and XhoI restriction enzymes (New England Biolabs). In yeast, the linear plasmid is resealed together with the PCR products by the recombination system exploiting the sequence homology at the end of the fragments.

After the in vivo cloning, plasmid DNA was recovered from yeast transformants, transformed into Escherichia coli competent cells (XL-1 Blue) using the Potassium-Calcium-Magnesium method (bakerlab/Cells.htm), extracted from E.coli transformants (QIAprep Spin Miniprep Kit,
QIAGEN), verified by DNA sequencing (BMR Genomics, Padua, Italy) and used in the yeast functional assay. All (4/4) the cDNA collected from *Xenopus laevis* presented the Tyr179Ser sequence variant, which may represent a natural polymorphism. We cannot formally exclude that this variant can have some impact on Xl_p53 transactivation specificity or temperature sensitivity.

**Development of chimeric Dm_p53, Cep-1, and Ta_p53 expression plasmids in yeast**

Two different types of chimeras were generated. In the type-one chimera, a portion of the human N-terminal region (1–63 aa, hN63, corresponding to the presence of both TADs) was fused at the 5’ of the full-length sequence of Dm_p53 or Cep-1. The type-two of chimera was instead generated replacing the predicted corresponding N-terminal regions of Dm_p53 or Cep-1 with the entire human N-terminal portion, 1–92 aa hN92. The two regions corresponding to hN63 and hN92 were amplified by PCR from the pTSG-Hs_p53, using the Q5 Hot Start High-Fidelity DNA Polymerase (New England Biolabs). Primers were designed to amplify the Hs_p53 portions of interest and to obtain amplicons that recombine at 5’ with the homologous sequences in the multicloning site of the pTSG vector and at the 3’ with the homologous sequences within Dm_p53 or Cep-1 ORF (Eurofins MWG Operon; sequences available upon request). The recombination occurs *in vivo* in yeast cells through a Gap repair transformation assay using pTSG plasmids single digested with BamHI HF restriction enzyme (New England Biolabs). The amplicons of interest were specifically inserted downstream the GAL1 promoter and, either upstream the Dm_p53 or Cep-1 ORF, or replaced to their corresponding N-terminal region. The resulting type-one chimera plasmids were named as pTSG-hN63-Dm_p53 and pTSG-hN63-Cep-1. Alternatively, the resulting type-two chimera plasmids were named as pTSG-hN92-ΔNDm_p53, pTSG-hN92-ΔNCep-1A, pTSG-hN92-ΔNCep-1B and pTSG-hN92-ΔNCep-1C. According to our alignment analysis and previous reports [34][35][36], the predicted N-terminal region of Dm_p53 encompasses residues 1–73. We identified three putative N-terminal portions for Cep-1: (A) predicted from our analysis, residues 1–164; predicted from previous study: residues 1–205 (B) [35]; residues 1–220 (C) [37]. After the *in vivo* cloning, plasmid DNA was recovered from yeast transformants and processed as described above.

**Luciferase quantitative assay**

yLFM yeast strains were transformed with the appropriate pTSG plasmid or the empty vector pRS314 (as vehicle control) and selected for 48 h at 30°C on SDtA plates [30]. Single colonies were then patched on SDtA plates and grown for additional two days at 30°C. Cells from patches were inoculated on a 96-well plate in 60 μl of media containing 2% raffinose (Fluka, Sigma-Aldrich, Milan, Italy)-SRtA- as carbon source or 2% raffinose supplemented with two different galactose (Sigma-Aldrich) concentrations (0.008% and 0.064%) to induce the GAL1 promoter and modulate p53 protein levels. Small-volume format yeast-based luciferase assay was performed after vigorous shaking for 6 hrs at three different temperatures: 24°C, 30°C, 37°C. For every transformant four different biological samples (corresponding to four patches) were tested. Following the 6hrs-incubation, 10 μl of cells were transferred on a white 384-well plate and incubated in agitation for 15’ together with 10 μl of Passive Lysis Buffer 2X (PLB 2X; Promega, Milan, Italy) to permeabilize yeast cells. The firefly luciferase activity was then measured luminescence after adding 10 μl of luciferase substrate (Bright-Glo Luciferase Assay System; Promega). Optical density (OD) was measured at 600nm and used as normalization factor to obtain Relative Light Units, followed by blank subtraction. The analysis was performed using the microplate reader Infinite M200 (Tecan, Milan, Italy). Blank controls were also included.
Phylogenetic analysis

p53 protein sequences were downloaded from OMA browser [38] based on the latest dataset. The multiple sequence alignment was performed using the alignment editor BioEdit v7.1.11 and visualized in Jalview [39]. The percentage of identity matrix was calculated using the software Clustal Omega [40]. The freeware MEGA6.0 (molecular evolutionary genetics analysis) tool [41] was used to infer radial phylogenetic trees based on the multiple sequence alignments. Phylogenetic analysis was performed using the neighbor-joining (NJ) method, and confidence limits of branch points were estimated by 1,000 bootstrap and Poisson substitution model. A 50% cut-off value for consensus tree was used to collapse nodes and re-rooting the tree. A species phylogenetic tree was originated using the NCBI Taxonomy Browser and visualized using MEGA6.0. The tanglegram algorithm, implemented in Dendroscope 3 [42] was used to compare the phylogenetic trees.

Results

p53 sequence evolution

We generated an updated comparative analysis of 47 p53 protein sequences in relation to metazoan phylogeny using the online OMA browser [38] (S1 File). Sequence alignment, pairwise comparison of sequence identity and the conservation of critical amino acids for DNA contact, oligomerization or thermodynamic stability [43] [44] were also performed on p53 sequences from *Homo sapiens* (Hs_p53, human p53) and commonly used animal models *Mus musculus* (Mm_p53, mouse p53), *Xenopus laevis* (Xl_p53, frog p53), *Danio rerio* (Dr_p53, zebrafish p53), *Drosophila melanogaster* (Dm_p53, fruit fly p53) and *Caenorhabditis elegans* (Cep-1, round worm p53) (Fig. 1 and S1 File) (Table 1 and 2). As expected, Hs_ and Mm_p53 share the highest overall identity (~78%), while for Dr_ and Xl_p53 the identity decreases to ~50% for all pairwise comparisons, whereas Dm_p53 and Cep-1 are the most divergent p53 proteins relative to chordates examined and share also less than 20% identity (Table 1A). The percentage of identity was higher considering exclusively the DBD (Table 1B and Fig. 1). The C-terminal region exhibited higher level of conservation respect to the N-terminal portion (Table 1C, Table 1D, and S1 File).

Many of the known p53 mutations hit the core domain and this further emphasizes the importance of the p53 sequence-specific transactivation function and hence of the DBD [45]. All residues that can establish direct contacts with RE sequences were invariant in the four chordate p53 proteins (Table 2A). Arg280, Lys120, Ala276 and Cys277 allow p53 binding to DNA bases within the major groove; whereas Ser241, Arg273, Arg248, Lys120 and Ala276 form direct contacts to the DNA backbone [44]. In Dm_p53 most of the residues fundamental for the DNA contact are also conserved (Table 2A), with the exception of Arg273, that is replaced by the corresponding Lys259, and of Ala276, replaced by Thr262, as previously reported [36] [46] [45]. Arg248, Arg273, Ala276 and Arg280 appeared to be conserved even in Cep-1 (Table 2A), highlighting the functional importance of these residues for DNA binding specificity, thus supporting previous data [46] [45] [37]. Looking at the conservation of zinc coordinating residues [35] [37], residues important for DBD stability [36] [44] and residues involved in the non-polar protein-protein interface [45] [44], we noticed that, among the chordates studied, all these residues are conserved (Fig. 1 and Table 2). Many of them are also conserved in Dm_p53 whereas Cep-1 was the most divergent sequence. [35] [37] [44] [45] [36]. The residues Arg175, Met243, Gly245, Arg248, Arg249, Arg273, and Arg282 are classified as either contact residues or residues that influence thermo-stability. Moreover, Met133, Val203, Asn239 and Asn268 are important for thermo-stability of the human p53 core domain, as specific mutations at
these residues can produce a “super-stable” p53 form \[44\] \[47\]. Surprisingly, in our alignment analysis these residues are not highly conserved among chordates, with the only exception of Asn239. Curiously, a methionine at position 133, corresponding to the start codon for ΔN133 p53 isoform, is present only in the human sequence, substituting a leucine that is present in all chordates.}

**Fig 1.** p53 DNA binding domain sequence alignment of Homo sapiens, Mus musculus, Xenopus laevis, Danio rerio, Drosophila melanogaster and Caenorhabditis elegans. Sequences were aligned using ClustalW tool \[74\] and the alignment was visualized using Jalview (http://www.jalview.org) \[39\]. The three shades of blue highlight different percentage agreement, respectively from darker to lighter in this order: >80%, >60%, >40%. A percentage agreement <40% is not highlighted. The alignment shows the conservation of human functional residues involved in DNA-protein interaction (solid box), zinc-binding (dashed box), DBD stability (asterisk, *), protein-protein interface (empty circle, #) and DBD thermostability (hash mark, #). Human zinc-binding and DNA-binding residue positions are also typed. On the top of the alignment, human p53 Loop, Sheet, Helix motifs (L, dashed lines; S, left-right double arrows; H, two-dimensional arrows) are also presented. Conservation and consensus graphs are shown. Conservation is visualized as a histogram with the relative score for each column. Conserved columns are indicated with an asterisk, and columns with mutations, where all properties are conserved, are marked with a plus. Consensus is displayed as the percentage of the modal residue per column. The plus symbol is used instead of displaying multiple characters in a single character space. A consensus logo is also generated and the scale of the letter is in agreement with the conservation of the residues.

\[doi:10.1371/journal.pone.0116177.g001\]
the other five species analyzed (Fig. 1 and Table 2E). This indicates a recent acquisition of ΔN133 isoform expression in the functional evolution of the p53 transcriptional network [48]. Lastly, human Val203 and Asn268 are replaced by alanine and arginine, respectively, in both Dr_ and Xl_p53.

**Yeast as a tool to investigate p53 functional evolution**

Yeast-based transactivation assay was carried out using the six p53 proteins in Fig. 1 and a total of ten luciferase reporter strains, of which five are based on natural human p53 REs and five are variations of the p53 consensus sequences (Table 3) [14] [16]. This panel of sequences was chosen taking into account the p53 consensus RE matrix shared by p53 proteins from different species [45] to scan a wide range of DNA binding affinity and transactivation potential for human p53, and to include both canonical and non-canonical REs [16]. All REs are placed in the same chromatin landscape, hence a matrix of results could be obtained where differences in

---

**Table 1. Percentage of identity matrix of p53 proteins.**

| A) whole sequence | Hs_p53 | Mm_p53 | Xl_p53 | Dr_p53 | Dm_p53 | Cep-1 |
|-------------------|--------|--------|--------|--------|--------|-------|
| Hs_p53            | 100.00 | 77.78  | 52.94  | 51.40  | 19.82  | 16.02 |
| Mm_p53            | 77.78  | 100.00 | 53.39  | 52.84  | 21.26  | 17.83 |
| Xl_p53            | 52.94  | 53.39  | 100.00 | 55.79  | 23.55  | 19.53 |
| Dr_p53            | 51.40  | 52.84  | 55.79  | 100.00 | 21.74  | 18.08 |
| Dm_p53            | 19.82  | 21.26  | 23.55  | 21.74  | 100.00 | 18.50 |
| Cep-1             | 16.02  | 17.83  | 19.53  | 18.08  | 18.50  | 100.00 |

| B) DBD sequence   | Hs_p53 | Mm_p53 | Xl_p53 | Dr_p53 | Dm_p53 | Cep-1 |
|-------------------|--------|--------|--------|--------|--------|-------|
| Hs_p53            | 100.00 | 88.44  | 67.34  | 69.85  | 24.23  | 17.09 |
| Mm_p53            | 88.44  | 100.00 | 67.84  | 71.36  | 24.23  | 18.59 |
| Xl_p53            | 67.34  | 67.84  | 100.00 | 70.85  | 24.62  | 13.50 |
| Dr_p53            | 69.85  | 71.36  | 70.85  | 100.00 | 24.74  | 16.08 |
| Dm_p53            | 24.23  | 24.23  | 24.62  | 24.74  | 100.00 | 13.30 |
| Cep-1             | 17.09  | 18.59  | 13.50  | 16.08  | 13.30  | 100.00 |

| C) N-terminal sequence | Hs_p53 | Mm_p53 | Xl_p53 | Dr_p53 | Dm_p53 | Cep-1 |
|------------------------|--------|--------|--------|--------|--------|-------|
| Hs_p53                 | 100.00 | 53.49  | 29.85  | 18.33  | 1.56   | 16.30 |
| Mm_p53                 | 53.49  | 100.00 | 27.27  | 24.14  | 3.28   | 20.93 |
| Xl_p53                 | 29.85  | 27.27  | 100.00 | 11.76  | 15.69  | 10.45 |
| Dr_p53                 | 18.33  | 24.14  | 11.76  | 100.00 | 11.11  | 15.00 |
| Dm_p53                 | 1.56   | 3.28   | 15.69  | 11.11  | 100.00 | 13.70 |
| Cep-1                  | 16.30  | 20.93  | 10.45  | 15.00  | 13.70  | 100.00 |

| D) C-terminal sequence | Hs_p53 | Mm_p53 | Xl_p53 | Dr_p53 | Dm_p53 | Cep-1 |
|------------------------|--------|--------|--------|--------|--------|-------|
| Hs_p53                 | 100.00 | 67.90  | 42.50  | 38.10  | 9.52   | 8.89  |
| Mm_p53                 | 67.90  | 100.00 | 38.16  | 27.50  | 5.00   | 4.94  |
| Xl_p53                 | 42.50  | 38.16  | 100.00 | 45.56  | 11.63  | 3.33  |
| Dr_p53                 | 38.10  | 27.50  | 45.56  | 100.00 | 8.79   | 9.09  |
| Dm_p53                 | 9.52   | 5.00   | 11.63  | 8.79   | 100.00 | 11.58 |
| Cep-1                  | 8.89   | 4.94   | 3.33   | 9.09   | 11.58  | 100.00 |

The identity matrix values were obtained using the Clustal Omega tool (https://www.ebi.ac.uk/Tools/msa/clustalo/) based on the full length sequence (A); the DBD (B); the N-terminal region (C) or the C-terminal region (D) of Homo sapiens (Hs_p53), Mus musculus (Mm_p53), Xenopus laevis (Xl_p53), Danio rerio (Dr_p53), Drosophila melanogaster (Dm_p53) and Caenorhabditis elegans (Cep-1) p53 proteins.

doi:10.1371/journal.pone.0116177.t001
Table 2. Conservation of key p53 functional residues.

|     | DNA-protein interaction |     |     |     |     |     |     |
|-----|-------------------------|-----|-----|-----|-----|-----|-----|
|     | A)                      |     |     |     |     |     |     |
|     | Hs_p53                  | Lys120 | Ser241 | Arg248 | Arg273 | Ala276 | Cys277 | Arg280 |
|     | Mm_p53                  | -      | -      | -      | -      | -      | -      | -      |
|     | Xl_p53                  | -      | -      | -      | -      | -      | -      | -      |
|     | Dr_p53                  | -      | -      | -      | -      | -      | -      | -      |
|     | Dm_p53                  | -      | -      | Lys    | Thr    | -      | -      | -      |
|     | Cep-1                   | Ser    | Lys    | -      | -      | -      | Tyr    | -      |
|     | B)                      |     |     |     |     |     |     |     |
|     | Hs_p53                  | Cys176 | His179 | Cys238 | Cys242 |
|     | Mm_p53                  | -      | -      | -      | -      |
|     | Xl_p53                  | -      | -      | -      | -      |
|     | Dr_p53                  | -      | -      | -      | -      |
|     | Dm_p53                  | -      | -      | -      | -      |
|     | Cep-1                   | Asn    | Leu    | -      | -      |
|     | C)                      |     |     |     |     |     |     |     |
|     | Hs_p53                  | Pro177 | His178 | Arg181 | Met243 | Gly244 |
|     | Mm_p53                  | -      | -      | -      | -      | -      |
|     | Xl_p53                  | -      | -      | -      | -      | -      |
|     | Dr_p53                  | -      | -      | -      | -      | -      |
|     | Dm_p53                  | Gln/Asn| Asn/-  | Ser/Val| Ile    |
|     | Cep-1                   | Leu    | Ser    | Ile    | Leu    | Asp    |
|     | D)                      |     |     |     |     |     |     |     |
|     | Hs_p53                  | Arg175 | Tyr220 | Gly245 | Arg249 | Arg282 |
|     | Mm_p53                  | -      | -      | -      | -      | -      |
|     | Xl_p53                  | -      | -      | -      | -      | -      |
|     | Dr_p53                  | -      | -      | -      | -      | -      |
|     | Dm_p53                  | -      | N/A    | Leu    | N/A    | Lys    |
|     | Cep-1                   | Ser    | Ile    | Leu    | -      | Trp    |
|     | E)                      |     |     |     |     |     |     |     |
|     | Hs_p53                  | Met133 | Val203 | Asn239 | Asn268 |
|     | Mm_p53                  | Leu    | Pro    | -      | Asp    |
|     | Xl_p53                  | Leu    | Ala    | -      | Arg    |
|     | Dr_p53                  | Leu    | Ala    | -      | Arg    |
|     | Dm_p53                  | Leu    | -      | Gln    | His    |
|     | Cep-1                   | Leu    | Met    | Gln    | Tyr    |

Presented is a summary for Homo sapiens (Hs_p53), Mus musculus (Mm_p53), Xenopus laevis (Xl_p53), Danio rerio (Dr_p53), Drosophila melanogaster (Dm_p53) and Caenorhabditis elegans (Cep-1). See also Fig. 1. Reported are residues involved in DNA-protein interaction (A); zinc-binding (B); protein-protein interactions (C); DBD stability (D) and DBD thermostability (E).

doi:10.1371/journal.pone.0116177.t002
transactivation potentials are directly dependent on the nature of the RE sequence, the type of p53 protein being expressed, the level of expression of the p53 proteins, and the growth temperature of yeast strains [27].

cDNAs for the six p53 proteins were cloned into yeast expression vectors using gap repair approaches (see Materials and Methods). To perform the functional assay, transformants were incubated for six hours in media containing two different concentrations of galactose, to achieve moderate (0.008%) and high (0.064%) p53 cDNA expression (see Materials and Methods).

A 10 RE panel to investigate the impact of different features on p53 transactivation potential

The relative transactivation potential was measured for each protein towards the 10 REs (Fig. 2) and changes in relative transactivation specificity were compared (Fig. 3). This latter comparison is largely independent from relative p53 protein expression, stability or nuclear localization and is the focus of this study. Consensus (CON) REs are artificial REs chosen to investigate more directly the impact of sequence changes in the CWWG core motif (TA, AT or AA) (Table 3 with relative predicted Kd). The non-canonical REs CON-J and CON-K were tested to examine protein capacity towards low affinity REs [49]. In particular, CON-J is composed of a 1/2 site contiguous to a 1/4 site (defined as 3/4 site), whereas CON-K consists of a 1/2 and 1/4 site separated by a 5-nt spacer (Table 3). The functional ranking of these REs with Hs_p53 (Fig. 2) confirmed previous results [16] [30]. In particular, CON-J was the most responsive followed by CON3 and CON2, while CON-J and CON-K were the weakest particularly at the lower galactose concentration. As expected, the highly conserved Mm_p53 showed comparable results with respect to the human ortholog. Xl_p53, and particularly Dr_p53, instead exhibited reduced discrimination towards the REs, more evident at the higher expression level (Fig. 2). CON-K remained the weakest RE. Dm_p53 showed very low level of activity compared to the chordate p53s tested. Surprisingly, responsiveness of CON-K was nearly comparable to CON1. Cep-1, the most evolutionary divergent p53 protein examined, was instead not functional towards all consensus REs tested.

| Table 3. List of the consensus and natural RE sequences used in our experiments. |
|-----------------------------------|-------------------|
| **CONSENSUS SEQUENCE** | **RRRCWWGYYY-RRRCWWGYYY** | **Kd** |
| CON1 | GGCGATGTCC-GGCGATGTCC | -7.56 |
| CON2 | GGCGATGTCC-GGCGTATGTC | -6.82 |
| CON3 | GGCGATGTCC-GGCGAAGTCC | -7.19 |
| CON-J | GGCGATGTCC-GGGCAC-CTTTTGTCC | -6.70 |
| CON-K | GGCGATGTCC-1GTTTGTCC | -6.48 |

| **NATURAL HUMAN p53 RE** | **RRRCWWGYYY-RRRCWWGYYY** | **Kd** |
|-------------------------|-------------------|---|
| p21 | GAACATGTCC-cAACATGTTg | -7.11 |
| PUMA | cGCAAGTCC-1GACCTTGTCC | -6.86 |
| MDM2 P2 (RE1 + RE2) | GGGCAAGTTC-AGACATGTCC-tGAGTAgTCC-TGACTTGTCT | N/A |
| MMP2 | AGACGAAGCCT-GAACTTGTCC | -7.01 |
| BAI1 | tGGC-TGCT-GGACATGTTC | -6.31 |

Reported are i) the RE name, ii) the nucleotide sequence with relative position of p53 monomer-binding sites (arrows) and iii) the calculated dissociation constant based on [49]. Non-consensus bases are reported in lowercase. n = spacer length.

doi:10.1371/journal.pone.0116177.t003
Analysis was further extended to five p53 REs derived from natural human p53 target genes (p21, PUMA, MDM2 P2, MMP2, and BAI1; Table 3). These REs contain mismatches from the consensus (except for MMP2) and differ among each other also at the Rs and Ys flanking the CWWG core. BAI-1 is a 3/4 site whereas MDM2 comprises two REs (P2) separated by a spacer that can act cooperatively in transactivation [16]. The predicted Kd's are in the same range as
for the consensus REs (Table 3). MDM2 P2 was the most responsive natural RE to Hs_p53 (Fig. 2), followed respectively by p21, PUMA and MMP2, and BAI1 as the weakest. Surprisingly, Mm_p53 was particularly active towards MMP2 but less towards MDM2 P2 (Fig. 2). Xl_p53 showed a transactivation pattern comparable to that of Hs_p53 with the exception of p21 RE, which was less active at the higher level of expression. Dr_p53 confirmed a lower degree of discrimination among REs and high activity at lower p53 expression. With the exception of BA1, transactivation was not correlated with galactose concentrations. Dm_p53 exhibited again a low overall transactivation potential, was completely inactive towards BAI1, while p21 was the most responsive RE. Cep-1 was also inactive towards all the natural human p53 REs tested (Fig. 2).

**Widespread evolutionary divergence of p53 transactivation specificity**

To compare directly relative changes in transactivation specificity for each p53 protein examined, we summarized the results from the functional assay using radar charts (Fig. 3). As a matter of fact, relative light unit (RLU) values cannot be directly compared given the lack of quantification of nuclear p53 proteins level. A visualization of relative transactivation specificity was obtained setting as 1 (100%) the results obtained for the most responsive RE with human p53 (CON1) at either level of expression. The result for human p53 was then overlaid with those obtained for the other p53 proteins (Fig. 3).

The profile of transactivation specificity was consistent for the data obtained with two concentrations of galactose inducer (Fig. 3 and S1 File), although weaker REs, such as CON2, CON-J or BA1-1 were more responsive at higher galactose concentration, possibly reflecting...
differences in p53 oligomeric state that can be affected by protein concentration. While the general conservation of transactivation potential is apparent, as most of the REs were active with the different p53 proteins, the radar plots indicated that each p53 protein tested exhibited a unique profile of transactivation specificity. \( \text{Mm}_p53 \) and \( \text{XL}_p53 \) were the most similar to \( \text{Hs}_p53 \) (S1 File). However, \( \text{Mm}_p53 \) and \( \text{XL}_p53 \) exhibited an apparent change in relative transactivation specificity that resulted in lack of discrimination between PUMA and BAI1 RE and higher relative activity towards MMP2. \( \text{Dr}_p53 \) profile resembled more the \( \text{Hs}_p53 \) one at low galactose concentration (S1 File), but had no apparent discrimination between the REs at higher galactose dose. \( \text{Dm}_p53 \) showed a unique pattern of transactivation specificity with high activity for all natural REs except for BAI-1.

**Xenopus laevis p53 is highly temperature sensitive**

Since we studied p53 proteins derived from either homeothermic (mouse and human) or poikiloithermic (frog, zebrafish and fly) animals, we decided to carry out the yeast transactivation assay both at 37°C and 24°C using the five natural p53 REs (see Materials and Methods). Results were then compared to the previous data acquired at 30°C (Fig. 4). \( \text{Hs}_p53 \) had a comparable activity at all three different temperatures tested. Unexpectedly, \( \text{Mm}_p53 \), showed some temperature sensitivity, especially at low expression levels. \( \text{Dr}_p53 \) was also temperature sensitive particularly at lower protein expression (Fig. 4), where a “ladder” effect consisting in a progressive decrease in transactivation capacity, was observed with increasing temperatures. No significant, temperature-induced differences were recorded with \( \text{Dm}_p53 \). \( \text{Cep-1} \) remained inactive also at 24°C and 37°C.

\( \text{XL}_p53 \) was the most sensitive to temperature for all the REs tested and with both galactose concentrations used, consistent with previous reports [50], whereas a significant gain in transactivation capacity was observed at 24°C (Fig. 4).

**A chimeric transactivation domain strongly enhances Drosophila melanogaster p53 transcriptional activity in yeast**

As mentioned above, despite the low level of conservation in the DBD, biochemical assays suggest that both \( \text{Cep-1} \) and \( \text{Dm}_p53 \) can bind to canonical p53 REs in vitro [36] [45]. Since the N-terminal transactivation domain is poorly conserved, and given that different types of transactivation domains may function less effectively in yeast cells [51], we generated chimeric constructs to provide an acidic transactivation domain to these p53 proteins derived from protostome animals. Hence, we created chimeric proteins for both \( \text{Dm}_p53 \) and \( \text{Cep-1} \), as described in Material and Methods. In these constructs (see also Fig. 5A), human p53 N-terminal portions were either used to replace the N-terminal of \( \text{Cep-1} \) and \( \text{Dm}_p53 \) or added at the 5’ of full-length p53 from both species. These chimeras were tested towards the five natural human p53 REs.

The two \( \text{Dm}_p53 \) chimeric proteins showed much higher transactivation capacities compared with the non-chimeric counterpart, reaching similar transactivation values as obtained with the p53 proteins from chordates (Fig. 5B and 5C). As a comparison, human p53 activity was also included. \( \text{Dm}_p53 \) chimeric proteins possess a transactivation potential similar to human p53 towards the REs p21, PUMA and MMP2. Interestingly, \( \text{Hs}_p53 \) is instead more active than the chimeric \( \text{Dm}_p53s \) towards MDM2 P2 and BAI1, indicating that the responsiveness of these chimeric constructs is likely more dependent on the affinity of their DBDs towards the REs. The human transactivation domain increased \( \text{Dm}_p53 \) transactivation capacity in our system without impacting the overall transactivation potential. Indeed, both type of chimeric constructs showed similar transactivation specificity, comparable also to the non-
chimeric Dm_p53, as highlighted by the radar plots (Fig. 5D). Taken together, these findings indicate that chimeric strategy can be used in our system to study TFs specificity.

Four different chimeric Cep-1 constructs were tested (see Materials and Methods) but all remained inactive in the transactivation assay (data not shown), as the non-chimeric construct.

Cep-1 is active towards the ced-13 p53 RE in yeast

We reasoned that one possibility for lack of Cep-1 activity could be ascribed to a markedly diverged DNA binding specificity of this protein. To date, no functional Cep-1 binding sites have been reported or annotated. Thus, we constructed a reporter strain (see Materials and Methods) based on the evidence of a potential p53-binding sites described in the promoter of ced-13, composed of two AAACATGTTT palindromic half sites separated by a 28nt spacer [28]. None of the REs previously tested in our work had these A/T-rich motif flanking the CWWG core. Surprisingly, while the strain was not responsive to Hs_p53 (as expected because of the long spacer) non-chimeric full-length Cep-1 led to modest but significant transactivation...
of the reporter (Fig. 6). To our knowledge, this is the first evidence of a functional binding site responsive to Cep-1 in transactivation assays. Hence, the lack of transactivation towards human or artificial REs by natural or chimeric Cep-1 proteins is likely to be dependent on a lack of affinity of the DBD. This result is in agreement with the hypothesis of co-evolution of p53 DBD and binding sites and warrants further investigation in the future.

Discussion

A role of cis-regulatory elements (CREs) evolution and of co-evolution of CREs and TFs in determining phenotypic diversity among species have been recently emphasized [2]. Conserved enhancer elements that can be bound by TFs are thought to be functionally essential and are usually enriched in proximity of key target genes, such as those controlling body plan development. On the other hand, lack of conservation of TF binding sites do not necessary imply lack of function but rather can underlie diversity in the regulation of gene expression. For example, the loss of a DNA binding element can be counterbalanced by the gain of a different element acting on the same transcriptional start site(s) [52]. Importantly, TF binding sites are usually degenerated. Hence, the presence of TF binding sites within orthologous promoters can be conserved but the actual sequence of the elements can differ, resulting in variation in relative binding affinities that can play an important role in expression divergence between species [2] [16]. On the other end, evolutionary divergence or conservation of gene regulatory modules can be strongly influenced by the promoter context, i.e. the gene regulatory network impacting
on the activation of a target promoter, or the integration of signals influencing biological outcomes [53]. Mobile repetitive elements can contribute to the spread of DNA binding elements, thereby contributing to the expansion of gene regulatory networks [24] [54]. Moreover, convergent evolution is another way to gain binding events in proximity of genes involved in the same functions or expressed in the same tissues between two organisms. A fascinating case linked p53-dependent regulation of genes involved in neuronal development with upstream insertion of specific transposons carrying p53 binding elements. The insertion of these elements convergently happened in both human and zebrafish genomes [55].

**Apparent evolutionary divergence of p53 transactivation specificity**

We sought out a versatile experimental system that could evaluate the transactivation potential and specificity of p53 proteins and, in principle, of any sequence-specific transcription factor. We chose a well-established yeast-based assay that minimizes the impact of variables such as chromatin state, promoter landscape and cofactors’ influences. In fact, all REs are placed in the same genomic locus, at single copy, and at a target chromosomal site upstream of the same minimal core promoter in completely isogenic reporter strains. The tool is versatile also because the p53 protein of interest can be expressed at different levels through an inducible promoter. In addition the assay can be performed in a small volume format, potentially amenable to automation [27].

Given that presently our approach does not take into account relative nuclear protein levels for the different p53 proteins expressed in yeast, comparisons can be drawn on changes in transactivation specificity of a given p53 protein towards different REs (Fig. 3). The REs were chosen to sample a wide range of sequence variability (Table 3), DNA binding affinity [49] and transactivation potential with human p53 [16]. Some of them are considered non-canonical, as they do not provide contact sites for all four monomers of the p53 tetramer [13] [56].
In our analysis, mouse and human p53 show high level of conservation, that does not necessarily preclude functionally relevant changes in protein structure and function. However, the mouse p53 DBD could be substituted by the human DBD in a chimeric transgenic model, without apparent phenotypic consequences [57]. Divergence is instead apparent for p53 REs between human and mouse genomes. Approximately two-thirds of the known human p53 binding sites are not alignable with their mouse counterparts, leading to a large divergence between the cis-regulatory networks, although the murine homologous genes still maintain the same function as in humans [52] [23]. Moreover, a class of p53 REs was identified in Alu elements and is thought to be primate-specific, an additional source of CRE differences between humans and mice [54]. Overall, all these findings support the idea of a different qualitative (target gene networks) and quantitative (strength or responsiveness) regulation of the p53 pathway between humans and rodents. Nevertheless, most p53-induced transcriptional responses, such as cell cycle arrest and induction of apoptosis, predate the common ancestor of human and rodents.

While there was undoubtedly a conservation of sequence-specific transactivation as most of the REs were responsive to all p53 proteins tested, with the exception of Cep-1, differences in transactivation specificities were apparent for each protein even in the comparison between Hs_ and Mm_p53 (Fig. 3). XL_p53 was similar in terms of relative transactivation specificity to human and mouse p53 (Fig. 3 and S1 File). A consistent trend was observed for XL_p53, Mm_p53 and particularly Dr_p53, which have higher relative transactivation potential towards lower affinity [49] (Table 3), non-canonical [16], or structurally more rigid [58] binding elements. In fact Dr_p53, especially when expressed at higher levels, lacked RE discrimination in terms of relative transactivation potential, a result not strictly related to saturating p53 protein level, as shown at 24°C. This peculiar transactivation capacity can be explained by the fact that although Dr_p53 is structurally and functionally similar to its human counterpart, it still retains some differences at key residues in the C-terminal helix, at the level of the tetramerization domain [59] [60].

Among the p53s tested in our assay, Dm_p53 was active only on a subgroup of REs. *Xenopus laevis* and, to a less extent, *Danio rerio* p53 are temperature sensitive proteins

*Thermodynamic stability has been extensively investigated for p53, in relation to its DNA-binding properties [20] [61] [47]. As a matter of fact, the melting temperatures (and consequently the stability) of p53 orthologs reflect the respective body temperature [61].

The transactivation assay confirmed severe temperature sensitivity for XL_p53. Transactivation was nearly abolished when the growth temperature of yeast reporter strains was raised to 37°C. To note, these results are in agreement with previous observations that XL_p53 is fully active at 30°C but inactive at 37°C [50]. Conversely, we also showed a clear gain in transactivation potential at 24°C. Indeed, *Xenopus laevis* standard laboratory rearing temperature is usually ~22–24°C [62] [63] and an increasing of temperature to 30°C is sufficient to activate a heat shock response [64] [65]. The observed XL_p53 temperature sensitivity could suggest a possible mechanism evolved to inactivate p53 activity when environmental temperature increases.

Dr_p53 also showed some degree of temperature sensitivity that was appreciable especially at lower protein expression and with a concomitant gain at 24°C (Fig. 4).

Thus, we propose that our assay could be specifically employed for thermo-sensitivity studies and to investigate or identify mutations altering temperature-dependent transactivation of TFs.
Chimeric Dm_p53 show higher transactivation potential without changes in specificity

*In vitro* EMSA assays showed that monomeric p53 proteins, even from distant species with overall low homology such as Cep-1 and Dm_p53, exhibit conserved DNA binding affinity (with similar dissociation constant) with respect to human p53 [61] [36] [66]. Crystal structure analysis overall confirmed this observation [67] [37]. However, the correlation between *in vitro* DNA binding affinity and transactivation potential for p53 family proteins can vary and recent data indicate that changes in the oligomeric state or in the cooperative interaction between DBD domains within a tetramer, induced by post-translational changes or due to sequence divergence, can strongly impact on transactivation specificity [14] [68] [69] [67].

Our results were not in agreement with these observations. Indeed, Dm_p53 exhibited weak transactivation activity towards some of REs tested, and Cep-1 was inactive with all the ten REs investigated (Fig. 2) It is worth noting that the assay we used does not directly measure DNA binding but rather p53-dependent modulation of transcription. This means that Dm_p53 and Cep-1 could in fact bind the REs but are inefficient or incapable to recruit transcriptional components essential for the activation of the minimal promoter driving the expression of the luciferase reporter. Earlier studies demonstrated that some classes of transactivation domains are not active in yeast [51] and this was also confirmed for human cardiac TFs [70]. To explore this possibility, and potentially solve this shortcoming of the experimental approach, we constructed chimeric proteins harboring the human p53 transactivation domains (hTADs) that are active in yeast and that either replaced the predicted corresponding N-terminal region or were added upstream the full-length sequence of Dm_p53 or Cep-1.

The activity of Dm_p53, in both chimeric proteins, was strongly enhanced. hTADs were sufficient to gain a transactivation capacity comparable to the human full length protein, even when just fused upstream to the Dm_p53 N-terminal domain. Importantly, the chimeric construct did not induce significant changes in transactivation specificity. Instead Cep-1 chimeric constructs remained inactive for transactivation. However we found wild type Cep-1 to be active towards its putative RE from the ced-13 gene (Fig. 6). These results pointed out that the sequence and structural differences between Cep-1 and the other chordate p53s are sufficient to impact its transcriptional activity, opening up possible future investigations about the evolutionary conservation of p53 DNA binding affinity and its coevolution with p53 binding sites and regulatory network. Of note, Cep-1 presents also marked structural and size differences in its C-terminal domain compared to the chordate p53 homologs. An oligomerization domain (OD) and a SAM domain are present at the C-terminal portion of Cep-1 protein, conferring a higher degree of protein stability than human p53 [61] [71].

In conclusion, the *cis*-regulatory network divergence could affect functional experiments when performed to examine the *in vivo* or *in vitro* effects of TF activation, deletion or a mutant TF. We propose that our yeast-based approach could be integrated in the plethora of tools commonly used in functional evolutionary studies as it allows robust assessment of relative transactivation specificity in a defined experimental set-up. Here we used, as a proof of concept, the tumor suppressor p53. Overall, we addressed differences in p53 transactivation capacity and specificity using p53 proteins derived from commonly used animal models. Unexpectedly, we observed widespread evolutionary divergence of p53 transactivation specificity, from Dr_p53 possessing similar transactivation potential towards different types of REs, to Hs_p53 exhibiting a wide range of transactivation potential, depending on the RE. These differences might be related to changes in the p53 transcriptional network and regulated functions in the course of metazoan evolution [36] [72] [73]. We could not investigate thoroughly the potential for p53 REs coevolution in the genomes of the six-species analyzed, which could constitute a development of this approach. In this regard, as an example we report results obtained on two putative Xl_p53
REs (S1 File). We performed an in silico pattern search analysis that led us to identify potential p53 REs in the promoter region of mdm2 and cdkn1a genes of Xenopus laevis (S1 File). This mdm2 RE was similarly responsive to both Hs_p53 and Xl_p53, whereas cdkn1a RE was inactive. These explorative results established no clear evidence for p53 protein and binding site co-evolution among Hs_p53 and Xl_p53. In the future, we also plan to compare p53 proteins derived from a larger range of species, to investigate how p53 transactivation specificity has been shaped during evolution and after the emergence of the three-gene p53 family.

Supporting Information

S1 File. Supporting Information Figures, Tables, and Legends. Contains Supporting Information on p53 protein sequence evolution. (ZIP)

Acknowledgments

We thank Drs Simona Casarosa and Licio Collavin for the cDNA sources used to clone Xenopus and Drosophila p53, respectively; prof. Daniel Huson, for his help in the Tanglegram analysis; prof. Thierry Soussi, Dr. Gilberto Fronza, Dr. Nicola Segata, Dr. Paolo Sordino and Dr. Alessandro Romanel, for critical comments on the manuscript.

Author Contributions

Conceived and designed the experiments: ML IR OJ AI. Performed the experiments: ML IR SD. Analyzed the data: ML IR SD OJ YC AI. Wrote the paper: ML IR OJ YC AI.

References

1. Maeso I, Irimia M, Tena JJ, Casares F, Gomez-Skarmeta JL (2013) Deep conservation of cis-regulatory elements in metazoans. Philos Trans R Soc Lond B Biol Sci 368: 20130020. doi:10.1098/rstb.2013.0020 PMID: 24218633
2. Rubinstein M, de Souza FS (2013) Evolution of transcriptional enhancers and animal diversity. Philos Trans R Soc Lond B Biol Sci 368: 20130017. doi:10.1098/rstb.2013.0017 PMID: 24218630
3. Wittkopp PJ, Kalay G (2012) Cis-regulatory elements: molecular mechanisms and evolutionary processes underlying divergence. Nat Rev Genet 13: 59–69. doi:10.1038/nrg3095 PMID: 22143240
4. Wagner GP, Lynch VJ (2008) The gene regulatory logic of transcription factor evolution. Trends Ecol Evol 23: 377–385. doi:10.1016/j.tree.2008.03.006 PMID: 18501470
5. Bode AM, Dong Z (2004) Post-translational modification of p53 in tumorigenesis. Nat Rev Cancer 4: 793–805. PMID: 15510160
6. Levine AJ, Oren M (2009) The first 30 years of p53: growing ever more complex. Nat Rev Cancer 9: 749–758. doi:10.1038/nrc2723 PMID: 19776744
7. Vousden KH, Prives C (2009) Blinded by the Light: The Growing Complexity of p53. Cell 137: 413–431. doi:10.1016/j.cell.2009.04.037 PMID: 19410540
8. Lion M, Bisio A, Tebaldi T, De Sanctis V, Menendez D, et al. (2013) Interaction between p53 and estri-diol pathways in transcriptional responses to chemotherapeutics. Cell Cycle 12: 1211–1224. doi: 10.4161/cc.24309 PMID: 23518503
9. Raimondi I, Ciribilli Y, Monti P, Bisio A, Pollegioni L, et al. (2013) P53 family members modulate the expression of PRODH, but not PRODH2, via intronic p53 response elements. PLoS One 8: e69152. doi: 10.1371/journal.pone.0069152 PMID: 23861960
10. Espinosa JM (2008) Mechanisms of regulatory diversity within the p53 transcriptional network. Oncogene, pp. 4013–4023. doi: 10.1038/onc.2008.37 PMID: 18278067
11. Menendez D, Inga A, Resnick MA (2010) Potentiating the p53 network. Discov Med 10: 94–100. PMID: 20670604
12. Wang B, Xiao Z, Ren EC (2009) Redefining the p53 response element. Proc Natl Acad Sci U S A 106: 14373–14378. doi: 10.1073/pnas.0903284106 PMID: 19597154
13. Menendez D, Inga A, Resnick MA (2009) The expanding universe of p53 targets. Nat Rev Cancer 9: 724–737. doi: 10.1038/nrc2730 PMID: 19776742

14. Ciribilli Y, Monti P, Bisio A, Nguyen HT, Ethayathulla AS, et al. (2013) Transactivation specificity is conserved among p53 family proteins and depends on a response element sequence code. Nucleic Acids Res 41: 8637–8653. doi: 10.1093/nar/gkt657 PMID: 23892287

15. Beckerman R, Prives C (2010) Transcriptional regulation by p53. Cold Spring Harb Perspect Biol 2: a000935. doi: 10.1101/cshperspect.a000935 PMID: 20679336

16. Jordan JJ, Menendez D, Inga A, Noureddine M, Bell DA, et al. (2008) Noncanonical DNA motifs as transactivation targets by wild type and mutant p53. PLoS Genet 4: e1000104. doi: 10.1371/journal.pgen.1000104 PMID: 18714371

17. Menendez D, Nguyen TA, Freudenberg JM, Mathew VJ, Anderson CW, et al. (2013) Diverse stresses dramatically alter genome-wide p53 binding and transactivation landscape in human cancer cells. Nucleic Acids Res 41: 7286–7301. doi: 10.1093/nar/gkt504 PMID: 23775793

18. Bisio A, De Sanctis V, Del Vescovo V, Denti MA, Jegga AG, et al. (2013) Identification of new p53 target microRNAs by bioinformatics and functional analysis. BMC Cancer 13: 552. doi: 10.1186/1471-2407-13-552 PMID: 24256616

19. Horvath MM, Wang X, Resnick MA, Bell DA (2007) Divergent evolution of human p53 binding sites: cell cycle versus apoptosis. PLoS Genet 3: e127. PMID: 17677004

20. Pagano B, Jama A, Martinez P, Akanho E, Bui TT, et al. (2013) Structure and stability insights into tumour suppressor p53 evolutionary related proteins. PLoS One 8: e76014. doi: 10.1371/journal.pone.0076014 PMID: 24124530

21. Lozano G (2010) Mouse models of p53 functions. Cold Spring Harb Perspect Biol 2: a001115. doi: 10.1101/cshperspect.a001115 PMID: 20452944

22. Gordon KL, Ruvinsky I (2012) Tempo and mode in evolution of transcriptional regulation. PLoS Genet 3: e1002432. doi: 10.1371/journal.pgen.1002432 PMID: 22291600

23. Jegga AG, Inga A, Menendez D, Aronow BJ, Resnick MA (2008) Functional evolution of the p53 regulatory network through its target response elements. Proc Natl Acad Sci U S A 105: 944–949. doi: 10.1073/pnas.0704694105 PMID: 18187580

24. Simeonova I, Lejour V, Bardot B, Bouarich-Bourimi R, Morin A, et al. (2012) Fuzzy tandem repeats containing p53 response elements may define species-specific p53 target genes. PLoS Genet 8: e1002731. doi: 10.1371/journal.pgen.1002731 PMID: 22761580

25. Wang T, Zeng J, Lowe CB, Sellers RG, Salama SR, et al. (2007) Species-specific endogenous retroviruses shape the transcriptional network of the human tumor suppressor protein p53. Proc Natl Acad Sci U S A 104: 18613–18618. PMID: 18003932

26. Resnick MA, Inga A (2003) Functional mutants of the sequence-specific transcription factor p53 and implications for master genes of diversity. Proc Natl Acad Sci U S A 100: 9934–9939. PMID: 12909720

27. Andreotti V, Ciribilli Y, Monti P, Bisio A, Lion M, et al. (2011) p53 transactivation and the impact of mutations, cofactors and small molecules using a simplified yeast-based screening system. PLoS One 6: e20643. doi: 10.1371/journal.pone.0020643 PMID: 21674059

28. Schumacher B, Schertel C, Wittenburg N, Tuck S, Mitani S, et al. (2005) C. elegans ced-13 can promote apoptosis and is induced in response to DNA damage. Cell Death Differ 12: 153–161. PMID: 15605074

29. Gietz RD, Schiestl RH, Willems AR, Woods RA (1995) Studies on the transformation of intact yeast cells by the LiAc/SS-DNA/PEG procedure. Yeast 11: 355–360. PMID: 7785336

30. Inga A, Storici F, Darden TA, Resnick MA (2002) Differential transactivation by the p53 transcription factor is highly dependent on p53 level and promoter target sequence. Mol Cell Biol 22: 8612–8625. PMID: 12446780

31. Sikorski RS, Hieter P (1989) A system of shuttle vectors and yeast host strains designed for efficient manipulation of DNA in Saccharomyces cerevisiae. Genetics 122: 19–27. PMID: 2659436

32. Monti P, Ciribilli Y, Bisio A, Foggetti G, Raimondi I, et al. (2014) DN-P63alpha and TA-P63alpha exhibit intrinsic differences in transactivation specificities that depend on distinct features of DNA target sites. Oncotarget 5: 2116–2130. PMID: 24926492

33. Flaman JM, Frebourg T, Moreau V, Charbonnier F, Martin C, et al. (1995) A simple p53 functional assay for screening cell lines, blood, and tumors. Proc Natl Acad Sci U S A 92: 3963–3967. PMID: 7732013

34. Ollmann M, Young LM, Di Como CJ, Karim F, Belvin M, et al. (2000) Drosophila p53 is a structural and functional homolog of the tumor suppressor p53. Cell 101: 91–101. PMID: 10778859

35. Schumacher B, Hofmann K, Boulton S, Gartner A (2001) The C. elegans homolog of the p53 tumor suppressor is required for DNA damage-induced apoptosis. Curr Biol 11: 1722–1727. PMID: 11696333
36. Herzog G, Joerger AC, Shmueli MD, Fersht AR, Gazit E, et al. (2012) Evaluating Drosophila p53 as a model system for studying cancer mutations. J Biol Chem 287: 44330–44337. doi: 10.1074/jbc. M112.417980 PMID: 23152666

37. Huyen Y, Jeffrey PD, Derry WB, Rothman JH, Pavletich NP, et al. (2004) Structural differences in the DNA binding domains of human p53 and its C. elegans ortholog Cep-1. Structure 12: 1237–1243. PMID: 15242600

38. Altenhoff AM, Schneider A, Gonnet GH, Dessiomoz C (2011) OMA 2011: orthology inference among 1000 complete genomes. Nucleic Acids Res 39: D289–294. doi: 10.1093/nar/gkq1238 PMID: 21113020

39. Waterhouse AM, Procter JB, Martin DM, Clamp M, Barton GJ (2009) Jalview Version 2—a multiple sequence alignment editor and analysis workbench. Bioinformatics 25: 1189–1191. doi: 10.1093/ bioinformatics/btp033 PMID: 19151095

40. Sievers F, Wilm A, Dineen G, Gibson TJ, Karplus K, et al. (2011) Fast, scalable generation of high-quality protein multiple sequence alignments using Clustal Omega. Mol Syst Biol 7: 539. doi: 10.1038/msb.2011.75 PMID: 21988335

41. Tamura K, Stecher G, Peterson D, Filipski A, Kumar S (2013) MEGA6: Molecular Evolutionary Genetics analysis version 6.0. Mol Biol Evol 30: 2725–2729. doi: 10.1093/molbev/mst197 PMID: 24132122

42. Huson DH, Scornavacca C (2012) Dendroscope 3: an interactive tool for rooted phylogenetic trees and networks. Syst Biol 61: 1061–1067. doi: 10.1093/sysbio/sys062 PMID: 22780991

43. Joerger AC, Fersht AR (2010) The tumor suppressor p53: from structures to drug discovery. Cold Spring Harb Perspect Biol 2: a000919. doi: 10.1101/cshperspect.a000919 PMID: 20516128

44. Kitayner M, Rozenberg H, Kessler N, Rabinovich D, Shaulov L, et al. (2006) Structural basis of DNA recognition by p53 tetramers. Mol Cell 22: 741–753. PMID: 16793544

45. Brandl T, Petrovich M, Joerger AC, Veprintsev DB (2009) Conservation of DNA-binding specificity and oligomerisation properties within the p53 family. BMC Genomics 10: 628. doi: 10.1186/1471-2164-10-628 PMID: 20030809

46. Dotsch V, Bernassola F, Coutandin D, Candi E, Melino G (2010) p63 and p73, the ancestors of p53. Cold Spring Harb Perspect Biol 2: a004887. doi: 10.1101/cshperspect.a004887 PMID: 20484398

47. Nikolova PV, Henckel J, Lane DP, Fersht AR (1998) Semirational design of active tumor suppressor p53 DNA binding domain with enhanced stability. Proc Natl Acad Sci U S A 95: 14675–14680. PMID: 9843948

48. Marcel V, Dichtel-Danjoy ML, Sagne C, Hafsi H, Ma D, et al. (2011) Biological functions of p53 isoforms through evolution: lessons from animal and cellular models. Cell Death Differ 18: 1815–1824. doi: 10.1038/cdd.2011.120 PMID: 21941372

49. Veprintsev DB, Fersht AR (2008) Algorithm for prediction of tumour suppressor p53 affinity for binding sites in DNA. Nucleic Acids Res 36: 1589–1598. doi: 10.1093/nar/gkm1040 PMID: 18234719

50. Bensaad K, Le Bras M, Unsai K, Strano S, Blandoing G, et al. (2003) Change of conformation of the DNA-binding domain of p53 is the only key element for binding of and interference with p73. J Biol Chem 278: 10546–10555. PMID: 12519788

51. Kennedy BK (2002) Mammalian transcription factors in yeast: strangers in a familiar land. Nat Rev Mol Cell Biol 3: 41–49. PMID: 11823797

52. Dowell RD (2010) Transcription factor binding variation in the evolution of gene regulation. Trends Genet 26: 468–475. doi: 10.1016/j.tig.2010.08.006 PMID: 20864205

53. Sullivan KD, Gallant-Behm CL, Henry RE, Fraikin JL, Espinosa JM (2012) The p53 circuit board. Biochim Biophys Acta 1825: 229–244. doi: 10.1016/j.bbabio.2012.01.004 PMID: 22333261

54. Cui F, Sirotin MV, Zhurkin VB (2011) Impact of Alu repeats on the evolution of human p53 binding sites. Biol Direct 6: 2. doi: 10.1186/1745-6150-6-2 PMID: 21208455

55. Micale L, Loviglio MN, Manzoni M, Fusco C, Augello B, et al. (2012) A fish-specific transposable element shapes the repertoire of p53 target genes in zebrafish. PLoS One 7: e46642. doi: 10.1371/journal.pone.0046642 PMID: 23118857

56. Lu WJ, Amatruda JF, Abrams JM (2009) p53 ancestry: gazing through an evolutionary lens. Nat Rev Cancer 9: 758–762. doi: 10.1038/nrc2732 PMID: 19776745

57. Besaratinia A, Pfeifer GP (2010) Applications of the human p53 knock-in (Hupki) mouse model for human carcinogen testing. FASEB J 24: 2612–2619. doi: 10.1096/fj.10-157263 PMID: 20371617

58. Beno I, Rosenthal K, Levine M, Shaulov L, Haran TE (2011) Sequence-dependent cooperative binding of p53 to DNA targets and its relationship to the structural properties of the DNA targets. Nucleic Acids Res 39: 1919–1932. doi: 10.1093/nar/gkq1044 PMID: 21071400

59. Storer NY, Zon LI (2010) Zebrafish models of p53 functions. Cold Spring Harb Perspect Biol 2: a001123. doi: 10.1101/cshperspect.a001123 PMID: 20679337
60. Joerger AC, Rajagopalan S, Natan E, Veprintsev DB, Robinson CV, et al. (2009) Structural evolution of p53, p63, and p73: implication for heterotetramer formation. Proc Natl Acad Sci U S A 106: 17705–17710. doi: 10.1073/pnas.0905867106 PMID: 19815500

61. Brandt T, Kaar JL, Fersht AR, Veprintsev DB (2012) Stability of p53 homologs. PLoS One 7: e47889. doi: 10.1371/journal.pone.0047889 PMID: 23112865

62. Nagasawa K, Tanizaki Y, Okui T, Watarai A, Ueda S, et al. (2013) Significant modulation of the hepatic proteome induced by exposure to low temperature in Xenopus laevis. Biol Open 2: 1057–1069. doi: 10.1242/bio.20136106 PMID: 24167716

63. James RS, Tallis J, Herrel A, Bonneau A (2012) Warmer is better: thermal sensitivity of both maximal and sustained power output in the iliotibialis muscle isolated from adult Xenopus tropicalis. J Exp Biol 215: 552–558. doi: 10.1242/jeb.063396 PMID: 22246264

64. Lang L, Miskovic D, Lo M, Heikkila JJ (2000) Stress-induced, tissue-specific enrichment of hsp70 mRNA accumulation in Xenopus laevis embryos. Cell Stress Chaperones 5: 36–44. PMID: 10701838

65. Mercier PA, Foksa J, Ovsenek N, Westwood JT (1997) Xenopus heat shock factor 1 is a nuclear protein before heat stress. J Biol Chem 272: 14147–14151. PMID: 9162043

66. D’Erchia AM, Tullo A, Lefkimmiatis K, Saccone C, Sbisa E (2006) The fatty acid synthase gene is a conserved p53 family target from worm to human. Cell Cycle 5: 750–758. PMID: 16582625

67. Kitayner M, Rozenberg H, Rohs R, Suad O, Rabinovich D, et al. (2010) Diversity in DNA recognition by p53 revealed by crystal structures with Hoogsteen base pairs. Nat Struct Mol Biol 17: 423–429. doi: 10.1038/nsmb.1800 PMID: 20364130

68. Ethayathulla AS, Nguyen HT, Viadiu H (2013) Crystal structures of the DNA-binding domain tetramer of the p53 tumor suppressor family member p73 bound to different full-site response elements. J Biol Chem 288: 4744–4754. doi: 10.1074/jbc.M112.408039 PMID: 23243311

69. Deutsch GB, Zielonka EM, Coutandin D, Weber TA, Schafer B, et al. (2011) DNA damage in oocytes induces a switch of the quality control factor TaPa63alpha from dimer to tetramer. Cell 144: 566–576. doi: 10.1016/j.cell.2011.01.013 PMID: 21335238

70. Reamon-Buettner SM, Cribilli Y, Inga A, Borlak J (2008) A loss-of-function mutation in the binding domain of HAND1 predicts hypoplasia of the human hearts. Hum Mol Genet 17: 1397–1405. doi: 10.1093/hmg/ddn027 PMID: 18276607

71. Ou HD, Lohr F, Vogel V, Mantele W, Dotsch V (2007) Structural evolution of C-terminal domains in the p53 family. EMBO J 26: 3463–3473. PMID: 17581633

72. Rutkowski R, Hofmann K, Gartner A (2010) Phylogeny and function of the invertebrate p53 superfamily. Cold Spring Harb Perspect Biol 2: a001131. doi: 10.1101/cshperspect.a001131 PMID: 20595397

73. Pan Y, Ma B, Levine AJ, Nussinov R (2006) Comparison of the human and worm p53 structures suggests a way for enhancing stability. Biochemistry 45: 3925–3933. PMID: 16548519

74. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, et al. (2007) Clustal W and Clustal X version 2.0. Bioinformatics 23: 2947–2948. PMID: 17846036