Review

TP53: an oncogene in disguise

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The standard classification used to define the various cancer genes confines tumor protein p53 (TP53) to the role of a tumor suppressor gene. However, it is now an indisputable fact that many p53 mutants act as oncogenic proteins. This statement is based on multiple arguments including the mutation signature of the TP53 gene in human cancer, the various gains-of-function (GOFs) of the different p53 mutants and the heterogeneous phenotypes developed by knock-in mouse strains modeling several human TP53 mutations. In this review, we will shatter the classical and traditional image of tumor protein p53 (TP53) as a tumor suppressor gene by emphasizing its multiple oncogenic properties that make it a potential therapeutic target that should not be underestimated. Analysis of the data generated by the various cancer genome projects highlights the high frequency of TP53 mutations and reveals that several p53 hotspot mutants are the most common oncoprotein variants expressed in several types of tumors. The use of Muller’s classical definition of mutations based on quantitative and qualitative consequences on the protein product, such as ‘amorph’, ‘hypomorph’, ‘hypermorph’ ‘neomorph’ or ‘antimorph’, allows a more meaningful assessment of the consequences of cancer gene modifications, their potential clinical significance, and clearly demonstrates that the TP53 gene is an atypical cancer gene.

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Facts

- p53 mutants are among the most common protein variants expressed in cancer cells.
- Classifying TP53 status in human cancer as ‘inactivated’ or ‘loss-of-function’ is misleading.
- Many mutant p53 variants are oncogenic with multiple GOF activities essential for neoplastic transformation.

Open Questions

- How does the diversity of oncogenic p53 variants contribute to the heterogeneity of the malignant phenotype?
- What is the contribution of p53 protein accumulation in human tumors to the GOF of mutant p53?
- Is there a tissue specificity of mutant p53 GOF?
- What will be the best strategy to target oncogenic p53 mutants for improved cancer therapy?
- Should the binary classification oncogene—tumor suppressor gene be replaced with a functional classification based on the activity of the mutated variant?

The development of next-generation DNA sequencing methods has led to a burst of information, including the release of the sequences of >5000 cancer genomes, an achievement inconceivable only 15 years ago.\textsuperscript{1,2} With the aim of understanding tumor development and finding novel genetic biomarkers to improve patient care or define new therapeutic targets, these analyses have led to the creation of lists describing the ‘most significant mutated genes’ in various types and subtypes of cancer.\textsuperscript{3–5} Well known cancer genes such as TP53, PIK3CA (phosphoinositide-3-kinase, catalytic, alpha polypeptide), adenomatous polyposis coli gene (APC) and KRAS (v-Ki-ras2 Kirsten rat sarcoma 2 viral oncogene homolog) are still at the top of these lists, but novel genes with potential clinical value have also been identified. However, their relatively infrequent mutation and/or association with specific tumor groups could limit their value as biomarkers or therapeutic targets.\textsuperscript{6}

In general, cancer genome studies highlight oncogenes as the most promising targets for drug development mainly because these hyperactive protein variants appear to be more easily druggable than the products of tumor suppressor genes, inactivated by heterogeneous nonsense or frameshift mutations.\textsuperscript{7} Therefore, drug discovery and development programs in academia and industry have mostly focused on kinase inhibitors and other approaches designed to inhibit activated oncoproteins, whereas relatively few therapeutic strategies to reactivate tumor suppressors have been developed to date.

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Abbreviations: LOH, loss of heterozygosity; LOF, loss-of-function; GOF, gain-of-function; AML, acute myeloid leukemia; CHIP, C-terminal of Hsc70 Interacting Protein; MQ, methylene quinolinone; RETRA, reactivation of transcriptional reporter activity; KRAS, v-Ki-ras2 Kirsten rat sarcoma 2 viral oncogene homolog; IDH1, isocitrate dehydrogenase 1; PIK3CA, phosphoinositide-3-kinase, catalytic, alpha polypeptide; TP53, tumor protein p53; PTEN, phosphatase and tensin homolog gene; APC, adenomatous polyposis coli gene

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The first evidence for the existence of tumor suppressor genes came from cell fusion experiments performed by Harris et al.8 in the 1960s, showing that fusion of normal cells with tumor cells resulted in a normal phenotype. Loss of specific chromosomes from the hybrid cells led to reappearance of the tumor phenotype, suggesting the existence of critical genes able to suppress the tumor phenotype. The tumor suppressor gene concept was also supported by Knudson’s epidemiological studies on familial and sporadic retinoblastoma and the subsequent loss of heterozygosity (LOH) analysis that identified chromosomal deletions in tumors.9,10 Haber and Harlow defined tumor suppressor genes as ‘genes that sustain loss-of-function (LOF) mutations in the development of cancer’.11 This classification has evolved over time and novel classes such as caretakers, gatekeepers or landscapers have been added to take into account the function of novel genes and their association with tumor development.4,12 Although oncogene and tumor suppressor gene alterations may have direct consequences on cell growth, caretakers and landscapers act indirectly by promoting either genetic instability or an abnormal cellular environment that will foster neoplastic transformation. This classification, based on the diversity of the functional activity of the wild-type product, should not be confused with the consequences of their alteration.

For example, LOF is still the main characteristic of tumor suppressor genes, but the Haber and Harlow definition does not take into account the heterogeneity of the different protein variants or other activities such as dominant negative (DN) effects on wild-type protein or gain-of-function (GOF). Furthermore, for some genes such as Notch, this definition can be cell-type specific, as the spectrum of mutations is different in blood and solid tumors, with different consequences for the protein.13,14 Although the two-box oncogene/tumor suppressor categorization remains a central concept in tumor biology, it is becoming increasingly obvious that it is difficult to put some genes into one box. As cancer research develops rapidly, it is essential to ensure highly flexible data classification so that labels do not paralyze the emergence of new concepts.

The status of TP53 as a tumor suppressor gene can be used as a paradigm. Wild-type TP53 clearly acts as a negative regulator of cell growth, but considering TP53 mutations solely as LOF mutations would prevent a full understanding of how TP53 mutations drive tumor growth, as TP53 status in human cancer is often defined in binary terms, wild-type versus inactivated, despite accumulating evidence that the majority of mutant p53 proteins are heterogeneous oncogenic proteins with multiple GOF activities and with potential as therapeutic targets.

In this review, we will show that, despite the important diversity of TP53 mutations, some specific p53 mutations are among the most frequent variants expressed in human cancer. In addition, we have gathered various lines of evidence accumulated over 35 years of research that illustrate how TP53 is an out-of-the-box entity. Finally, using Muller’s classical definition of mutations based on quantitative and qualitative consequences on the protein product, we will discuss how this classification would be more appropriate to define the heterogeneous effects of cancer gene mutations.

Pattern of TP53 mutations in human cancer: oncogenic hotspot mutant TP53 is one of the most frequently expressed protein variants in human cancer

A unique feature of the TP53 gene compared with other tumor suppressor genes is its mode of inactivation.15–18 More than 80% of somatic and germline TP53 alterations are missense mutations that lead to the synthesis of a stable mutant protein that accumulates in the nucleus of tumor cells (Figure 1). This high frequency of substitutions is highly analogous between the various types of cancer despite a different spectrum of mutation due to variable exposure to carcinogens.

Oncogenes are typically activated by missense mutations that target specific key residues of the protein (Figure 1). Tumor suppressor genes display either out-of-frame insertions and deletions (indel) or nonsense mutations, both leading to loss of protein expression (Figure 1). These patterns of cancer-associated mutations have been used to classify cancer genes as either oncogenes or tumor suppressor genes in the absence of data on the functional consequences of the mutations. In a recent review on cancer genomics, Volgelstein et al.4 proposed a ratiometric method to classify cancer genes based on the frequency of recurrent mutations and type of alterations (missense versus nonsense or frame-shift). Although their method can separate classical oncogenes from tumor suppressor genes in general, the TP53 gene would still straddle the boundary between the two classes of genes.

The strong selection to maintain expression of full-length p53 protein in tumors is highly suggestive of a vital role in transformation, including DN activity and/or GOF. As >90% of TP53 mutations are localized in the core domain of the protein (residue 100 to 300), all p53 protein isoforms are affected by these alterations.19

The distribution of mutations in the p53 protein is also unique among all cancer genes, including oncogenes and...
tumor suppressor genes. All but 7 residues of the 393 amino acid residue p53 protein have been the target of at least 1 mutation in human cancer, and in the core domain that contains the DNA-binding region, each residue has been found to be mutated at least 5 times in independent tumors, and up to >2,000 times for hotspot mutants. This vast scattering of TP53 mutations is due to the marked fragility of the core domain, which can be destabilized by amino acid substitutions at many different positions.

The Cancer Genome Atlas Pan-Cancer project has released an integrated set of genomic data from 3,200 cancer patients with 12 tumor types. The data include genomic, epigenomic, transcriptional and proteomic information. Not surprisingly, TP53 was found to be the most frequently mutated gene in this new set of data (Figure 2 and Supplementary Figure 1). This leading position for TP53 was already obvious before the various cancer genome sequencing projects were launched and it remains unchallenged today, as most novel cancer genes are either cancer-specific (IDH1 and IDH2), mutated at low frequency (FBXW7 or GATA3) or both. One of the drawbacks of using a classification based on gene mutation frequency is that it does not take into account the diversity of the different protein variants for each gene. Although KRAS mutations are restricted to a few codons, those in other genes may be scattered along the coding region, which leads to multiple protein variants and hinders potent information on mutant diversity. Using mutational data released by the Pan-Cancer project, we have been able to perform an integrated analysis focusing directly on each protein variant (Figure 3). Three PIK3CA variants (p.H1047R, p.E545K and p.E542K) were the most frequent mutants found in the 12 tumor types; they were found in 9.9% of patients, corresponding to ~300,000 cancer cases worldwide. This observation clearly supports the importance of this gene as a target for therapy.

Mutant p53 heterogeneity: LOF and GOF

Before addressing the various GOF activities of mutant p53, it is essential to discuss the reasons for one of the most striking

Figure 2  Genes most frequently mutated in various types of cancer in the Pan-Cancer study. Only the 40 most significantly mutated genes in the Pan-Cancer study are shown on this graph. The PAN-CANCER study included glioblastoma multiforme (GBM), lymphoblastic acute myeloid leukemia (LAML), head and neck squamous carcinoma (HNSC), lung adenocarcinoma (LUAD), lung squamous carcinoma (LUSC), breast carcinoma (BRCA), kidney renal clear-cell carcinoma (KIRC), ovarian carcinoma (OV), bladder carcinoma (BLCA), colon adenocarcinoma (COAD), uterine cervical and endometrial carcinoma (UCEC) and rectal adenocarcinoma (COADREAD). Pan-cancer: integrated data with all cancer types. Data were generated by analysis of the mutations released by Kandoth et al.
features of mutant p53, namely p53 protein accumulation in tumor cells. This topic was overlooked for a long time, perhaps due to the widespread belief that this feature was inherent to the mutant protein which displayed an increased half-life. More recently, this issue has been revisited, as targeting mutant p53 accumulation could be a very promising approach for the development of therapy.

The increased levels of nuclear p53 in tumor cells were described more than 3 decades ago but the relation with TP53 mutation became clear 10 years later and immuno-histochemistry analysis of tumors using various p53 monoclonal antibodies was developed as a surrogate for p53 variant analysis. Several observations argued that p53 protein accumulation was not an intrinsic property of the TP53 protein. Approximately 70% of individuals with the Li-Fraumeni syndrome carry a heterozygous germline mutation in the TP53 gene and express both wild-type and mutant p53 in their normal cells. Despite this genotype, accumulation of mutant p53 protein can only be visualized in tumors, whether or not the wild-type p53 allele is retained.

Terzian et al. used a knock-in mouse model to analyze the expression of mutant p53 in various tissues. Their first observation was that mouse tumors behaved like human tumors; they displayed mutant p53 accumulation in the nucleus, whereas mutant p53 was undetectable in normal cells. This feature was observed in both heterozygous (one mutant and one wild-type allele) and homozygous (two mutant alleles) animals. This is an important finding because it rules out the possibility that the inability of mutant p53 to act as a transcription factor and induce sufficient levels of MDM2 protein, a p53-specific E3 ubiquitin ligase that targets p53 for modification and subsequent degradation by the 26 S proteasome, is a cause for mutant p53 hyperstability.

However, additional ubiquitin ligases have recently been identified that participate in the degradation of the tumor suppressor, including Pirh2 (p53-induced protein with...
a RING-H2 domain), Trim24 (transcription intermediary factor 1-alpha), COP1 (constitutive photomorphogenesis protein 1) and CHIP (C-terminus of Hsc70 interacting protein). The CHIP E3 ligase participates in p53 degradation by forming a large complex that includes HSP90 and HSP70. Mutant p53 forms stable complexes with both HSP, CHIP and MDM2, inhibiting the ligase activity of both. A large survey of tumor cell lines expressing wild-type and mutant p53 showed that stabilization of endogenous mutant protein is due to a complete lack of ubiquitination. Inhibition of HSP90, either via knockdown or by specific drugs, alleviates the formation of such complexes, leading to reactivation of both MDM2 and CHIP, and mutant p53 degradation.

The heterogeneity of p53 mutants was discovered 25 years ago with the description of the so-called 'structural' mutants, in which the DNA-binding domain is unfolded (e.g. p.R175H), and 'DNA contact' mutants, in which residues interacting directly with DNA are substituted (e.g., p.R273H). Nuclear magnetic resonance, circular dichroism and X-ray diffraction analyses have confirmed and expanded this description and suggested that multiple thermodynamic stages are associated with the various mutations depending on their position. This heterogeneity extends to the biological activity of mutant p53 proteins. Analysis of the transcriptional activity of 2500 p53 mutants on 8 different target genes representative of different TP53 functions demonstrated heterogeneous penetrance of the various p53 mutants with >50% presenting only partial loss of activity.

Mutant p53 GOF has been extensively investigated in multiple in vitro and in vivo systems (see refs 18 and 44–47 for reviews). GOF was hypothesized as early as 1993, when Dittmer et al. showed that the introduction of different p53 mutants into TP53-null cells resulted in an oncogenic phenotype such as enhanced tumorigenic potential. It is also possible to date GOF several years earlier, when mutant TP53 misidentified as wt TP53 was shown to transform cells. Whether or not reinterpretation of these results in the light of new data can predate subsequent observations is an endless semantic debate. Further studies have largely supported this observation. Mutant p53 GOF includes enhanced tumorigenesis, metastasis, resistance to therapy and genomic instability. The genomic instability effect was further supported in a recent analysis of 3000 tumors showing that TP53 mutations were strongly associated with a high frequency of copy number changes. GOF in p53 mutants is highly heterogeneous, an observation that may be related to the various conformations of p53 mutants. GOF mechanisms may be the result of changes in the specificity of the DNA-binding activity of the p53 mutant, leading to the induction of novel transcriptional programs, or changes of its interaction with other cellular proteins directly or indirectly related to the regulation of gene expression. Several transcriptional programs are specifically activated by p53 mutants, most of them resulting in increase of tumorigenicity but whether this is mediated by a direct binding of the mutated protein to a specific DNA sequence or via an interaction with other transcription factors remains to be elucidated (see ref. 53 for review). A recent study from Myers and co-workers shows that several p53 mutants exert a transcription-independent GOF by downregulating the AMPK (AMP-activated protein kinase), an observation adding one more grain to the important relation between TP53 and metabolism. This observation is also in line with the study by Zhang et al., showing that p53 mutants stimulates the Warburg effect in cancer cells.

The discovery that the p53 family consists of three members (TP53, TP63 and TP73) increased the complexity of this network, as the two p53 homologs might also contribute to its oncogenic potential. TP63 and TP73 are both expressed as many isoforms due to alternative usage of promoters for transcription and alternative splicing. Long isoforms (p73 or p63 containing the transactivation domain (TA-p73 or TA-p63)) are able to transactivate the same target genes as p53 and induce apoptosis, while short forms (amino-deleted p63 or p73 isoforms (DN-p63 or DN-73)) have an opposite activity via DN mechanisms. TP63 and TP73 are able to cooperate with TP53 to induce apoptosis, suggesting the existence of a complex network of interactions between the products of these three genes. p53 mutants with unfolded structure, but not DNA contact mutants, bind specifically to p63 and p73 and impair their apoptotic activity (see refs 58–60, 61 for review).

An important issue that has never been fully analyzed is the consequence of accumulation of mutant p53 protein in tumor cell nuclei. Studies have shown that this accumulation may be as much as 10-100-fold higher than that of wild-type protein, which raises questions concerning the specificity of some mutant p53 activities and how some nonspecific squelching effects may be associated or confounded with these novel properties. Furthermore, this aspect also raises the issue of tumor heterogeneity, as p53 accumulation in human tumors is highly heterogeneous, even for single p53 variants, and could be associated with the genetic background of the tumor and the individual.

The general idea that loss of transcription in mutant p53 is the driving force selected during tumorigenesis also needs to be reevaluated, as it is far from straightforward. Surprisingly, mice expressing p53 mutants transcriptionally defective for the three canonical pathways, growth arrest, senescence and apoptosis, are not prone to cancer. Cells from mice deficient for the three major TP53 target genes, CDKN1A (the gene encoding PUMA) and PMAIP1 (the gene encoding Noxa) (p21 null/−/− mice) are deficient in their ability to undergo p53-mediated apoptosis, G1/S cell-cycle arrest, and senescence, but these animals also remain tumor-free. Taken together, these observations suggest that TP53 driver mutations are selected to impair specific TP53 pathways that remain to be identified, and only mutants defective in such transcriptional programs will be selected in human tumors.

**Mouse models.** The first TP53 knockout mice were generated by removing exons encoding the DNA-binding domain, thus impairing the expression of all p53 isoforms. Homozygous TP53−/− mice are highly prone to cancer, in particular T-cell lymphoma and sarcoma. Although these knockout mice supported the model of TP53 as a tumor suppressor gene, they were not fully satisfactory, since most cancers in humans are carcinomas, which were observed at very low frequency in TP53 knockout mice.
Improvements in the production of transgenic mice led to the creation of knock-in mice expressing various hotspot mutants that include structural or DNA contact mutants and result in more ‘human-like’ tumors.\(^{69,70}\) Compared with TP53\(−/−\) mice, these novel knock-in models displayed a higher degree of heterogeneity in the spectrum of tumor types with more frequent carcinomas. Furthermore, these tumors were highly invasive and metastatic, a feature absent in TP53\(−/−\) mice. Two characteristics of these mice models support the notion of heterogeneous GOF for p53 mutants. First, the spectrum of tumors differed according to the various mutant alleles used in these mice. For example, the ‘DNA contact’ mutant p.R270H (p.R273H in humans) results in a high frequency of carcinomas, whereas the ‘structural’ mutant p.R172H (p.R175H in humans) predominantly results in osteosarcomas. Second, the tumor spectra of heterozygous TP53 p.R172H/− or TP53 p.R270H/− mice were different from those of TP53\(+/−\) or TP53 p.R172H/− mice, which argues against any differences due to a DN activity toward wild-type p53. Some p53 mutants such as p.R175P are defective in activating genes associated with apoptosis without impairing growth arrest.\(^{71}\) Onset of tumors in knock-in mice expressing this mutant (p.R172P in mice) is delayed compared with TP53\(−/−\) mice or mice expressing hotspot mutations.\(^{72}\) In contrast to tumors expressing other p53 mutants, these tumors were mostly diploid, indicating that the growth arrest function of TP53 could be essential for maintaining genetic stability, but is not the primary tumor suppressor function of wild-type TP53.

**Targeting mutant p53 for novel cancer therapy.** Many investigators have initiated efforts to develop novel strategies for pharmacological reactivation of mutant p53 in cancer cells.\(^{73,74}\) This is a major challenge, for several reasons. Mutant p53 is clearly a different kind of target compared with those of successful novel anticancer agents such as Herceptin (trastuzumab) and Gleevec (imatinib) that block critical oncogenic kinases overexpressed in various tumors. In the case of mutant p53, the main aim is to refold and reactivate a dysfunctional tumor suppressor (Figure 4). Moreover, the protein target in this case is a DNA-binding transcription factor, a type of protein target that has been considered not easily ‘druggable’. To make things more complicated, the structural heterogeneity of mutant p53 proteins raises concerns as to the feasibility of designing or identifying therapeutic agents that can rescue more than one specific mutant form of p53 or a subset of mutants. However, the realization that mutant p53 can have GOF activity may open possibilities for therapeutic strategies designed to inhibit these functions, which may be easier to achieve than full restoration of wild-type function.

Researchers have approached the challenge of mutant p53 reactivation in various ways (Figure 4). Fersht and co-workers\(^{76}\) used detailed structural information from NMR and/or X-ray crystallography as the starting point for rational design of small molecules that can bind and stabilize the wild-type conformation of the p.Y220C hotspot p53 mutant, which occurs in \(~75\) 000 cases of cancer per year worldwide. Structural studies have shown that the substitution of cysteine for tyrosine at position 220 gives rise to a destabilizing crevice in the p53 protein. Fersht et al.\(^{76}\) have designed compounds such as the carbazole derivative PK083 that bind to the crevice and raise the melting temperature of the mutant protein.

The compound PK088 also raises the melting temperature of the p.Y220C mutant, and triggers cell-cycle arrest and apoptosis in tumor cells in a p.Y220C-dependent manner.\(^{77}\) PK088 increases the fraction of correctly folded p.Y220C mutant p53 in cells and enhances expression of the TP53 targets p21 and Noxa. Synergy with the MDM2-inhibiting compound Nutlin-3 was also observed, supporting the proposed mechanism of action.

NSC319726 (ZMC1) is another mutant-specific p53 reactivator. Carpizo and co-workers\(^{78}\) identified this thiosemicarbazone compound based on analysis of the NCI database and showed that it specifically targets the p53 hotspot mutant p.R175H. NSC319726 can restore wild-type conformation and function to this mutant and trigger p.R175H-dependent cell death by apoptosis.\(^{78}\) Moreover, NSC319726 suppresses growth of tumor xenografts expressing the p.R175H mutant in mice. NSC319726 has zinc ion-chelating properties that seem to be important for its ability to reactivate p.R175H mutant p53. A subsequent study provided further data on the zinc-metallochaperone activity of NSC319726, and showed that the compound can rescue several other p53 mutants with impaired zinc binding, for example, p.G245S.\(^{79}\) This study also highlighted the ability of NSC319726 to induce reactive oxygen species (ROS) in cells.

The small molecules PRIMA-1 (p53 reactivation and induction of massive apoptosis-1) and MIRA-1 (mutant p53-dependent induction of rapid apoptosis-1) were identified in a cell-based screen of the NCI Diversity set.\(^{80,81}\) PRIMA-1 and its structural analog PRIMA-1Met (APR-246) refold mutant p53, enhance expression of several TP53 targets, including Bax, Puma and Noxa, and inhibit human tumor xenografts in SCID mice and mouse tumors in syngeneic hosts. These effects are observed with a range of mutant p53 proteins. APR-246 synergizes with several chemotherapeutic drugs, for
example, adriamycin and cisplatin, to induce mutant p53-dependent tumor cell death. Both PRIMA-1 and APR-246 are converted to the Michael acceptor methylene quinuclidine (MQ) that binds covalently to cysteines in the p53 core domain. MQ binding to p53 is sufficient for mutant p53 reactivation, as shown by protein transfer experiments with APR-246/MQ-treated recombinant p53. It is currently not clear to which p53 cysteines MQ binds, although docking simulations and functional studies in cells have indicated that C124 is one possible target. APR-246 has been tested in a phase I/II clinical study in patients with hematological malignancies or hormone-refractory prostate cancer. This study showed that APR-246 is safe and has a favorable pharmacokinetic profile. Biological effects consistent with p53 reactivation were observed in patient leukemic cells after treatment, and clinical responses were observed in two patients, including a 10-year-old boy with a p53 V173M core domain mutation.

In addition to targeting p53, APR-246 has been shown to inhibit thioredoxin reductase (TrxR1) and convert the enzyme to an NADPH oxidase, thereby increasing cellular ROS levels. Moreover, Tessoulin et al. found that APR-246 can decrease GSH levels and thus impair the redox balance in multiple myeloma cells independently of p53.

Various other small molecule reactivators of mutant p53 have also been identified, including CP31398, Ellipticine, P53R2, SCH529074 and stictic acid reviewed in the study by Bykov and Wiman. Their mechanisms of action are not fully understood and they have not yet been tested in the clinic.

In light of the growing evidence in favor of GOF activities of mutant p53, a more modest but still attractive aim for therapeutic targeting of mutant p53 is inhibition of the oncogenic properties of mutant p53 (Figure 4). This may be easier to achieve than full restoration of wild-type function. One such approach consists of disruption of mutant p53 binding to TP63/p73. This has been shown for the small molecule RETRA (reactivation of transcriptional reporter activity). Disruption of complexes between mutant TP53 and TP63/p73 by RETRA leads to restoration of expression of TP53 target genes and tumor suppression.

A significant fraction (8%) of TP53 mutations are nonsense mutations that give rise to expression of a truncated and inactive p53 protein. The c.637C>T (p.R213*) mutant is the most common nonsense mutation in TP53, and is actually more common than many missense mutations in human tumors (see also http://p53.free.fr). Restoration of nonsense TP53 mutations will obviously require different approaches than those described above for missense mutations. Interestingly, aminoglycosides, for example, gentamicin and G418, have been shown to induce read-through of the c.637C>T (p.R213*) mutant p53 and expression of full-length p53. Although clinical use of these drugs is limited by their toxicity, the results nonetheless demonstrate that induction of read-through of premature stop codons in nonsense mutant p53 is feasible and suggest that high throughput screening for more efficient and less toxic read-through-inducing compounds should be carried out. Questions remain as to the activity of the full-length p53 protein induced on translational read-through and to what extent pharmacological induction of read-through of premature stop codons will induce read-through of natural stop codons. If these problems can be solved, induction of read-through could be a useful strategy not only for reactivation of nonsense mutant p53, but also for reactivation of other tumor suppressors that are frequently inactivated by nonsense mutations, for example APC and PTEN (phosphatase and tensin homolog gene).

**Cancer gene classification.** In Muller suggested a classification of mutations based on quantitative analysis of wild-type ‘characters’. He proposed the terms ‘amorph,’ ‘hypo-morph,’ ‘hyper-morph,’ ‘neomorph’ or ‘antimorph’. The significance of these terms was subsequently modified, as they were proposed at a time when the relationship between a gene and its product was not clearly established. These terms can now be used to classify cancer genes to more clearly understand the consequences of cancer gene alterations. This system has enormous benefits, as it is based on the outcome of the mutation regardless of the initial biological function of the gene. It could greatly facilitate the design of optimal drug development strategies.

The term amorph (or LOF) mutation can be applied mainly to tumor suppressor genes, whose function(s) must be totally impaired to drive tumorigenesis, whereas a hypomorph mutation only leads to partial reduction of activity. From a genetic point of view, true amorphic alterations can be easily associated with genes that sustain biallelic deletions such as PTEN or retinoblastoma gene (RB1). Other mechanisms, such as promoter methylation, loss of expression via microRNA dysregulation or frameshift and nonsense mutations, can also lead to total LOF, but in many cases it is difficult to exclude residual activities that will lead to heterogeneous hypomorphic variants, and for many genes it is likely that true amorphic variants are not as frequent as hypomorphic variants (Table 1). For the APC gene, the size of the truncated proteins has a major influence on several clinical features or genetic events that target the remaining wild-type allele, suggesting that several mutants are hypomorphic. Dosage reduction via haplodeficiency can also be included among hypomorphic mutations. Although many hotspot TP53 mutations are transcriptionally inactive, the majority of the remaining TP53 missense variants are hypomorphic, as they display heterogeneous loss of transcriptional activity.

Antimorphic or DN mutations, as subsequently described by Herskowitz, have been defined as disruption of the wild-type activity by the mutant polypeptide (Table 1). DN activity was initially defined by hetero-oligomerization of wild-type and mutant protein alleles leading to the formation of an inactive hetero-oligomer, a definition which is restricted to proteins with potential oligomerization activities. The use of the term ‘DN activity’ has been largely extended to any type of indirect inactivation of the wild-type allele function or its pathway via the product of the mutant allele. This broad definition can take into account the consequences of multiple mechanisms such as binding and sequestration of cofactors, limiting their availability or the occupancy of transcription promoters by factors that contain inactive transcription domains and therefore act as repressors. Furthermore, DN activity is fairly difficult to assess and distinguish from haplo-insufficiency or GOF. Only mouse models using a combination of knock-in or knockout mutations can resolve these issues. Multiple DN activities have been described for tumor suppressor genes but...
only for a few genes, such as TP53 or WT1 (Wilm's tumor 1), has it been interpreted in terms of a specific interaction between the wt and mutant allele. The formation of heterotetramers between wild-type and mutant p53 has been well documented and many in vitro analyses indicate that the activity of poised tetramers, based on transcription assays, is severely impaired. Formation of p53 tetramers obeys a particular kinetic with rapid formation of dimeric molecules followed by slow association of these dimers into tetramers.94 Consequently, poised dimers that would contain either a single wild-type or mutant subunit have not been observed and only wt2/Mut2 tetramers are formed. Several mouse models have confirmed the DN activity of certain p53 mutants towards the protein expressed by the wild-type allele, including the hotspot mutant p.R175H.95 It remains to be explained why the second TP53 allele, localized on chromosome 17p, is so frequently deleted in human and mouse tumors. Either the DN effect is not complete and full inactivation of TP53 is mandatory for tumor progression, or other genes, localized in the vicinity of TP53, are the true targets of these LOH events. Other tumor suppressor genes such as eukaryotic translation initiation factor 5A (EIF5A) or the potassium channel tetramerization domain containing 11 proteins (KCTD11) are co-deleted with TP53.96 Analysis of tumors at this transition from TP53 Mut/wt to TP53Mut/− would be highly informative in this regard.

The 'hypermorph' label is the most explicit term to describe many protein variants expressed from mutated oncogenes, as most of them are hyperactive proteins. For amplified genes such as N-myc, MDM2 or CCND1 (the gene encoding cyclin D1), or genes upregulated via chromosomal translocations such as c-myc, this definition appears to be obvious at first glance, as it refers to the abnormal accumulation of wild-type proteins (Table 1). Nevertheless, it is possible that, at a certain threshold level, the accumulated protein will impair unrelated

| Protein | Amorph | Hypomorph | Antimorph | Hypermorph | Neomorph | Mutation type |
|---------|---------|-----------|-----------|-----------|-----------|---------------|
| TP53    | Yes     | Yes       | Yes       | Yes       | Yes       | Mis, N, F    |
| Rb1     | Yes     | Yes       | No        | No        | No        | D, Mis, N, F, S |
| CDKN2A  | Yes     | No        | No        | No        | No        | D, Mis, N, F, S, Met |
| PTEN    | Yes     | No        | Yes       | No        | No        | D, Mis, N, F, S, Met |
| MLH1    | Yes     | No        | No        | No        | No        | Mis, N, D, F |
| FBXW7   | Yes     | No        | No        | No        | No        | F, N, S |
| GATA3   | Yes     | No        | No        | No        | No        | D, N, F, S |
| KDM6A   | Yes     | No        | No        | No        | No        | N, F, S |
| STAG2   | Yes     | No        | No        | No        | No        | Mis, N, F, S |
| STK11   | Yes     | No        | No        | No        | No        | D, Mis, N, F |
| RPL22   | Yes     | No        | No        | No        | No        | F |
| RUNX1   | Yes     | No        | No        | No        | No        | T, Mis |
| APC     | Yes     | Yes       | No        | No        | No        | D, Mis, N, F, Met |
| NF1     | Yes     | No        | No        | No        | No        | D, Mis, N, F |
| ARID1A  | Yes     | No        | No        | No        | No        | N, Mis, F, S, D |
| BAP1    | Yes     | No        | No        | No        | No        | N, Mis, F, S, O |
| CDH1    | Yes     | No        | No        | No        | No        | N, Mis, F, S |
| MAP2K4  | Yes     | No        | No        | No        | No        | D, Mis, N |
| MLL3    | Yes     | No        | No        | No        | No        | Mis, S, F |
| PRDM1   | Yes     | No        | No        | No        | No        | Mis, N, F, S, D, S |
| BRCA1   | Yes     | Yes       | No        | No        | No        | D, Mis, N, F, S, Met |
| VHL     | Yes     | Yes       | No        | No        | No        | D, N, F, S |
| APC     | Yes     | Yes       | No        | No        | No        | D, Mis, N, F, S, Met |
| RET     | Yes     | Yes       | No        | No        | No        | T, Mis, N, F |
| DNMT3A  | Yes     | No        | Yes       | No        | No        | Mis, F, N, S |
| NOTCH1  | Yes     | No        | No        | Yes       | No        | T, Mis, O |
| BRAF    | No      | No        | No        | Yes       | No        | Mis, T, O |
| KRAS    | No      | No        | No        | No        | Yes       | Mis |
| MYC     | No      | No        | No        | No        | No        | A, T |
| MDM2    | No      | No        | No        | No        | No        | A |
| AKT1    | No      | No        | No        | Yes       | No        | Mis |
| MYCN    | No      | No        | No        | Yes       | No        | A |
| CTNNB1  | No      | No        | No        | Yes       | No        | Mis, T |
| EGFR    | No      | No        | No        | Yes       | No        | A, O, Mis |
| MALAT1  | No      | No        | No        | Yes       | No        | T, Amp |
| NFE2L2  | No      | No        | No        | Yes       | No        | T, F |
| NPM1    | No      | No        | No        | Yes       | No        | Mis |
| NRAS    | No      | No        | No        | Yes       | No        | Mis |
| PIK3CA  | No      | No        | No        | Yes       | No        | Mis |
| PIK3R1  | No      | No        | No        | Yes       | No        | Mis, F, O |
| SETBP1  | No      | No        | No        | Yes       | No        | Mis |
| IDH1    | No      | No        | No        | Yes       | No        | Mis |
| IDH2    | No      | No        | No        | Yes       | No        | Mis |
| FLT3    | No      | No        | No        | Yes       | No        | Mis, O |

Abbreviations: D, gene deletion; F, frameshift mutation (small insertion and deletion); Met, loss of expression via promoter hypermethylation; Mis, missense mutation; N, nonsense mutation; O, other (splice mutations, intronic mutation); S, synonymous mutation; T, gene translocation

Mutation type for each cancer gene was extracted from the cancer gene census at the COSMIC database except for the methylation status (http://cancer.sanger.ac.uk/cancergenome/projects/cell_lines/). Fusion proteins generated via chromosomal translocation are not included in this list.
Conflict of Interest

KGW is a co-founder, shareholder and board member of the company Aprea AB that develops p53-based cancer therapy, including APR-246. TS declares no conflict of interest.

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pathways via nonspecific mechanisms. The view that mutated oncogenes such as KRAS or PIK3CA are only hyperactive proteins must be modified, as several studies have described additional activities suggestive of neomorphic (GOF) activity, but as these observations were based on cell transfection with protein overexpression more studies are needed to assess the exact behavior of all of these genes.97,98 The boundaries between hypermorphic and neomorphic mutations are not clear cut and alterations in a single gene can lead to variants that can have either one or both characteristics.

Although chromosomal translocations leading to fusion genes and the synthesis of chimeric proteins with novel functional specificity are the most obvious and easy-to-demonstrate neomorphic mutations, it is more difficult to demonstrate missense mutations acting as neomorphic mutations. Nevertheless, neomorphic mutations have recently been demonstrated in isocitrate dehydrogenases 1 and 2 (IDH1 and IDH2), which are mutated in several cancer types such as glioma and AML (Table 1). In both genes, a single missense mutation affecting an amino acid residue localized in the catalytic region of the protein accounted for > 90% of reported events and were shown to change the substrate specificity of these enzymes.99 As discussed in the previous section, several hotspot p53 mutants are obviously neomorphic with a marked diversity in the activity gained by each variant.

Conclusion

In vitro, mouse in vivo and clinical studies all point toward the importance of selection for oncogenic p53 mutants in human tumors. Although classification of cancer genes into onco- genes or tumor suppressor genes can be very helpful, it is becoming increasingly obvious that the boundary between the two classes is not as clear as previously thought, and that some genes may straddle these two categories. Furthermore, tissue type, genetic background and many other factors also influence the phenotype induced by a specific mutation. Nonetheless, we can predict that, except for rare cases, most cancer gene alterations will give rise to products with pleiotropic activities. Using the classification based on Muller’s proposition more than 80 years ago, it is possible to establish a comprehensive and meaningful categorization of the various cancer genes that illustrates how TP53 does not fit the classical definition of a tumor suppressor gene. Beyond the rhetorical aspect of this statement, there is a genuine need to avoid an overly simplistic, binary, ‘wild-type versus inactivated’ classification; defining p53 mutants as oncogenes with heterogeneous GOFs that affect multiple pathways must be considered. Designing drugs to target specific p53 mutants seems an attractive strategy, as several of these mutants are among the most frequent protein variants found in several cancer types and associated with the death of > 120,000 patients worldwide. Indeed, small molecules capable of targeting hotspot mutants p.R175H or p.Y220C have been identified. Further drug discovery efforts should focus on other frequent p53 mutants, such as p.R273H, p.R248W and p.R249S. With drugs targeting a wide range of common p53 mutants in our future arsenal, our chances of efficiently fighting cancer may be greatly improved.100

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