Cell cycle checkpoint control: The cyclin G1/Mdm2/p53 axis emerges as a strategic target for broad-spectrum cancer gene therapy - A review of molecular mechanisms for oncologists

ERLINDA M. GORDON1-3, JOSHUA R. RAVICZ1, SEIYA LIU4, SANT P. CHAWLA1 and FREDERICK L. HALL2,3

1Cancer Center of Southern California/Sarcoma Oncology Center, Santa Monica, CA 90403; 2Aveni Foundation and 3DELTA Next-Gen, LLC, Santa Monica, CA 90405; 4Department of Cell Biology, Harvard University, Cambridge, MA 02138, USA

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Correspondence to: Dr Erlinda M. Gordon, Cancer Center of Southern California/Sarcoma Oncology Center, 2811 Wilshire Boulevard, Suite 414, Santa Monica, CA 90403, USA
E-mail: erlinda.gordon@gmail.com

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Abstract. Basic research in genetics, biochemistry and cell biology has identified the executive enzymes and protein kinase activities that regulate the cell division cycle of all eukaryotic organisms, thereby elucidating the importance of site-specific protein phosphorylation events that govern cell cycle progression. Research in cancer genomics and virology has provided meaningful links to mammalian checkpoint control elements with the characterization of growth-promoting proto-oncogenes encoding c-Myc, Mdm2, cyclins A, D1 and G1, and opposing tumor suppressor proteins, such as p53, pRb, p16INK4A and p21WAF1, which are commonly dysregulated in cancer. While progress has been made in identifying numerous enzymes and molecular interactions associated with cell cycle checkpoint control, the marked complexity, particularly the functional redundancy, of these cell cycle control enzymes in mammalian systems, presents a major challenge in discerning an optimal locus for therapeutic intervention in the clinical management of cancer. Recent advances in genetic engineering, functional genomics and clinical oncology converged in identifying cyclin G1 (CCNG1 gene) as a pivotal component of a commanding cyclin G1/Mdm2/p53 axis and a strategic locus for re-establishing cell cycle control by means of therapeutic gene transfer. The purpose of the present study is to provide a focused review of cycle checkpoint control as a practicum for clinical oncologists with an interest in applied molecular medicine. The aim is to present a unifying model that: i) clarifies the function of cyclin G1 in establishing proliferative competence, overriding p53 checkpoints and advancing cell cycle progression; ii) is supported by studies of inhibitory microRNAs linking CCNG1 expression to the mechanisms of carcinogenesis and viral subversion; and iii) provides a mechanistic basis for understanding the broad-spectrum anticancer activity and single-agent efficacy observed with dominant-negative cyclin G1, whose cytotoxic mechanism of action triggers programmed cell death. Clinically, the utility of companion diagnostics for cyclin G1 pathways is anticipated in the staging, prognosis and treatment of cancers, including the potential for rational combinatorial therapies.

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Correspondence to: Dr Erlinda M. Gordon, Cancer Center of Southern California/Sarcoma Oncology Center, 2811 Wilshire Boulevard, Suite 414, Santa Monica, CA 90403, USA
E-mail: erlinda.gordon@gmail.com

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1. Introduction

Decades of research in the fields of biochemistry, molecular biology, molecular genetics, virology, genetic engineering, functional genomics and cancer gene therapy, have converged in identifying the executive components of a commanding regulatory axis of mammalian cell cycle control: Providing new mechanistic insights, biochemical pathways, unifying concepts and checkpoint control elements with profound implications in the prevention, diagnosis and treatment of cancer (Fig. 1). The challenge remains, however, to successfully integrate these celebrated epochs of gene discovery and biochemical pathway characterization into a practical understanding of cell cycle control befitting the actual praxis and applied pharmacology of contemporary clinical oncologists. This focused review, prepared by contributing scientists and clinical practitioners in the field of genetic medicine, is intended to present the current state-of-the-art in applied cell cycle checkpoint control as it relates to cancer management.

2. Tumor initiation, tumor promotion and Ser/Thr protein phosphorylation: Then and now

From a clinical perspective, the molecular mechanisms of chemical co-carcinogenesis first came to light in the 1940s with the pioneering studies of croton oil (i.e., phorbol esters) and 3,4-benzpyrene in a now-classic mouse skin model (1), in which increased tumor production was only observed when the inflammatory phorbol ester followed, rather than preceded, the application of the DNA-damaging carcinogen, thereby defining the separable and discernible stages of tumor initiation, promotion and progression (2). The subsequent discovery that protein kinase C (PKC), which plays a major role in signal transduction and cell proliferation, is the cellular receptor for the tumor-promoting phorbol esters (3) ushered in a wave of pharmaceutical interest in selective PKC inhibitors, only to be thwarted by the general multifunctionality of PKC, the multiplicity of PKC isoenzymes, the limited specificity of PKC modulators and the remaining unanswered questions and intricacies of PKC function, which stifled the promise of targeting PKCs for cancer therapeutics (4). Nevertheless, the importance of discrete Ser/Thr kinase activities, which operate via recognition-site-specific protein phosphorylation events, in the control of biochemical pathways governing mammalian cell growth and proliferation, as well as tumorigenesis, invasion and metastasis, has emerged as a major regulatory theme.

3. Focus on cyclin-dependent targeting of proline-directed protein phosphorylation

Basic research in yeast genetics characterized a number of cell division cycle (Cdc) mutants, thereby identifying important genes, notably Cdc28 in baker’s yeast (S. cerevisiae) and its homologue Cdc2 in fission yeast (S. pombe), which encode a unique serine/threonine protein kinase that is not only critical for mitosis in yeast, but is highly conserved in both structure and function throughout the evolution of all eukaryotes, including Homo sapiens (5). The molecular cloning and characterization of the Cdc2/Cdc28 kinase (CDK1 in mammals) and its implicit role in governing the defined stages and checkpoints of the eukaryotic cell division cycle supported by the independent discovery of cyclins A and B as prominent oscillating proteins of unknown function in sea urchin embryos (A. punctata), which were eventually cloned, biochemically characterized and determined to be positive-acting regulatory subunits of the executive Cdc2/Cdc28 kinase, thereby linking nascent protein synthesis (i.e., cyclin proteins) to the ordered stepwise progression of the cell division cycle through its defined phases (6). From this point onwards, the term ‘cyclin-dependent’ kinase (CDK) has been used to characterize the vertebrate homologs of this key regulatory enzyme. Recognized as a triumph of basic research in simple model systems, as well as a major advance in terms of elaborating the executive enzymology regulating cell cycle transitions, the principal investigators, Hartwell, Nurse and Hunt, were awarded the Nobel Prize for Medicine in 2001 (7). Armed with the DNA coding sequences for both the catalytic subunit (kinase) and the putative regulatory subunits (cyclins) of this executive protein kinase from primitive eukaryotic model systems, the molecular cloning and characterization of homologous coding sequences in higher animals began in earnest (8,9), enabling scientists to solve a lingering paradox in modern cell biology, i.e., the molecular basis for deconstructing cell cycle regulation in higher animals into two separable and distinct protein kinase activities, each with very different substrate specificities: The purified mitosis-promoting factor (MPF), which controls the G2-to-M phase transition (10) vis-à-vis the S phase-promoting factor (SPF), which orchestrates the G1-to-S phase transition (11,12).

Mechanistic insights were derived from international studies of site-specific protein phosphorylation in rat pheochromocytoma, which led to the identification and characterization of the growth factor-sensitive proline-directed protein kinase (PDPK), based on its ability to phosphorylate a unique site on tyrosine hydroxylase, a key regulatory enzyme in the synthesis of adrenal catecholamines (13). These collaborative studies led to the discovery that the preferred amino acid sequence recognized by this unique interphase kinase activity, in terms of substrate specificity, is X-Ser/Thr-Pro-X-X, thereby predating the requirement for a single orienting proline residue immediately adjacent to the actual phosphorylation site (14), affirming the appellation ‘proline-directed’ protein kinase, and presenting this unique Ser/Thr kinase activity (which is biochemically distinguishable from the M phase-specific histone H1 kinase activity of purified MPF) as a likely candidate for the illusive SPF in mammals (15). Structurally, the high frequency of S-P-X-X and T-P-X-X motifs (presumably type-I β-turns) found in gene regulatory proteins is not only indicative of a new DNA-binding unit (16), but also of proline-directed protein phosphorylation sites that have been canonical by natural selection, along with this site-specific protein kinase activity.

Molecular cloning and characterization of homologous kinase subunits in vertebrates revealed that the biochemical and physiological activities of MPF are attributable to CDK1/cyclin B complexes, the Ser/Thr kinase activity of which regulates the cytoskeletal dynamics associated with mitosis (10,12). In 1991, Hall et al characterized the subunits of the purified PDPK as a complex of CDK1 and cyclin A (17); when CDK2, a second homologue of the yeast Cdc2/Cdc28 kinase, was identified in humans, this homologous kinase, which is expressed somewhat earlier in the cell cycle compared...
with CDK1, was also found to partner with cyclin A and is enzymatically active as a CDK2/cyclin A heterodimer (18). Moreover, in addressing the paradox of differential substrate specificities, it was determined that the cyclin A subunit of these CDK complexes not only acts as a positive regulatory subunit, in terms of kinase activation, but it is the inducible ‘cyclin’ subunit that determines the substrate specificity of the active protein kinase. In this case, the cyclin A subunit physically targets the cyclin A/CDK holoenzymes to the Retinoblastoma (Rb) tumor suppressor protein (19), where progressive site-specific phosphorylation of pRb serves to inactivate the tumor suppressor (i.e., transcription/E2F repressor) (20), thereby linking the molecular activation of G1-phase transcription in humans to the expression of specific cyclin proteins (21). The cyclin-targeted CDK activities serve to overcome the suppressive function of Rb-related ‘pocket’ proteins (pRb, p107 and p130) that govern the feed-forward mechanics of the cell cycle, i.e., the coupling of protein phosphorylation and gene transcription, which drives cell cycle progression (22,23).

4. Focus on G1-phase regulation: Oncogenic cyclins vis-à-vis tumor suppressive gatekeepers

A fundamental characteristic of cancer genetics is the molecular dysregulation of cell cycle checkpoint control elements, which normally ensures the orderly progression of cell growth, DNA synthesis and mitotic cell division, while actively ensuring genomic fidelity. Among the manifold genetic alterations known to contribute to the pathogenesis of cancer in humans, including the molecular genetic disruptions of tumor viruses, the majority of these mutations are observed in genes that regulate progression through the G1 phase of the cell division cycle, including pRb-related tumor-suppressor proteins, which govern cell cycle progression, and the much-studied p53 tumor suppressor (mutated in >50% of human cancers), which serves as a molecular ‘guardian’ of DNA fidelity and an ‘executioner’ via its pro-apoptotic function (24). Alterations in the enzymatic machinery that controls the decisions to progress from a resting state (G0) into the cell cycle (G0-to-G1 transition) and/or to progress from the G1 to the S phase led to the identification of a growing family of human cyclins and their CDK partners that are directly implicated in the mechanisms of tumorigenesis and cancer (25-27). Thus, it is becoming abundantly clear that the major tumor suppressive elements (i.e., pRb and p53) that control progression through the mammalian cell cycle (28) are themselves a target for molecular inactivation by the expression and growth-promoting activities of two distinct types of potentially oncogenic cyclin proteins: The G1 cyclins (D-, E- and A-type cyclins) targeting CDKs to the Rb-Axis, and the perplexing cyclin G1, which disables the functions of p53 (Fig. 2).

The regulatory importance of human G1 cyclin expression in the promotion of cell proliferation and cancer is indicated by the telltale manner of their discovery and cloning in humans, which is directly linked to the pathogenesis of neoplastic disease. Human cyclin A was first identified biochemically as a co-precipitate, along with the pRb tumor suppressor, in a screen for proteins that bind with high affinity to the cell-transforming adenoviral E1A oncoprotein (29); soon thereafter, the molecular cloning of the human cyclin A gene was achieved by tracking the various insertion sites where the hepatitis B virus (HBV) physically integrates into the human genome (30). Importantly, it was determined that the HBV-induced dysregulation of the human cyclin A locus not only enforces a generalized overexpression of the gene, but it also physically mutated/truncated the N-terminal segment of the cyclin A protein, removing signal sequences necessary for its cyclical proteolytic degradation and producing a hybrid HBV-cyclin A transcript encoding a ‘stabilized’ cyclin A protein (hence interphase PDPK activity), which may indeed participate in the development and progression of hepatocellular carcinoma (HCC) (31). In subverting the cyclin A locus, a key component of the mammalian cell cycle machinery, the transforming HBV had apparently created a ‘dominant-positive’ mutant gene product to essentially commande cell cycle
control by maintaining a critical growth-promoting function of the cyclin A subunit (i.e., kinase activation via the retained ‘cyclin box’), while deleting the natural proteolytic processing, which normally eliminates cyclin A entirely, upon completion of each cell division cycle. Such viral subversion and oncogenic activation of a growth-regulatory element identifies human cyclin A as a bona fide proto-oncogene (15, 32), and hence a useful biomarker for characterizing tumor cell proliferation in histochemical detail (33).

The identification of the D-type cyclins (D1, D2 and D3) as oncogenes forged new and important links between mitogenic signal transduction, D-type cyclin gene expression and tumorigenesis (34): The growth factor-sensitive cyclin D1 was initially identified as a PRAD1/bcl-1 proto-oncogene, which is subject to gene amplification and/or rearrangement, resulting in cyclin D1 overexpression in a wide spectrum of human cancers, including B-cell neoplasms, carcinomas of the head and neck, various sarcomas and human breast cancers (>50%); cyclin D2 was identified as the integration site of a murine leukemia virus, while the cyclin D2 and D3 genes are amplified and overexpressed in numerous human cancers, including leukemias, lymphomas, glioblastomas, colorectal, renal and pancreatic carcinomas (26, 35). Partnering promiscuously as heterodimers with a number of CDK subunits (CKD1, 2, 3, 4 and 6), D-type cyclins function in early G1 to target the CDK complexes to the pRb-related tumor suppressor proteins, where site-specific inhibitory phosphorylation of pRb, to a limited extent, primes pRb (36) and advances cell cycle progression to a critical control point in the late G1 phase, known as the restriction point, or R-point (37): This is where the induction of cyclin E-dependent kinase activity contributes to the hyper-phosphorylation and inactivation of Rb, thereby releasing the E2F transcription factors and driving cells irreversibly through the G1-to-S phase transition, beyond which additional extracellular signals in the form of mitogenic growth factors are no longer required (38, 39). Taken together as an important class of growth-promoting proto-oncogenes, the orderly and progressive expression of specific G1 cyclins (D-type cyclins first, followed by cyclin E and cyclin A) are now viewed as inducible rate-limiting activators of G1 phase progression, the dysregulation of which is potentially oncogenic.

Just as the executive CDKs of G1 progression are engaged by positive regulatory subunits with profound oncogenic potential (i.e., G1 cyclins), G1 progression is also negatively controlled by polypeptide CDK inhibitors (CDKIs) whose expression is linked with the classical tumor suppressor p53. The finding that p53, a sequence-specific DNA-binding transcription factor, selectively induces p21 \( ^{WAF1/CIP1} \) (a universal CDKI) as a mediator of p53-initiated cell cycle arrest, has attracted interest in CDKIs as a strategic locus for prospective therapeutic interventions (40). Indeed, two families of endogenous CDKIs were found to function operationally as tumor suppressors: i) The WAF1/CIP/KIP family of CDKIs (p21, p27 and p57) appear to inhibit the activity of all major CDKs, while the INK4 family of CDKIs (p16, p15, p18 and p19) more specifically inhibit the cyclin D-dependent kinases CDK4 and CDK6, which phosphorylate/inactivate the pRb suppressor in early G1 (35, 41). Of note, genetic alterations involving the p16\( ^{INK4a} \) locus (chromosome 9p21) have been identified as germline mutations in melanoma-prone patients, and as deletions or mutations in a large percentage of primary human tumors, including sarcomas, lymphomas, leukemias, squamous cell carcinomas and pancreatic adenocarcinomas. In addition, genetic engineering of homologous deletions in mice confirmed the overall importance of this locus in the suppression of tumorigenesis (41, 42).

5. Targeting CDKs and CDK inhibitors (CDKIs) for cancer therapy: Current status, issues

Elucidation of the ‘executor’ roles of G1 cyclins, CDKs and CDKIs, as they are critical regulatory components of the cell cycle, and the development of effective strategies for their manipulation, hold promise for the rational design of anti-cancer therapies.
cycle machinery that are frequently altered in human cancers, prompted renewed interest in the development of specific kinase inhibitors expected to block cell cycle progression and/or induce growth arrest, thereby providing a prospective abundance of targets for novel antineoplastics (43).

Unfortunately, but not unexpectedly (see PKC issues above), the first generations of pharmaceutical pan-CDKIs failed to meet such expectations, due to the general lack of target specificity (44). Even the p21WAF1/Cip1 protein, which is the classic pan-CDKI that mediates p53-dependent cell cycle arrest, allowing for DNA repair, is sufficiently complex, in terms of the regulation of its expression, binding activities and subcellular localization, to function operationally, either as a tumor suppressor under certain circumstances (cell cycle block) or as a tumor-promoting oncogene (by preventing apoptosis), depending on the cellular context (45).

Over the past 20 years, a substantial number of prospective pan-CDKIs have been developed as potential cancer therapeutics and tested in numerous clinical trials in a variety of different tumor types, only to fail upon final analysis to meet objective clinical endpoints with an acceptable profile of systemic toxicities (46, 47). The principal reasons provided to explain the general failure of non-selective pan-CDKIs in the clinical setting are as follows: i) Lack of clear understanding of the physiological mechanisms of action as to which CDKs are actually being inhibited in vivo; ii) lack of patient selection and stratification on the basis of pertinent companion biomarkers that may help identify a subset of good clinical responders; and iii) lack of a therapeutic window, meaning the inability to achieve therapeutic levels of the drugs due to their intrinsic inability to discriminate between cancerous and healthy tissues (47). Despite the disappointing early attempts to develop pharmaceutical CDKIs for cancer therapy, a more specific focus on the cellular activities of CDK4 and CDK6, which are activated by the oncogenic D-type cyclins, has recently achieved limited success (47, 48). Nonetheless, the major challenges of precise target specificity, rapidly acquired drug resistance, untoward toxicities in normal tissues, optimal patient selection (companion biomarkers) and efficient drug delivery remain.

6. Cyclin G1, a non-canonical cyclin, opposes p53, the fragile ‘guardian of the genome’

Cyclin G1 is a non-canonical cyclin that i) structurally looks like a cyclin (cyclin box), ii) behaves like the earliest of all G1 cyclins (induction: G0-to-G1 boundary) and iii) physically targets a key regulatory enzyme (Mdm2) to a critical checkpoint substrate (p53) in the executive regulation of the cell division cycle. While the observed growth-promoting function of human cyclin G1 is somewhat analogous to the progressive cyclin D, E, A/CDK/pRb/E2F axis controlling cell cycle progression (Fig. 2), it is well-known that cyclin G1 operates via a separable and distinct regulatory pathway, namely the commanding cyclin G1/Mdm2/p53 axis, where precision targeted Ser/Thr phosphatase activity (i.e., cyclin G1-dependent Ser/Thr phosphatase activity) is intimately linked, through biochemical activation of the transforming protein of the MDM2 oncogene, to the commanding checkpoint control function(s) of the p53 tumor suppressor protein (Fig. 3).

The human cyclin G gene, now cyclin G1 or CCNG1 (49), was initially cloned by Hall et al at Children’s Hospital Los Angeles (Los Angeles, CA, USA), where it was determined to be overexpressed in human osteosarcoma cells (50), providing the first of several important links to cancer. The observed
overexpression of the cyclin G1 gene in human osteosarcoma, which remained constitutive throughout the cell division cycle in synchronized MG-63 cells, was rather perplexing, since murine cyclin G was identified in mice in a molecular screen for transcriptional targets of the p53 tumor suppressor, thereby (mistakenly) suggesting a growth-inhibitory function rather than a growth-promoting function for cyclin G1 (51). However, it was experimentally determined that the enforced overexpression of cyclin G1 in either normal or neoplastic cell lines did not cause the anticipated cell cycle arrest, nor did experimental overexpression of cyclin G1 induce apoptosis (51), as the p53 tumor suppressor (referred to as the molecular guardian of the genome and policeman of the oncogenes) is known to do under various conditions (52). By contrast, the ectopic overexpression of cyclin G1 is reported to promote the clonal expansion of normal human fibroblasts and to accelerate cell growth in RKO (p53+/−) colon carcinoma cells (53), whereas the molecular suppression (or knockout) of cyclin G1 expression by antisense strategies was found to be uniformly lethal (54). These studies revealed an essential growth-promoting function for cyclin G1, thus identifying CCNG1 as an accessible gene locus that may, through its suppression, demonstrably inhibit the growth of human tumor xenografts (upon intra-tumoral injection) in a nude mouse model of cancer (55,56).

In 1995, Hall et al., in collaboration with E.M. Gordon, Medical Director of the Vector Production Unit, USC Gene Therapy Laboratories (Los Angeles, CA, USA) altered the focus of his research laboratory from the biochemistry of proline-directed protein kinases, CDK pathway characterization and related gene discovery to work together in translational research: Their goal was to develop a targeted and injectable gene delivery vehicle (vector) that would be efficient, effective and safe for repeated use in clinical practice. While their pioneering work on a tumor-targeted retrovector was progressing, Hall and Gordon utilized available retroviral vector-mediated gene transfer technologies to investigate functional knockouts of cyclin D1 (antisense CycD1) and cyclin G1 (antisense CycG1) in MG-63 osteosarcoma cells, in comparison with the enforced expression of p21WAF1/CIP1, a universal CDKI expressed in sense orientation (54). After examining the comparative experimental results in terms of both cytostatic (all three constructs inhibited cell proliferation) and cytotoxic activities (i.e., TUNEL staining for overt apoptosis), Hall and Gordon focused on cyclin G1 knockout for potential clinical development. At that time, this was a ‘road less traveled’ in terms of regulatory biology, which did indeed make a difference in the clinical setting.

7. The commanding cyclin G1/Mdm2/p53 axis operating in normal and neoplastic cells

A critical ‘executor’ element in the cyclin G1/Mdm2/p53 axis is the human homologue of the murine double minute gene 2 (Mdm2) oncogene. Originally characterized as an amplified gene locus encoding a potent transforming (i.e., tumorigenic) oncoprotein in mice, the human Mdm2 gene was found to be amplified in numerous human cancers, including soft tissue sarcomas (20%), osteosarcomas (16%) and esophageal carcinomas (13%) (57,58). More recently, links to the pathogenesis of breast cancer confirm that abnormally high levels of the Mdm2 oncoprotein are detected in at least one-third (38%) of human breast cancers, which cannot be explained by gene amplification alone (59). Shortly after its characterization, it was discovered that the Mdm2 oncoprotein forms a tight physical complex with the p53 tumor suppressor, thereby inhibiting p53-mediated transactivation events (60). In addition to inhibiting p53-dependent transcription, it was determined that the Mdm2 oncoprotein functions biochemically as a specific E3 ubiquitin ligase that is responsible for the ubiquitination and degradation of the p53 tumor suppressor protein (61). Thus, the Mdm2 oncoprotein antagonizes the p53 tumor suppressor on two levels: By ubiquitin-independent repression of p53 transactivation, and by ubiquitin-dependent proteolysis of the p53 protein (62), as seen in Fig. 3 (MOA II, tumor promotion).

This antagonistic association of the Mdm2 oncoprotein with p53 function and stability suggested that overexpression of the Mdm2 gene is yet another molecular mechanism, in addition to deletion or inactivating mutations in the TP53 gene per se, that may functionally inactivate the p53 tumor suppressor in the acquisitive process of neoplastic transformation. Therefore, it was reasoned that pharmaceutical agents targeting the Mdm2/p53 interaction in transformed cells retaining wild-type TP53 status may be a promising new approach to cancer therapy (63), which is considered particularly important in the context of hematological malignancies, where the frequency of TP53 mutations is relatively low (in comparison with solid tumors), while Mdm2 is frequently amplified and overexpressed (64). Pharmacological targeting of the pivotal Mdm2/p53 interaction with natural compounds, small molecules, and even stapled (helix-linked) peptides, has been extensively investigated in an effort to restore the tumor suppressor activity of wild-type p53 (65). Years of preclinical experimentation have identified promising drug candidates for further clinical evaluation; however, issues of drug-related toxicity and acquired resistance persist, and it was concluded that this particular class of compounds would definitely benefit from rational combinations with a second, potentially synergistic, pharmacological strategy (64).

In the elaboration of a pivotal Mdm2/p53 axis controlling both DNA-fidelity and cell fate, the oncogenic function of the non-canonical cyclin G1 was revealed: From a biochemical perspective, the critical functions of the Mdm2 oncoprotein, which is the major-negative regulator of p53, are intricately controlled by multiple site-specific protein phosphorylation events (66), thereby necessitating a site-specific Ser/Thr dephosphorylation event in order for the Mdm2 oncoprotein to be conformationally activated to carry out the destabilization of the p53. Moreover, this crucial Mdm2-activating dephosphorylation event was found to be cyclin G1-dependent. It was determined that the cyclin G1 protein forms a stable complex with the Ser/Thr protein phosphatase designated 2A (PP2A) (67,68), specifically with the B'-class of regulatory subunits that determine both the substrate specificity and the subcellular localization of PP2A complexes (69). Conceptually, cyclin G1 binds tightly to the Mdm2 oncoprotein in vitro and in vivo, where it recruits PP2A to the Mdm2 oncoprotein, thereby stimulating the ability of the PP2A to dephosphorylate and activate Mdm2 at the critical regulatory site (68). In this capacity, cyclin G1 acts both as a targeting subunit and as a selectivity factor that stimulates and propels PP2A catalytic
activity toward Mdm2. The discovery that cyclin G1 associates physically with both PP2A and Mdm2 and, furthermore, that this physical association regulates the accumulation and degradation of the p53 protein (70), provides new and important insights into the oncogenic function of cyclin G1 and suggests that the main role of cyclin G1 may be to activate the Mdm2 oncoprotein to override the cell cycle checkpoint control functions of p53 (71). These findings provide a mechanistic explanation for the observation that cyclin G1 expression is associated with growth promotion rather than with growth arrest (54,72). This conclusion is further supported by studies of genetic engineering in mice, where it was discovered that cyclin G1-deficient mice not only survive, but exhibit a reduced incidence of hepatic tumors upon exposure to hepatocarcinogens followed by partial hepatectomy (73). This decrease in tumor susceptibility associated with loss of cyclin G1 function is attributed to a resulting increase in p53 levels and a corresponding increase in p53 tumor suppressor activity (73). Taken together, these findings raise the possibility that the strategic modulation of cyclin G1 function in the commanding cyclin G1/Mdm2/p53 axis may be targeted at the molecular level for the development of novel anticancer agents.

8. Targeting cyclin G1 function in experimental hyperplasia: Applications to cancer control

Upon examining the role of cyclin G1 in normal proliferative cell populations, which exhibit normal levels of wild-type TP53 and Mdm2 gene expression, it was determined that downregulation of cyclin G1 (CCNG1) by retrovirus-mediated antisense gene transfer inhibits vascular smooth muscle cell proliferation and subsequent neointima formation (a form of hyperplasia) seen in rat carotid arteries following balloon catheter injury (74). Similar cytostatic and cytocidal effects of antisense cyclin G1 treatment were observed in proliferative cell populations of both rabbit keratocytes and human fetal lens epithelial cells (75). Immunohistochemical staining for cyclin G1 expression in balloon-injured arteries revealed a marked induction of gene expression, from non-detectable levels in quiescent arteries to relatively high levels of immunoreactivity in the proliferative neointimal masses (Fig. 4A). The inducible expression of cyclin G1 observed in vascular injury/restenosis is reminiscent of the marked induction of cyclin G1 in the regenerating liver following partial hepatectomy (76), rising rapidly from exceedingly low basal levels measured in the quiescent liver to appreciable steady-state levels. In contrast to the normally low levels seen in the liver, the expression of cyclin G1 in human tumor xenografts appears to be constitutively elevated (Fig. 4A).

Targeting cyclin G1 function with a dominant-negative (dnG1) mutant construct. After realizing that antisense ‘knockdown’ of cyclin G1 expression in normal and neoplastic cells was indeed informative and promising, yet reasoning that partial or incomplete suppression of CCNG1 gene expression may still be problematic under physiological conditions (where enzymatic phosphotransferase rates are measured in msec and compensatory changes in gene expression are likely to occur), it was considered preferable to develop and deploy a ‘dominant-negative mutant’ construct of the cyclin G1 protein, the enforced expression of which would serve to counteract vital aspects of the cyclin G1 structure, binding and/or metabolism, thereby acting as a cyclin G1 antagonist, even in the presence of an abundance of the wild-type protein (Fig. 4B).
Working with structural deletions and point mutations in the human cyclin G1 coding sequences, based on the homology of the repeating helical domains comprising what is referred to as the cyclin box, and the conserved amino acid residues predicted from crystallography to serve as contact points of the cyclin A/CDK2 complex (49,50), an N-terminal deletion mutant was selected for development as the most potent negative-acting mutant of cyclin G1 (77). This dominant-negative mutant construct of human cyclin G1 (designated dnG1) is devoid of the extended N-terminal domain proximal to the cyclin box domain, and is missing the first two helical segments (α1 and α2 helices) of the cyclin box itself (Fig. 9).

9. Development of a targeted retroviral vector to efficiently deliver the dnG1 ‘designer gene’

The turn of the 21st century brought major biotechnological advances in the design and clinical applications of actively-targeted gene delivery vehicles, specifically patho-tropic (disease-seeking) gene therapy vectors, which enabled the first systemically administered, tumor-targeted retroviral vectors to be validated in clinical medicine (78). While the gristly pathobiology of the tumor extracellular matrix (ECM), as well as the molecular mechanisms of retroviral vector-mediated gene transfer and expression, are beyond the scope of this focused review, the potential clinical implications of an ‘active’ tumor-targeted gene delivery vehicle are wide-ranging. Focusing on the abnormal collagenous proteins (i.e., lesion signature proteins) that are pathologically exposed during the process of tissue injury (such as balloon angioplasty), Hall and Gordon adaptively engineered a physiological surveillance function embodied within the complex structure of von Willebrand coagulation Factor (vWF), which normally guides platelets to sites of significant tissue injury, into the surface envelope protein (gp70) of the Moloney murine leukemia virus, thereby creating a desirable ‘gain-of-function’ (pathological-matrix-targeting), without impairing the natural receptor-mediated binding and entry of the targeted viral particles into target cells, thus preserving the efficiency of gene transfer (79,80). Eventually, in anticipation of human gene therapy applications, the enabling tumor-targeting gain-of-function, first established in rodents, was genetically engineered into the 4070A ‘amphotropic’ murine leukemia virus envelope protein that is capable of transducing human cells (81,82).

Armed with two powerful enabling biotechnologies, i) the matrix/lesion-targeted retrovector and ii) the cytoidal dominant-negative mutant construct of cyclin G1 (dnG1), validation of both targeted gene delivery and single-agent efficacy was demonstrated in preclinical studies of vascular restenosis, where significant long-term inhibition of neointima formation in balloon-injured rat arteries was observed (77,83). This initial proof-of-concept was followed by preclinical studies on the safety and efficacy of the matrix/lesion-targeted vector bearing the inhibitory dnG1 construct (Mx-dnG1) applied as simple eye drops for the prevention of corneal haze induced by excimer laser-injury in rabbits (84). The encouraging results of this preclinical study prompted a formal evaluation of potential applications of anti-proliferative eye drops in humans by the NIH Recombinant DNA Advisory Committee (RAC), for a phase I/II intervention for superficial opacity/corneal scarring that occurs after phototherapeutic keratectomy (85). With these critical proofs-of-principle in hand (i.e., the efficacious preclinical management of experimental hyperplasia with Mx-dnG1), Hall and Gordon shifted their pioneering studies of lesion-targeted gene delivery to include the always pertinent and challenging models of metastatic cancer (86,87) (Figs. 5-7), where the physiological obstacles to tumor-targeted gene delivery (diffusion, filtration, turbulence, shear forces, inactivation) remain paramount, and the clinical management of cancer cell growth, as well as the clinical management of tumor-angiogenesis, represents a major unmet medical need.

Important insights into the mechanisms and efficiencies of tumor-targeted gene delivery were gained from a liver model of metastatic cancer, wherein human MiaPaca2 pancreatic cancer cells were infused into the portal vein to establish tumors in nude mice (Fig. 5). In this model of regional gene delivery to established liver metastases, marker studies demonstrated an informative and selective targeting of cancer cells within the liver during the earliest stages of the metastatic process (Fig. 5A and C; left panel). Of note, this exquisite targeting of collagenous matrix signature proteins (i.e., biochemical footprints) left by invasive cancer cells, long before a discernable tumor is formed within the liver, has considerable implications in the diagnosis and treatment of human cancers. Later on, when flagrant tumors are formed within the liver and robust tumor angiogenesis is apparent, the tumor targeting extended to the tumor vasculature, where the proliferative endothelial cells are transduced by the vector, as demonstrated by β-galactosidase transgene expression (Fig. 5D-F; left panel). First and foremost, when the marker gene was replaced by dnG1, as in the cancer gene therapy vector (Mx-dnG1), dose-dependent tumor eradication was observed upon repeated intravenous infusions (86) (Fig. 5; right panel).

Prospective clinical trials for this regional approach to cancer gene therapy underwent formal review (and approval) for use in humans by the Food and Drug Administration (FDA)/RAC, a tumor site-specific phase I/II evaluation of the safety and efficacy of hepatic arterial infusion of a matrix-targeted retroviral vector bearing a dnG1 construct as treatment for colorectal carcinoma metastatic to the liver (88). However, at this point, Hall and Gordon opted instead to explore the even greater potential for systemic gene delivery (87), in light of the potential for more wide-ranging clinical applications in the treatment of metastatic cancers (Figs. 6-7).

In a classic nude mouse model of metastatic cancer (Fig. 6; left panel), it was observed that tumor-targeted gene delivery is not only conceivable (A and B vs. C), it is apparent (D and E vs. F); in addition, it is evident that the physiological partitioning of the circulating vector to tumors is dependent on the high-affinity matrix (Mx)-binding domain that was genetically transposed from the D2 module of the vWF propeptide into the primary structure of the retroviral envelope protein (79-83,87). Notably, in order to accomplish tumor-targeted gene delivery within 1 h, the matrix/lesion-targeted vector infused into the tail vein of the mice necessarily transited the heart, the lungs, and the heart once more, prior to entering the systemic circulation, including the blood vessels feeding the tumor xenografts (Fig. 6A and B; left panel), where it appears to spread out, similar to Coomassie blue dye in a natural sponge, as it physically contacts and effectively
transduces proliferative tumor cells with a β-galactosidase marker gene (Fig. 6D and E; left panel).

When the marker gene was replaced with the inhibitory dnG1 construct (Fig. 6; right panel), it was confirmed that the cytocidal mechanism of action of dnG1, as observed in both hyperplastic lesions and cancerous xenografts, is attributed to the activation of DNA fragmentation via the process of apoptosis. This pioneering study demonstrated that the Mx/lesion-targeted dnG1 retroviral vector (Mx‑dnG1) deployed by peripheral vein injection i) accumulates rapidly in
tumor vasculature, ii) transduces tumor cells with exceedingly high-level efficiency and iii) enables demonstrable therapeutic gene delivery and long-term efficacy of dnG1 without eliciting appreciable toxicity (87). In addition to the marked apoptosis observed in Mx-dnG1 treated tumors (Fig. 6; right panel), characteristic histological changes in tumors after repeated systemic administration include: Zones of focal necrosis, overt anti-angiogenesis and reparative fibrosis accompanied by ECM deposition, which occupies an increasingly greater proportion of the residual tumor, as the cancer cells and the proliferative tumor vasculature are eradicated progressively by the repeated intravenous infusions of Mx-dnG1 (Fig. 7; left panel A-D control tumors vs. right panel A-D treated tumors).

Following the establishment of three preclinical proofs of principle, namely that i) tumor-targeted gene therapy is feasible via simple intravenous infusions, ii) tumor-targeted delivery of the dnG1 ‘killer gene’, which exhibits both anti-angiogenic and anti-tumor activities, is capable of altering the course of metastatic pancreatic cancer, an otherwise fatal disease and iii) the design features and general safety profile of the first targeted, injectable gene therapy vector had been critically evaluated by the FDA/RAC and formally approved for clinical use in humans, the first clinical studies were initiated.

10. Clinical studies demonstrate broad-spectrum anticancer activity and single-agent efficacy

In 2003, a clinical-grade retroviral expression vector bearing an inhibitory dnG1 construct of the cyclin G1 gene (designated Rexin-G) became the world’s first targeted, injectable vector to be approved for clinical trials in the treatment of intractable metastatic disease. Uniquely suited, by design, to perform this tumor-targeting function within the context of the human circulatory system, Mx-dnG1 (aka Rexin-G) is a pathotropic (disease-seeking) gene delivery vehicle bearing the dnG1 ‘killer’ gene, a therapeutic tumor-targeted nanoparticle that selectively seeks out and accumulates in metastatic lesions upon simple intravenous infusion. Preclinical validation of tumor-targeted gene delivery, accompanied by critical demonstrations of dose-dependent efficacy in pertinent cancer models, was followed by the first clinical studies, which aimed to demonstrate the potential for achieving broad-spectrum, single-agent, anticancer efficacy in the clinic. A summary of completed clinical studies is shown in Table I.

As with all targeted biologics, a major objective was to assess and establish the clinical safety of the vector system, while progressively seeking efficacy within the rule-based designs and dose escalation protocols used in phase I cancer clinical trials (89). Cooperative (US FDA sanctioned/BFAD-approved) international studies conducted in the Philippines enabled stepwise intra-patient dose escalations, thereby achieving clinical dose-dependent efficacy in an expedient manner (90,91), while the US FDA required more traditional studies of potential dose-limiting toxicities (DLT) and/or cumulative toxicities to be conducted at sub-therapeutic doses (92) before a formal determination of adequate safety would enable a progressive stepwise dose escalation. When clinical significant dose-dependent efficacy was demonstrated in US-based phase I/II studies of intravenous Rexin-G for advanced sarcoma and osteosarcoma (93-95), including
significant improvements in both progression-free and overall survival, the FDA approved an ‘across-the-board’ dose escalation allowance (applicable to all ongoing trials), thereby enabling these higher, more efficacious doses (i.e., optimal biological doses) to also be administered to breast and pancreatic cancer patients (96-98).

From these clinical studies, including the physiological dose-response kinetics, a predictive pharmacology of tumor-targeted genetic medicine began to emerge: A calculus of parity (or therapeutic equivalence) was introduced, in which the metastatic tumor burden of a given patient is evaluated and used in calculations, including vector targeting efficiencies and performance, in determining the effective intravenous dose(s) required to achieve tumor eradication (91). Advancements in bioprocess development enabled high-potency clinical grade vectors, which further simplified intravenous infusions and led to the accelerated approval of Rexin-G for all solid tumors by the Philippine FDA in 2007. This was followed by a rapid progression of clinical studies in the US, where Rexin-G received several Orphan Drug designations, including FDA Fast-Track status for metastatic pancreatic cancer, which stands at the cusp of phase III clinical trials (97,98). Long-term follow-up of an advanced phase I/II study using Rexin-G for chemotherapy-resistant sarcomas revealed the clinical value of the observed Rexin-G (dnG1) dose-response relationships in terms of overall patient survival: The 1-year survival rate increased from 0% at dose level I to 28.5% at dose level II-III, to an impressive 38.5% at dose level IV-V (with 31% of the patients alive at 2 years) following initiation of Rexin-G monotherapy, which may be considered as a gold standard in terms of objective clinical responses (96-98).

At the time of this review, Rexin-G had been administered safely to >270 patients worldwide, including >3,000 total intravenous infusions without serious side effects, vector issues, inactivating antibodies, or DLT. Documentation of significant anticancer activity, single-agent efficacy and objective clinical responses (including evidentiary histology), have been reported in a broad spectrum of cancers derived from all three germ layers (98-100). Clinical confirmation of i) tumor-targeted gene delivery, ii) a pro-apoptotic dnG1 mechanism of action and iii) characteristic changes in tumor histology (originally observed in preclinical models) are clearly recapitulated in the clinical setting, as is clearly seen in an opportunistic surgical (liver) biopsy obtained from a metastatic pancreatic cancer patient undergoing intravenous Rexin-G treatment (Fig. 8). In addition to statistically significant increases in patient survival, a considerable number of advanced-stage, chemotherapy-resistant cancer patients treated with repeated infusions of Rexin-G as monotherapy, including metastatic breast cancer, pancreatic cancer, malignant melanoma, osteosarcoma and soft tissue sarcoma patients, remain cancer-free or without active disease progression ≥9-12 years after the initiation of Rexin-G treatment (101).

11. Restoring long-lost tumor suppression with a dominant-negative cyclin-G1 antagonist

Strong support for the proposed growth-promoting function of cyclin G1 in the commanding cyclin G1/Mdm2/P53 axis came from functional genomics, which identified microRNAs (miRNAs) as a new class of endogenously expressed, small non-coding gene-regulatory RNA molecules that interact with the p53 transcription factor and its network of effectors at multiple levels (102,103), thereby impacting both the mechanics of cell cycle progression and the fate of proliferative cells. High-throughput screens investigating the role of miRNAs in the pathogenesis of human HCC identified miR-122 as the most predominant species of hepatic regulatory miRNA, which is
either lost or severely downregulated in ~70% of HCC cancers and in all HCC-derived cell lines (104). These studies identified cyclin G1 as a primary target of miR-122 suppression, which accounts for the inverse correlation between miR-122 and high levels of cyclin G1 expression observed in primary liver carcinomas. Loss of miR-122 expression in liver cancer correlates with a poor prognosis indicative of tumor progression, including a notable gain in invasive metastatic properties (105).

As depicted in Fig. 3, miR-122 regulates p53 protein through cyclin G1 (106): The miR-122/cyclin G1 interaction modulates p53 protein stability and transcriptional activity. Accordingly, enforced/restored expression of miR-122 decreases cell viability (107) and induces cell cycle arrest and apoptosis in HCC cell lines (108).

Important insights into the natural tumor suppressor function of miR-122 were gained from genetic engineering, where it was determined that miR-122 gene knockout mice are prone to hepatic tumor development, while AAV-mediated delivery of miR-122 to the livers of such mice inhibits tumorigenesis (109). The miR-122/cyclin G1 interaction modulates p53 protein stability and transcriptional activity. Accordingly, enforced/restored expression of miR-122 decreases cell viability (107) and induces cell cycle arrest and apoptosis in HCC cell lines (108).

The establishment of cyclin G1 (CCNG1) as a bona fide proto-oncogene and a powerful growth-promoting protein has profound implications for the clinical staging, prognosis and management of human cancers. Normally induced in response to tissue injury (G0/G1 boundary), as in hyperplasia (Fig. 4), cyclin G1 is frequently overexpressed in human breast and prostate cancers (111), osteosarcomas, colorectal cancers (112) and HCC (113), wherein CCNG1 overexpression is clearly correlated with a poor prognosis. As mentioned above, CCNG1-null mice survive and even exhibit decreased incidence, size and malignancy of tumors due to the resulting dominance of DNA-sensing p53-dependent tumor suppressor functions (73); however, in proliferative populations of tumor cells, including tumor neovasculature, the blockade of cyclin G1 expression or function is uniformly lethal. This apparent dependence of cancer cells on sustained cyclin G1 expression for cell survival, which is acquired during the course of diverse multistage carcinogenesis, is referred to as ‘oncogene addiction’ (114); indeed, this acquired state of dependency on cyclin G1 identifies CCNG1 as a vulnerable locus in human cancers, which can be targeted on several levels to gain a positive therapeutic advantage.

12. Cyclin G1 proto-oncogene promotes cell survival and progression over DNA fidelity

The establishment of cyclin G1 (CCNG1) as a bona fide proto-oncogene and a powerful growth-promoting protein has profound implications for the clinical staging, prognosis and management of human cancers. Normally induced in response to tissue injury (G0/G1 boundary), as in hyperplasia (Fig. 4), cyclin G1 is frequently overexpressed in human breast and prostate cancers (111), osteosarcomas, colorectal cancers (112) and HCC (113), wherein CCNG1 overexpression is clearly correlated with a poor prognosis. As mentioned above, CCNG1-null mice survive and even exhibit decreased incidence, size and malignancy of tumors due to the resulting dominance of DNA-sensing p53-dependent tumor suppressor functions (73); however, in proliferative populations of tumor cells, including tumor neovasculature, the blockade of cyclin G1 expression or function is uniformly lethal. This apparent dependence of cancer cells on sustained cyclin G1 expression for cell survival, which is acquired during the course of diverse multistage carcinogenesis, is referred to as ‘oncogene addiction’ (114); indeed, this acquired state of dependency on cyclin G1 identifies CCNG1 as a vulnerable locus in human cancers, which can be targeted on several levels to gain a positive therapeutic advantage.
Examination of the structure of cyclin G1 provides conceptual insights as to its multiplex biochemical functions (Fig. 9; MOA IV, right panel). The cyclin G1 protein is an unusually unstable protein, exhibiting a half-life of ~20 min; this instability is attributed to ubiquitin-mediated proteolysis involving the extreme N-terminus of the protein (115). The homology between cyclin G1 and known regulatory cyclins suggests that cyclin G1 may activate one or more CDK partners under certain conditions. Indeed, cyclin G1 is capable of physically interacting with CDK1, 2, 4, 5 and 6, the kinase activity of which appears to contribute physiologically to the short half-life of the cyclin G1 protein (116). However, neither Histone H1- nor Rb-kinase activity was detected in cyclin G1/CDK2 immunoprecipitates (116), suggesting that cyclin G1 association with a specific CDK partner, as well as the resulting CDK activity, may be more tightly regulated by additional factors and/or is uniquely targeted to one or more undetermined protein substrates. Subsequently, convincing aspects of cyclin G1 function in direct association with CDK5 have been revealed: Activation of CDK5 by cyclin G1 binding physically targets the active kinase complex to the c-Myc oncoprotein, the overexpression of cyclin G1 enables cancer cells to overcome radiation-induced (i.e., DNA-damage-induced) cell cycle arrest (117). Although the transcriptional targets of c-Myc include a number of DNA repair genes, thereby coupling DNA replication to the pathways and processes that preserve the integrity of the genome (118), the net effect of cyclin G1 function in association with CDK5 (or CDK2) is to abrogate DNA-fidelity checkpoint controls to promote cell survival, cell competence and cell cycle progression at the peril of increasing error-prone DNA synthesis, as is often found in cancers.

Significant clues into the function of cyclin G1/CDK complexes as a ‘survival factor’ were obtained in the nervous system, where cyclin G1 expression was induced by nerve injury in mature (post-mitotic) motor neurons and remained elevated during the early stages of nerve regeneration along with other ‘immediate-early’ genes associated with cell survival (119), suggesting that cyclin G1 functions as a survival factor in post-mitotic neurons, maintaining cell viability during the process of tissue repair. Following this mechanistic insight, we find that mild cognitive impairment, a transitional disease state that precedes more serious neurodegenerative problems associated with Alzheimer’s disease (AD), is accompanied by increased co-expression of cyclin G1, CDK2 and CDK5 in afflicted brain regions (120). This is in keeping with the theory that neuronal cell death associated with AD has, as its root cause, an ectopic re-entrance into the cell cycle (121), which results in the hyperphosphorylation of microtubule-associated tau proteins characteristic of AD models. The resulting cyclin G1/CDK5 phosphorylation of c-Myc on Ser-62 (X-Ser*-Pro-X) stabilizes the multifunctional transcription factor, resulting in transcriptional activation of a defined (E-Box-containing) set of growth-promoting genes, including both cyclins and CDKs, which are capable of driving cell proliferation (118). Mechanistically, c-Myc is a critical platelet-derived growth factor-inducible ‘competence gene’ that activates diverse cellular processes associated with entry and progression through the cell cycle, including the synthesis of cellular components in preparation for cell division. In this manner, by activating and selectively targeting CDK5 kinase activity to activate and stabilize the c-Myc oncoprotein, the overexpression of cyclin G1 enables cancer cells to overcome radiation-induced (i.e., DNA-damage-induced) cell cycle arrest (117). Although the transcriptional targets of c-Myc include a number of DNA repair genes, thereby coupling DNA replication to the pathways and processes that preserve the integrity of the genome (118), the net effect of cyclin G1 function in association with CDK5 (or CDK2) is to abrogate DNA-fidelity checkpoint controls to promote cell survival, cell competence and cell cycle progression at the peril of increasing error-prone DNA synthesis, as is often found in cancers.
of AD neurofibrillary tangles. Various neurotoxic insults induce hyperactivation of CDK5, with fatal consequences for the cell (122). On the other hand, lack of CDK5 activity results in neuronal cell death (123), suggesting that precise control of CDK5 activity is crucial for neuron survival via the site-specific phosphorylation of the apoptosis inhibitor Bcl-2 (124). Apart from cyclin G1/CDK5 complexes, which target and activate the c-Myc competence gene in cancer cells (117), a brain-specific regulatory subunit of CDK5 has been identified as p35, the N-terminal truncation of which (from p35 to p25) causes prolonged activation of CDK5, mis-localization and pathological phosphorylation of substrates; i.e., the p25/Cdk5 kinase hyperphosphorylates tau, disrupts the cytoskeleton and promotes cell death by apoptosis in primary neurons (122,125). In this same manner in which the proteolytic truncation (N-terminal deletion) of this p35 CDK5-activator protein converts the p35/CDK5 kinase complex from a survival factor to a cytotoxic p25/CDK5 kinase activity that promotes cell death by apoptosis in neurons, the ‘designer’ N-terminal truncation of cyclin G1 into a dnG1 mutant protein (Fig. 9) converts the survival functions (i.e., anti-apoptotic, pro-survival) of the CCNG1 oncogene (mediated via cyclin G1/CDK5-dependent c-Myc activation) into the pro-apoptotic, cytocidal functions of the truncated mutant dnG1 ‘killer’ gene. In this case, a potent proteolytically stable antagonist of wild-type CCNG1 gene function is generated.

A note to future biomedical researchers. While the functionality of cyclin G1 in the nervous system is beyond the scope of this oncological review (cancer cells are constitutively competent), this mechanistic link to cell survival and proliferative competence has important implications in terms of artificially/therapeutically promoting nerve regenerations and associated tissue repair, in light of the striking potentiality for tail and/or limb regeneration that is evident in lower vertebrates.

Whereas the early induction of cyclin G1 observed as cells exit from quiescence (G0-to-G1 transition) in experimental models of hyperplasia and liver regeneration is indicative of its role in establishing cellular competence to proliferate, the putative CDK kinase partner would thereby represent a G0/G1 competence-promoting factor (CPF), perfectly analogous to SPF and MPF. The target substrate for this cyclin G1-dependent kinase activity, CDK5 and/or possibly CDK2, (116,120) is the c-Myc oncogene, the activation of which is associated with the start of cellular proliferation, cell cycle progression, increased genomic instability, reduced adhesion, EMT, tumor invasion and metastasis. The overexpression of c-Myc in ~50% of human cancers correlates with poor patient survival (126), while c-Myc is often considered to be among the most desirable cancer targets, although it appears to be among the most ‘undruggable’ of molecular targets in all of cancer therapy (127).

In addition to i) promoting cell growth and survival as an enzymatic cyclin G1/CDK5 (or cyclin G1/CDK2) CPF in reparative tissues, and ii) disabling p53 checkpoint control via the cyclin-G1/Mdm2/p53 axis, resulting in error-prone DNA-synthesis that is characteristic of an advanced metastatic state, a cyclin-G1-dependent pathway also appears to mediate the troublesome EMT, which characteristically results in a more aggressive, invasive cancer phenotype (observed in distant metastases) and, hence, a worsening prognosis (113). Recently, CCNG1 was found to be amplified in a particularly aggressive subset of triple-negative breast cancer (TNBC) cells and in patients with chemotherapy-resistant TNBC, in which cyclin G1 overexpression was directly linked to the molecular mechanisms regulating both polyplody and chemotherapy resistance (128): Upon paclitaxel exposure, the pro-survival function of cyclin G1 promotes breast cancer cell survival by inducing polyplody, thereby limiting treatment options and outcomes involving taxanes, yet further validating cyclin G1 as a strategic target for medical intervention in the treatment of advanced metastatic breast cancer (Table I).

13. Combinatorial approaches and companion diagnostics for CCNG1 targeted therapies

Dysregulation of cyclin G1 is the driving oncogenetic event in a large number of human cancers. By restoring a critical locus of CCNG1 suppression, the enforced expression of miR-122, as well as cyclin G1 silencing, suppresses tumor growth and increases the sensitivity of HCC cells to doxorubicin (DOX) (106); thus, exciting new possibilities for combinatorial therapies are emerging. Indeed, cyclin G1 expression regulates and determines the outcome of taxane chemotherapy: Elevated cyclin G1 expression accompanies paclitaxel (PTX) -induced mitotic arrest and promotes cancer cell survival after PTX exposure, whereas cyclin G1 depletion by RNA interference delays mitotic slippage and enhances paclitaxel-induced apoptosis (129), providing molecular insights to guide future combinatorial treatment options. Moreover, additional insights into the molecular mechanisms of chemotherapeutic drug resistance have recently been uncovered: It was recently reported that miR-27b, a novel miRNA that regulates multidrug resistance (MDR) in gastric cancer, targets and suppresses CCNG1 expression and restores p53-dependent checkpoint activities. Ectopic expression of miR-27b increases the chemosensitivity of gastric cancer cells to several chemotherapeutic drugs by inhibiting CCNG1 (130). Likewise, ectopic expression of the tumor-suppressive miR-23b, which is downregulated in colon cancer and potently mediates multiple steps of metastasis, including tumor growth, invasion and angiogenesis, selectively targets CCNG1 for suppression (131): By binding with the 3’ untranslated region of CCNG1, miR-23b downregulates cyclin G1 mRNA and protein expression, thereby inhibiting ovarian tumorigenesis and cancer progression (132). A variety of molecular approaches targeting cyclin G1 expression and/or function for cancer control are shown in Table II.

Establishment of the once-paradoxical cyclin G1 proto-oncogene as a dominant, growth-promoting, potentially cancer-causing component of the commanding cyclin G1/Mdm2/p53 axis has enormous implications in terms of the future of clinical medicine, including cancer diagnostics, prognostics, and the potential for rational combinational therapies. The frequency of cyclin G1 overexpression observed in human cancers, as well as the newfound association to the molecular mechanisms of cancer metastasis (EMT) and
chemotherapy resistance, provide new avenues for patient profiling, staging and treatment, including chemosen- sitization by strategic targeting of cyclin G1 as a locus of diagnostic evaluation and clinical control. The advent of the injectable, tumor-targeted vector and its demonstrated safety in clinical trials, enables physicians of the future to think and to reach beyond previous limitations in clinical trial designs. The development of Mx-dnG1 (Rexin-G), which, by itself, induces apoptosis in cancer cells and tumor-associated vasculature (in the presence or absence of p53), is a powerful clinical tool in terms of applied cell cycle checkpoint control, which merits conscientious clinical development. Indeed, the functional characterization of discrete structural domains that are present within the cytocidal dnG1 have recently identified the ELAS1 peptide, which is based on a conserved sequence motif within the extended C-terminal domain of cyclin G1, as a therapeutically useful peptide drug (133). Reportedly, the ELAS1 peptide, which competitively blocks the physical association of cyclin G1 with the B’γ subunit of PP2A, sensitizes osteosarcoma cells to camptothecin and irinotecan, and induces apoptotic death in both prostate adenocarcinoma and squamous cell carcinoma, albeit in a wild-type T553-dependent manner. By comparison, the ability of dnG1 to activate apoptosis in the presence (hyper- plasia, neoangiogenesis) or absence (most metastatic cancer cells) of wild-type p53 predicates structure-function relationships involving programmed cell death, which remained to be identified within the truncated α-helices (molecular interfaces) of the dnG1 cyclin box domain. Further elaboration of the structure-function relationships of the cyclin box (134), which provide a mechanistic link to cell death/signaling mechanisms, came from screening studies of the ubiquitin-proteasome system, which has profound implications in cancer biology. In addition to degrading specific cyclin proteins during progression of the cell cycle, and generating peptide antigens that are bound and presented to the immune system by the major histocompatibility complex, the ubiquitin-proteasome system was also found to generate intracellular peptides with profound regulatory activity and importance (135,136). Accordingly, a comprehensive screen for such intracellular peptides in synchronized HeLa cells identified a small peptide derived from the proteolysis of cyclin D2, specifically the α5 helix of the cyclin box, which induces cell death via apoptosis and/or necrosis in all cell lines tested (i.e., broad-spectrum efficacy), including human cervical cancer, breast cancer, melanoma and thyroid tumor-derived lines (136). Moreover, when this cyclin D2 α5 helix-derived peptide is fused to a cell-penetrating peptide, the resulting construct, infused in vivo in a rat brain tumor model, reduced the volume of rat C6 glioblastomas by ~50% (136). Using a fluorescently labeled peptide monitored by real-time confocal microscopy in MDA-MB-231 breast cancer cells, the α5 helix-derived peptide entered the cells within min, localized to the nucleus as well as the cytoplasm, and induced cell death in breast cancer cells, specifically in G1/S-arrested cells and in cells that were actively progressing through S phase (137). Reasoning that the protein/protein interface and conformational changes required in the molecular mechanism of cyclin-dependent CDK activation would be potential targets for the design of specific inhibitors of cell cycle progression, a small peptide inhibitor (C4) derived from the α5 helix (aa 285-306) of cyclin A was found to be capable of binding to cyclin A/CDK2 complexes, thereby inhibiting the kinase activity of kinase complexes harboring CDK2 in a competitive manner (138). Intricate structural studies indicate that this C4 (α5 helix) peptide binds to the active site of CDK2 to interfere with the interaction of target phosphoacceptor substrates, independently of ATP binding in the catalytic site, thus overcoming one of the major drawbacks of inhibitors that target the ATP-binding site of protein kinases (which tend to result in poor selectivity) and introducing a novel class of designer
peptide inhibitors of cyclin-dependent kinase activation, with the potential to treat a wide range of tumor types (138). The cytotoxic efficacy of this C4 (α5 helix) peptide was validated in cellulo by demonstrating that proliferation of all the human tumor cell lines tested (breast cancer, liver cancer and T-cell leukemia) is blocked in a dose-dependent manner when the C4 (α5 helix) peptide is coupled to a cell-penetrating carrier peptide (138).

As both the crystal structure of cyclin A and the mechanism of cyclin A/CDK2 interaction have been extensively characterized (134), and the major contact points with the catalytic CDK subunit have been determined to be the α3, α4 and α5 helices of the cognate cyclin subunit, the demonstration that the ‘designer’ peptide inhibitor C4 (α5 helix) does not compete with cyclin A in terms of cyclin A/CDK2 complex formation, nor does it simply interact with monomeric CDK2; rather, the binding of cyclin A to CDK2 promotes a conformational change in CDK2 that exposes the substrate docking site on the CDK, thereby facilitating the binding of the C4 (α5 helix) across the catalytic cleft of CDK2 and blocking the phosphorylation of target substrates (138). Based on the profound mechanistic insights provided by this detailed structure-activity analysis, as well as the broad-spectrum anticancer efficacy demonstrated by the cyclin D1-derived α5 helix peptide, recently identified by functional proteomics (136,137), we may now present a clearer and more comprehensive picture of the molecular mechanisms by which the dnG1 construct, developed by direct experimentation (77) and embodied by Rexin-G in clinical medicine (98), is able to produce such broad-spectrum anticancer efficacy when administered repeatedly as a single therapeutic agent (Tables I and II).

Clearly, dnG1, which is structurally truncated, and yet retains the α3, α4 and α5 helices of the predicted substrate (c-Myc) docking site (Fig. 9), presents a rational, stable and competitive intracellular inhibitor of cyclin G1/CDK5 activity (and possibly cyclin G1/CDK2 activity), which blocks the phosphorylation/activation of c-Myc, thwarts proliferative competence and leads to cell death via apoptosis and necrosis in cancer cells derived from all three germ layers, save for artificially-immortalized ‘vector producer’ cell lines (e.g., 293T cells) that have been purposefully transformed by the strategic integration of the SV40 large T antigen, a dominant-acting oncprotein, which coincidentally targets pRb, p53 and PP2A for subversion (139) and often leads to malignant transformation. Retaining the structural interaction domains that bind Mdm2 and PP2A, the C-terminus of cyclin G1 not only exhibits the PP2A-B′γ association domain, it apparently ‘displays’ this domain at least an order of magnitude more efficiently as a protein than the ELAS1 peptide drug, which was designed to mimic, thus compete with, this interaction (133). Of note, this ELAS1 peptide (thus dnG1) was found to sensitize U2OS osteosarcoma cells to radiation therapy (RT), as well as camptothecan/irinotecan chemotherapy (140), thereby providing a new avenue for dnG1 (Rexin-G) to further improve the efficacy of chemoradiotherapy (CRT).

While the p53 tumor suppressor normally guards DNA fidelity and serves as the executioner (by inducing apoptosis) of aberrant cells, this locus of control is frequently lost in the process of cell transformation, tumorigenesis, invasion, EMT and/or metastasis, as the balance of power, i.e., the dominance of the commanding axis, shifts from p53 tumor suppression to the oncogenic cyclin G1/Mdm2 phenotype. From a therapeutic aspect, dnG1 functions mechanistically, hence clinically, to counteract the root proliferative competence of cancer cells, that being the pro-survival, anti-apoptotic, tumor-promoting actions of cyclin G1/CDK activity associated with the phosphorylation/activation of the c-Myc oncoprotein. In retaining the structural elements and interfaces of the commanding cyclin G1/Mdm2/p53 axis that interact with Mdm2 and PP2A, the dnG1 protein provides an additional means of sensitizing cancer cells to RT, cytotoxic chemotherapies and CRT. In the revealing crucible of clinical medicine, wherein investigational new drugs, such as Rexin-G, are initially tested under rather extreme conditions (e.g., late-stage, metastatic and/or chemotherapy-resistant cancers) wherein standard therapies have failed, the repeated clinical demonstrations of tumor eradication (in the presence or absence of p53) in multiform cancer types, including eradication of primary tumors, metastatic lesions, and even lymphatic metastasis (98,99), resulting in long-term dose-dependent survival benefits (101), provides a compelling molecular exemplar for the future of cancer gene therapy.

14. Conclusions along with a note concerning the future of clinical oncology

The commanding cyclin G1/Mdm2/p53 axis is presented herein with collegial appreciation for the multidisciplinary nature of the task: The decades of basic, translational and clinical cancer research that has brought this ‘accessible’ locus of checkpoint control to light. The model is purposefully simple yet powerful in its applications, intended to provide a logical framework for new information, including the development of pertinent companion diagnostics, and to serve as a guide for clinical oncologists to better understand the executive regulatory components that determine the efficacy and outcomes of cancer therapies through the increasingly complex molecular mechanisms of cell cycle checkpoint control. This review is presented in appreciation of the numerous cancer patients whose informed consent has served to teach and tell, and whose long-term, cancer-free survival stands as a meaningful milestone in clinical oncology.

The broad-spectrum, single-agent efficacy of Mx-dnG1 (Rexin-G) demonstrated in end-stage cancers, where standard therapies had failed, is an excellent step forward; and one can only imagine what improved clinical outcomes may arise when the physiological surveillance properties of the tumor-targeted Mx-dnG1 vector is eventually permitted to be administered worldwide at an earlier, less intractable stage of metastatic cancer (Figs. 5 and 8). Furthermore, these first pioneering demonstrations of clinical safety, efficacy and long-term survival of Mx-dnG1 (Rexin-G)-treated cancer patients serve as a challenge to those who may otherwise distort or obscure what has been historically accomplished, and to serve as a guidepost for the next generation of clinical oncologists who will assuredly be more knowledgeable and skilled in applied molecular medicine. Finally, this review article is presented well in advance of the time when genetic medicine will no
longer be viewed as a competitive (or disruptive) biotechnology, but will rather be considered a rational and beneficial addition to the combinatorial practice of the postmodern medical oncologist.

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Availability of data and materials

The materials included in this manuscript, including all relevant raw data, will be made freely available to any researchers who wish to use this for non-commercial purposes, while preserving any necessary confidentiality and anonymity.

Authors’ contributions

EG reviewed the published literature, wrote the manuscript, analyzed the critical stages in Rexin-G drug development, reviewed and edited the final manuscript. JR reviewed the published literature, wrote sections of the manuscript and the reference section, and edited the final manuscript. SL reviewed the published literature, wrote and edited the reference section, and edited the final manuscript. SC reviewed the published literature, analyzed the clinical trials using Rexin-G, reviewed and edited the final manuscript. FH reviewed the published literature, analyzed the molecular mechanisms of action of cyclin G1 in the cell cycle control pathway, wrote the manuscript, reviewed and edited the final manuscript. All authors approve the submission of this manuscript for publication.

Ethics approval and consent to participate

Not applicable.

Patient consent for publication

Not applicable.

Competing interests

Drs Gordon and Hall are co-inventors of Rexin-G targeted gene delivery system, which was developed at the University of Southern California Keck School of Medicine, and are co-founders of DELTA Next-Gen, LLC. Dr Chawla, Mr. Ravicz and Mr. Liu have no competing interests to disclose.

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