Targeting the telomere and shelterin complex for cancer therapy: current views and future perspectives

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Abstract

Aberrant telomere homeostasis is essential for cell immortality, enabling cells to evade telomere dependent senescence. Disruption of telomere structure and function in cancer cells is highly toxic as shown by detailed pre-clinical evaluation of telomerase inhibitors. Under telomerase inhibition, cells must divide sufficiently frequently to allow one or more telomeres to shorten to an unprotected length. Functioning telomeres are disguised from the DNA damage machinery by DNA remodelling and other activities of the telomere binding complex shelterin. Direct interference with shelterin has been shown to result in cell killing and small molecules directly targeting telomere DNA also have anti-tumour effects partially dependent on shelterin disruption. However, shelterin components have not generally been regarded as therapeutic targets in their own right. In this review, we explore the possibilities for therapeutic targeting of the shelterin complex.

Keywords: shelterin • telomere • telomerase • telosome • targeting • novel therapeutics • cancer • senescence

Introduction

In recent years, understanding of the molecular mechanisms underlying tumour cell biology has provided a variety of potential targets for drug development. One important current goal is to determine the best targets for therapeutic intervention given a particular molecular aetiology. For some targets it will be possible to develop highly specific and effective treatments, though it is likely that many pathways will prove refractory to targeting in the clinical setting. In particular, many leads with promising pre-clinical activity may ultimately prove effective only in specific patient subsets [1]. Therefore, agents with broad spectrum activity will presumably form the backbone of cancer therapy for some time to come.

Immortality is a near-universal phenotype of cancer cells that has attracted considerable interest recently in respect of therapeutic interventions [2]. Most normal somatic cells have a pre-defined maximum lifespan in vitro, bounded by the onset of replicative senescence. In contrast, tumour cells circumvent senescence and divide indefinitely. The complete network of signalling events that underlie senescence establishment and maintenance are poorly understood. However, the central role of telomere homeostasis is now firmly established [3].

Mammalian telomeric DNA comprises extended repeats of the sequence TTAGGG ending in a 3’ G-rich single stranded overhang on both termini. The natural end must be protected from recognition as damaged DNA to prevent cell cycle arrest in proliferating cells. In normal somatic cells the end replication problem, oxidative stress and processing by nucleases cause progressive telomere shortening during each round of cell division. Ultimately, compromised protection causes DNA damage signalling at critically shortened telomeres and telomere dependent senescence [3]. In contrast, most cancer cells aberrantly express telomerase to counteract telomere attrition, maintaining telomeres at a stable length [4].

Telomerase is present in the vast majority of human tumours and directly controls immortality, whereas its core RNA (hTR) and protein (hTERT) subunits are transcriptionally repressed in normal cells [2]. Thus, it is widely regarded as a highly attractive cancer target and has been the subject of intense drug development...
efforts. Thorough pre-clinical validation of multiple telomerase targeting approaches in cancer and normal cells has proven reproducibility of results across multiple in vitro and in vivo model systems in a large number of studies [2, 4]. Telomerase enzyme inhibition and targeting of hTR in cancer cells generally results in progressive telomere shortening and delayed onset senescence in a telomere length dependent manner, while a rapid growth inhibition and apoptosis induced by dysfunctional telomeres has been documented with hTERT targeting agents [5, 6]. In contrast, normal cells are usually unaffected. Encouragingly, several telomerase-directed therapies are now in clinical trial [2, 4].

Telomerase inhibition with the oligonucleotide enzyme inhibitor GRN163L provides indisputable pre-clinical proof of concept that induction of telomere dysfunction in cancer cells is an attractive therapeutic mechanism and there is good reason to be optimistic about its clinical prospects [2, 4]. However, evaluation is at an early stage and in a worst-case scenario that efficacy is not demonstrated, there are currently no alternative small molecule telomerase enzyme inhibitors scheduled for clinical trials.

A second class of agent directly targeting telomeric DNA secondary structure have also been investigated and found to cause toxicity in cancer cells (G-quadruplex (G4) targeting agents, GTAs). It was originally envisaged that these would block access of telomerase to the G-overhang. However, an emerging consensus is that GTAs elicit their effects at least in part by affecting the t-loop, it may provide 3’ end protection. Telomere repeat binding factor 2 (TRF2) affects formation of telomeric G4 in vitro and, conversely, G4 DNA may affect the function of shelterin components [7, 8, 9]. The main attempt so far to directly target telomeres in cancer cells is through the use of agents that bind and stabilize G4 DNA (GTAs). Because G4 DNA must be resolved to allow telomerase access, GTAs were originally envisaged as inhibitors of telomerase primer extension, though they are now also known to cause more rapid toxic effects. Many chemically diverse GTAs have been described [7]. Our focus is on the development paths of three agents that have come close to clinical testing.

A diamidoanthraquinone was the first reported GTA, showing slight preference for binding G4 over duplex DNA [10]. Intensive structure-activity optimization led to development of the trisubstituted acridine BRACO19 [11]. BRACO19 inhibits telomerase activity in cancer cells at sub-toxic concentrations and can reduce telomere length, though rapid telomere length independent growth suppression with increased telomere fusions has also been observed [12–14]. Growth suppression of uterine cancer and paclitaxel pre-treated vulval carcinoma xenografts was observed following delivery by the intraperitoneal route, though oral dosing was found to be inactive, presumably because the compound exhibits poor epithelial transport and is relatively unstable [12, 13, 15]. Derivative AS1410 was lead compound for a phase I trial planned by Antisoma Plc (London, UK). In 2005, the company indicated to investors that the trial would not commence because of compound toxicity unrelated to mechanism of action and that another lead would be selected (http://www.antisoma.com). In January 2008, collaboration was established with Argenta Discovery to deliver improved GTA candidates (http://www.argentadiscovery.com).

RHPS4 is a pentacyclic acridium salt with good specificity for binding triplex and G4 DNA over other conformations. It inhibits telomerase activity, has relatively low toxicity, is efficiently taken up into breast and lung cancer cells and is stable in culture medium [16]. Sub-toxic doses of RHPS4 inhibit telomerase in several cell lines but can also lead to rapid growth suppression and G1 arrest in the absence of significant telomere shortening [17]. However, analysis of the sensitivities of 36 tumour cell lines established that short telomere length does correlate with RHPS4 sensitivity [18].

The compound exhibits broad spectrum activity against a range of tumour cell lines in vitro and in xenograft models of melanoma and uterine, prostate, colorectal, breast and lung cancer [17–20]. Furthermore, it efficiently potentiates the activity of several other chemotherapy agents. However, context dependent effects have been observed: combination with paclitaxel was synergistic in MCF7 breast cancer cells but antagonistic in M14 melanoma cells [18, 19]. In 2006, Pharminox agreed in-licensing of rights to preclinical development of RHPS4 (http://www.pharminox.com). Two related acridinium salts were recently identified as potential backup leads on the basis of improved quadruplex binding specificity and low non-specific toxicity [21]. Additionally, a new and more flexible synthetic route has been described for RHPS4 and substituted derivatives [22].

Telomestatin, a natural macrocyclic pentaoxazole isolated from Streptomyces anulatus, is among the most efficient G4 stabilizing agents discovered [23]. Telomestatin inhibits telomerase activity and causes telomere shortening and apoptosis in a range of cancer cell lines in vitro and inhibits growth of leukaemia xenografts [24, 25]. Treatment also augmented apoptosis induced by daunorubicin, mitoxantrone and vincristine in human leukaemia cell lines and enhanced inhibition of colony formation by imatinib in primary chronic myeloid leukaemia (CML) cells [26]. In vivo evidence of telomestatin efficacy is currently limited, though suppression of human leukaemia cell xenografts has been shown [25].

The pharmaceutical company Sosei was to undertake collaborative pre-clinical development of telomestatin (GM-95/SOT-095)
shortening, degradation of the 3' overhang and loss of POT1 and/or TRF2 binding (reviewed in [6, 8]). Interestingly, GTAs also elicit a growth suppressive effect on cells utilizing the alternative recombination-based mechanism of telomere maintenance, alternative lengthening of telomeres (ALT) [6]. ALT is active in around 10–15% of human tumours and prevalent in certain tumour types with poor outcome including those of mesenchymal origin [30]. These cells do not express telomerase activity so are generally refractory to telomerase inhibition, making the effective targeting of ALT cells an additional advantage to GTAs over other telomerase inhibitors. RHPS4 was shown to induce short-term growth arrest in the ALT cell line GMB47 [17] in addition to interfering with telomere replication to induce telomere dysfunction and Ataxia Telangiectasia and Rad3 related (ATR)-dependent Ataxia Telangiectasia Mutated (ATM) signalling [31]. ALT cells are characterized by long heterogeneous telomeres, which provide a larger substrate for GTAs to bind. This may enhance the efficacy of G4 stabilization and prevent recombination required for telomere maintenance. Consistent with this telomestatin has been observed to block topoisomerase III recruitment to telomere G4 DNA in ALT cells resulting in telomere uncapping and DNA damage signalling [32].

Despite these potentially non-selective effects on telomerase there is growing evidence that GTAs exhibit tumour specificity. BRACO19 caused acute toxicity in breast, prostate, colorectal, ovarian, lung and gastric cancer cell lines at lower concentrations than in two normal lung fibroblast strains [33]. Concentrations of telomestatin effective against leukaemia cells had little effect on normal CD34+ cells or normal fibroblasts while concentrations causing acute toxicity and telomere uncapping in cervical and breast cancer cells did not affect normal fibroblast or breast epithelial cells [25, 26, 34]. Similarly, RHPS4 treatment induced DNA damage signalling, telomere dysfunction and toxicity in melanoma cells but not in normal fibroblasts [20]. Alongside promising results from in vivo models, it is possible that many tumour cells may be ‘primed’ for telomere dependent death, suggesting that targeting shelterin directly may also have specificity.

Composition of the shelterin complex

The core reactions of telomere homeostasis are controlled by the six-member complex shelterin, comprising the proteins TRF1 and TRF2, POT1, TRF1-interacting protein 2 (TIN2), transcriptional repressor/activator protein 1 (RAP1) and POT1- and TIN2-organizing protein (TPP1) [PTOP/TINT1/PIP1] [3]. Double stranded telomeric DNA is bound directly by homodimers of TRF1 and TRF2 which differentially recruit other proteins to the telosome, including shelterin proteins TIN2 and RAP1 (recruited by TRF1 and 2, respectively), in addition to accessory factors with roles in DNA damage signalling and repair such as Apollo [35]. The single stranded G-overhang is bound by the POT1/TPP1 complex, in which TPP1 stimulates POT1 recruitment to the telomere in vitro and in vivo [36]. Finally, the single strand and double strand binding subcomplexes are physically linked via interaction between TPP1 and TIN2 [3].

TRF1/2 share similar domain organization, though the N-terminus of TRF1 has an acidic region, whereas that of TRF2 is rich in Gly/Arg residues (basic region). The TRF2 N-terminus displays sequence non-specific DNA binding activity targeting to replication forks and Holliday junctions which may affect telomere replication and end protection [3, 37]. TRF1/2 perform a range of DNA remodelling activities: TRF1 bends DNA and promotes DNA looping and strand pairing by binding to non-adjacent half sites with conformational flexibility [3]. TRF2 binding to the junction of the G-overhang promotes formation of t-loops, which provide 3' end protection by sequestration of the G-overhang within the telomere duplex. Recent observations point to a mechanism involving DNA melting mediated by super-coiling [3, 38].

Shelterin proteins diversely affect telomere length, with TRF1, TIN2 and RAP1 implicated in negative length regulation. One aspect of TIN2 activity appears to be suppression of TRF1 PARsylation by tankyrase, a telomere-associated PARP which targets TRF1 to the proteasome pathway [39]. TRF2 was also characterized as a negative regulator in overexpression experiments; however, TRF2 inhibition, like POT1, results in rapid telomere deprotection [3, 6, 40]. POT1 appears to have a dual role in length regulation: protection of the single stranded region may prevent telomerase access and in this context POT1 has been implicated as the effector of TRF1 dependent negative length regulation [41]. However, TPP1 directly interacts with telomerase, recruiting it to the telomere [42] and the POT1/TPP1 complex stimulates telomerase processivity [36, 43].

In addition to length regulation, shelterin proteins play essential roles in suppression of DNA damage signalling and inappropriate repair by homologous recombination and non-homologous end-joining. Disruption of these protective functions (telomere ‘uncapping’) occurs in ageing normal cells or can be achieved by dominant negative and knockdown approaches targeting TRF2 or POT1 which are essential for suppression of ATM and ATR, respectively [44]. TRF2 or POT1 presumably act in part by blocking access of the Mre11/Rad50/Nbs1 complex (MRN) and replication protein A (RPA) damage sensors to telomeric DNA [3]. Similar to the ultimate outcome of telomerase inhibition, though with far more rapid effects, such interventions induce highly fus-
Shelterin targeting approaches

Due to the complexity of drug development even in its early phases, in order to have confidence that targeting the telomere will be fully validated in the clinic it is appropriate to explore a range of targets. Therefore, it is timely to consider strategies for compound discovery with shelterin as the focus. However, the protein components of shelterin are not generally regarded as potential targets in their own right, with intervention studies frequently focused on functional rather than translational questions [6]. Because shelterin components may prove difficult to drug, particularly in the case of potentially targeting protein–protein interactions, novel discovery solutions may need to be applied to identify promising leads. Encouragingly, there is a growing list of small molecule inhibitors of similar unconventional targets. Development of the Bcl2 homology domain 3 (BH3)-mimetic Bcl2 inhibitor ABT-737 using ‘structure/activity relationship (SAR) by nuclear magnetic resonance (NMR)’ provides notable proof-of-concept that fragment based screening coupled with structure led design is an appropriate approach [47].

As an example of the application of this general strategy to shelterin, we submitted the structure of the TRF1/TIN2-peptide interaction ([35] (Protein Data Bank code 3BOO) for computational solvent mapping on the FTMAP server (http://ftmap.bu.edu). The FTMAP algorithm is a computational solution to detect candidate druggable sites on protein surfaces building on observations from NMR-screening and ‘multiple solvent crystal structures’ techniques that druggable ‘hotspots’ can be identified on the basis of their ability to bind many individual small molecule fragments/probes irrespective of affinity [48]. Computational docking of 16 small molecule probes into PDB structures is performed with a fast Fourier transform correlation approach to identify and rank likely clustering sites. The algorithm correctly identifies known drug-binding sites in several proteins [48].

Nine probe clusters were found with the top ranking cluster localized in a cleft adjacent to the TIN2 peptide binding site (Fig. 1A). Mutation either of TRF1-F142 or of TIN2-L260 is sufficient to ablate TRF1/TIN2 interaction in 293T cells [35]. The predicted site is in close juxtaposition with F142 and at the lip of the hydrophobic pocket which interacts with L260. The same analysis performed for the TRF2/TIN2 structure 3BU8 [35] also revealed a candidate druggable pocket in TRF2 bounded by the stacked helices 1 and 3 and the loop region connecting helices 8 and 9. This site is distal from the TIN2 probe, though might conceivably play a role in other TRF2 protein interactions. Similarly a structure was recently reported for an SCF^B^x^4^–TRF1 interaction. Fb^x^4^, an F-box protein subunit of Skp, Cullin, F-box containing (SCF) ubiquitin E3 ligase, mediates ubiquitination and degradation of TRF1. This structure may represent an additional target for development of small molecule inhibitors [49].

Given the considerable structural understanding of several shelterin components [3], structure based design using similar algorithms seems a rational approach to identify suitable ligands.

Cell based assays could provide a second general approach to identify shelterin inhibitors [2]. Importantly, many interactions could be addressed without pre-existing knowledge of the pathway anatomy. We recently reported that promoter screening can identify inhibitors of hTERT expression [50]. Similar assays might be justified for some shelterin related targets such as TRF2, which is overexpressed in several human tumours and may play a role in drug resistance in some settings [6, 51, 52]. Similarly the telomere uncapping phenotype itself could be used in a screening approach to identify potential inhibitors of telomere dysfunction or telomerase itself. In addition, the availability of a good cellular model of telomere uncapping will enable further investigation of the signalling events underlying the telomere uncapping response and may identify new biomarkers of telomere dysfunction, which could aid future drug discovery strategies targeting the telomere (Fig. 2A). We designed a cell based screening assay, which utilizes an adenviral vector expressing a form of hTR mutated in the template region for reverse transcription by the telomerase catalytic subunit hTERT (Ad-hTR-mut). When incorporated into telomeres this template synthesizes mutant telomere repeat sequences of the template TATATATATAA [53], which are predicted to be incapable of binding key components of the shelterin complex, POT1, TRF1 and TRF2. Expression of this construct in cancer cell lines induces rapid telomere uncapping, DNA damage signalling and cytotoxicity in an hTERT-dependent manner [53–57], thereby circumventing the problem of phenotypic lag which has hampered identification of telomerase inhibitors through screening assays in the past. In the context of this screening assay any inhibitors of telomere uncapping or the downstream signalling events will result in increased cell viability as measured by 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay, therefore any compounds inducing non-specific toxic effects will be removed in the initial screening process. With an estimated proportion of 30% of drugs failing in clinical trial due to non-specific toxicity [58] this is a tantalizing prospect from a drug development point of view.

Three validated reagents, known telomerase inhibitors BIBR1532 [59, 60] and suramin [61] and commercially available general inhibitor of ATM/ATR kinase, postulated to rescue viral
cytotoxicity by blocking incorporation of mutant telomere repeats through inhibition of telomerase in the case of BIBR1532 and Suramin [60, 61], or by blocking the downstream DNA damage signalling that ensues from telomere uncapping in the case of the ATM/ATR inhibitor [62], increased cell viability compared to Ad-hTR-mut alone (Fig. 2B). These proof-of-concept studies show the power of such a model system in combination with well-defined reagents including small molecules or siRNA, as a screening approach and as a tool to probe signalling pathways downstream of telomere uncapping. More detailed investigation of shelterin expression profiles is clearly needed and well validated and standardized biomarker assays would be an advantage in this respect.
Inhibiting the enzymes that post-translationally modify shelterin could also be an attractive therapeutic mechanism, as has previously been suggested for telomerase. Tankyrase-1 is widely considered a good target in this regard, while a new candidate is the Pin1 prolyl isomerase which also contributes to TRF1 instability and might provide an alternative or complementary target [39, 63]. Other known modifications of TRF1 include phosphorylation by Plk1 which promotes DNA binding [64]. Therefore, blocking negative regulators of Plk1 might similarly enhance TRF1 telomere association. Because regulation of the stability and/or localization of shelterin components is of major functional importance for telomere biology, existing functional fluorescent tagged shelterin expression constructs could rapidly be put to use as screening reagents to interrogate this class of pathways [65, 66]. Importantly, screening could further enhance understanding of telomere homeostasis even if not all interactions constitute valid therapeutic targets.

**Conclusion**

To identify potential new anti-cancer agents, knowledge from mechanistic studies of tumour cell biology must be employed to evaluate potential targets in the translational setting. Increasing evidence suggests that directly targeting telomeric DNA has acceptable specificity for cancer cells. Therefore, agents directed against the shelterin complex may also have anti-tumour activity. A growing list of examples suggests that unconventional targets should not be ignored in drug discovery. Approaches such as structure-led design and cell-based screening might provide pharmacological access to ‘undruggable’ proteins in the shelterin complex. The most effective telomere directed therapies will be identified by exploration of all targets using suitable assays based on mechanistic understanding of telomere homeostasis and the unique biology of each candidate. However, well-designed drug discovery assays could generate novel small molecule tools to further probe telomere biology even if high quality clinical leads are not identified.

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**Conflict of interest**

The authors confirm that there are no conflicts of interest.
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