

# Unanticipated differences between $\alpha$ - and $\gamma$ -diaminobutyric acid-linked hairpin polyamide-alkylator conjugates

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## ABSTRACT

Hairpin polyamide–chlorambucil conjugates containing an  $\alpha$ -diaminobutyric acid ( $\alpha$ -DABA) turn moiety are compared to their constitutional isomers containing the well-characterized  $\gamma$ -DABA turn. Although the DNA-binding properties of unconjugated polyamides are similar, the  $\alpha$ -DABA conjugates display increased alkylation specificity and decreased rate of reaction. Treatment of a human colon carcinoma cell line with  $\alpha$ -DABA versus  $\gamma$ -DABA hairpin conjugates shows only slight differences in toxicities while producing similar effects on cell morphology and G<sub>2</sub>/M stage cell cycle arrest. However, striking differences in animal toxicity between the two classes are observed. Although mice treated with an  $\alpha$ -DABA hairpin polyamide do not differ significantly from control mice, the analogous  $\gamma$ -DABA hairpin is lethal. This dramatic difference from a subtle structural change would not have been predicted.

## INTRODUCTION

The ability to control gene expression through the use of DNA sequence-specific, cell-permeable molecules holds therapeutic promise. Based on the natural product distamycin A, pyrrole–imidazole polyamides are a class of molecules that can be programmed to bind a broad repertoire of DNA sequences with specificities comparable to naturally occurring DNA-binding proteins (1). They have also been shown to permeate living cells, localize to the nuclei (2,3) and regulate gene expression (4,5).

Pairing rules have been established for minor groove recognition, whereby an *N*-methylpyrrole/*N*-methylimidazole (Py/Im) pair recognizes C•G, the reverse (Im/Py) recognizes G•C and Py/Py specifies A•T or T•A (1,6). The replacement

of Py with  $\beta$ -alanine ( $\beta$ ) can enhance binding affinity by allowing the polyamide to reset its curvature with that of the DNA minor groove (7,8). Polyamides may be linked via a turn moiety to give a hairpin structure which binds in the ‘forward’ direction N→C with respect to the 5′→3′ direction of the DNA strand (9). In earlier studies, linkage of antiparallel three-ring subunits via a  $\gamma$ -aminobutyric acid moiety, resulting in a turn with three methylene units, was shown to enhance DNA-binding affinity ~100-fold over unlinked subunits (10). The  $\gamma$ -aminobutyric acid turn demonstrates selectivity for A•T or T•A base pairs over G•C and C•G (11). Use of a glycine amino acid, with one methylene turn unit, resulted in a 3-fold reduction in affinity relative to the  $\gamma$ -aminobutyric acid. Subsequent NMR studies revealed that the glycine amino acid substituted polyamide bound DNA in an extended, rather than a hairpin, conformation (12,13).

Substitution of the  $\gamma$ -aminobutyric acid with the chiral (*R*)-2,4-diaminobutyric acid [(*R*)- $\gamma$ -DABA] was later found to further increase DNA-binding affinity ~10-fold (14). In contrast, hairpin polyamides with the opposite enantiomer (*S*)- $\gamma$ -DABA result in an unfavorable steric clash with the DNA minor groove and decreased binding for the forward orientation. Remarkably, binding affinity to the opposite orientation, in which the polyamide binds C→N with respect to the 5′→3′ direction of the DNA strand, is energetically favorable due to relief of the steric clash.

Polyamides have been used to regulate gene expression by binding to promoter and enhancer elements, but they do not prevent RNA polymerase elongation when bound to the coding region of genes (1). Conjugation of an alkylating moiety to a sequence-specific DNA-binding molecule would be expected to result in a molecule capable of covalent DNA attachment that could arrest transcription (15,16). An early example of this is Tallimustine (FCE 24517), which is a conjugate of distamycin A and the nitrogen mustard chlorambucil (17). When attached to a DNA minor groove binder, chlorambucil is capable of alkylation at the N3 position of an adenine proximal to the binding site (18,19). Hairpin

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polyamide–chlorambucil conjugates have been demonstrated to target specific sequences of the HIV-1 promoter and simian virus 40 (19,20).

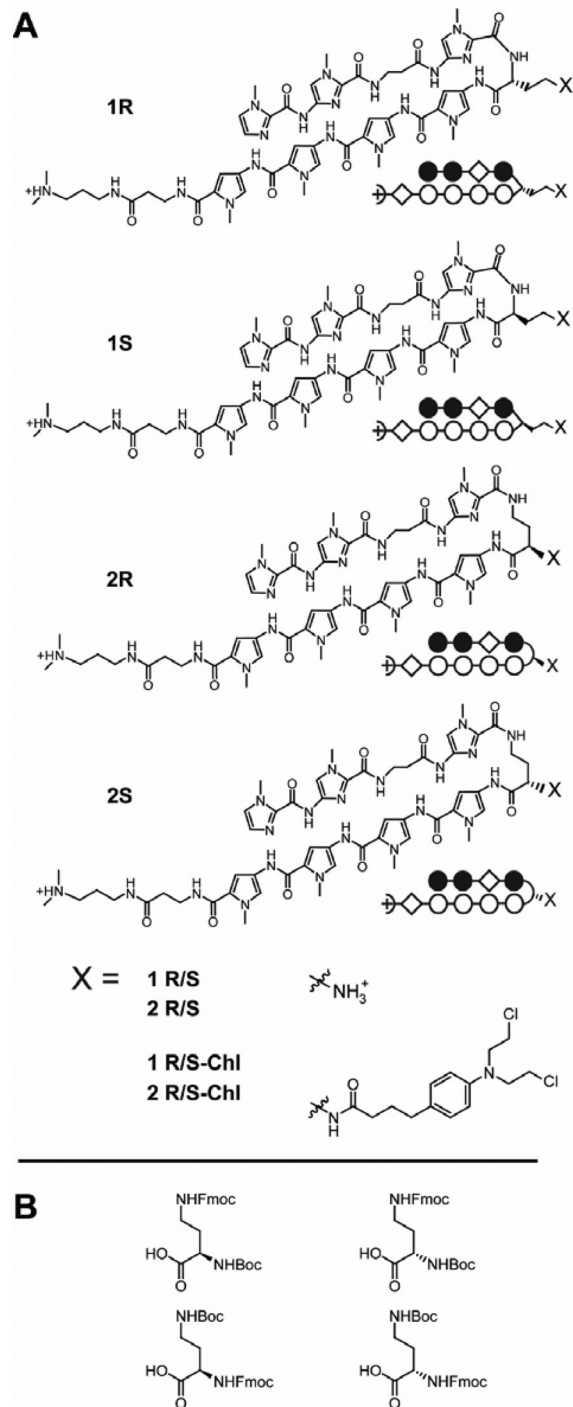
Recently, a hairpin polyamide–chlorambucil conjugate, synthesized at The Scripps Research Institute, was found to alkylate within the coding region of the histone H4c gene of the human colon carcinoma cell line SW620 and to block proliferation of these cells in culture (21). In addition, this conjugate was found to block tumor growth in a SW620 xenograft mouse model without apparent toxicity. When we attempted to scale-up additional conjugate at Caltech, the newly-synthesized molecule was found to be highly toxic in athymic nude mice, contrary to prior experiments (22). Subsequent studies showed that the two laboratories had utilized two different turn units in the synthesis of the conjugate: the conjugate synthesis at Scripps used Boc-D-Dab(Fmoc)-OH, while the conjugate synthesis at Caltech used Fmoc-D-Dab(Boc)-OH. The molecules are therefore constitutional isomers differing in the hairpin turn moiety. The original molecule made at Scripps continues to be called **1R-Chl** (22) and the new molecule is designated **2R-Chl**. **1R-Chl** utilizes an  $\alpha$ -DABA turn unit, whereas **2R-Chl** employs the standard  $\gamma$ -DABA turn unit (Figure 1). Linkage via the new  $\alpha$ -DABA turn unit results in one methylene group in the hairpin, whereas **2R-Chl** has three methylene groups. In addition, the linker between the polyamide core and the chlorambucil alkylator in **1R-Chl** is lengthened by two methylene groups relative to **2R-Chl**.

Polyamides utilizing the  $\alpha$ -DABA turn moiety have not been previously studied and the reasons for the unique biological properties displayed by **1R-Chl** are unknown. In the current study, we report the chemical and biological characterization of  $\alpha$ -DABA polyamides and compare them to the analogous  $\gamma$ -DABA polyamides. A series of four parent molecules and their corresponding chlorambucil conjugates were synthesized (Figure 1A). Each molecule contains the heterocyclic sequence: ImIm $\beta$ Im-HT-PyPyPyPy- $\beta$ Dp (where HT = hairpin turn and Dp = 3-(dimethylamino)-propylamine). Molecules in series **1** contain the new  $\alpha$ -DABA turn and molecules in series **2** utilize the  $\gamma$ -DABA turn. The *R* and *S* enantiomers of the parent polyamides and chlorambucil conjugates in each series have been generated for comparison. The DNA-binding affinities of the parent molecules were measured by DNase I footprinting and the alkylation profiles (sequence specificity and reactivity) of the polyamide–chlorambucil conjugates were established using thermal cleavage assays. The effects of polyamide treatment on SW620 cell morphology, proliferation, viability and cell cycle profiles were compared using phase contrast microscopy, cell counting, trypan blue exclusion assay and fluorescence-assisted cell-sorting (FACS) analysis, respectively. Importantly, toxicity effects of the polyamide constitutional isomers on mice were determined.

## MATERIALS AND METHODS

### Polyamide synthesis and characterization

Polyamides were synthesized on solid phase on Boc- $\beta$ -Pam resin using Boc-protected monomers and dimers according to protocols described previously (23). The following monomers/dimers were used in generating each of the



**Figure 1.** Polyamide structures and syntheses. (A) Chemical and ball-and-stick structures for polyamides **1R**, **1S**, **2R**, **2S** and their **Chl** conjugates. (B) Chemical structures of the four turn units used in the syntheses of the parent polyamides and polyamide-chlorambucil conjugates: Boc-D-Dab(Fmoc)-OH (upper left, used in the syntheses of **1R** and **1R-Chl**); Boc-Dab(Fmoc)-OH (top right, used in the syntheses of **1S** and **1S-Chl**); Fmoc-D-Dab(Boc)-OH (lower left, used in the syntheses of **2R** and **2R-Chl**); Fmoc-Dab(Boc)-OH (lower right, used in the syntheses of **2S** and **2S-Chl**).

molecules: Boc-Py-OBt (OBt = benzotriazol-1-yloxy), Boc- $\beta$ -Im-OH and Im-Im-OH. Boc deprotection was conducted at r.t. for 30 min using 80% TFA/DCM prior to the first coupling reaction and after addition of each heterocycle. Carboxylic

acids were activated with *N,N*-diisopropylethylamine (DIEA) and 2-(1H-benzotriazol-1-yl)-1,1,3,3-tetramethyluronium hexafluorophosphate (HBTU) for 30 min at 37°C. Monomer and dimer couplings were allowed to continue for 1 h (OBt esters) or 2 h (activated carboxylic acids). Polyamides were cleaved from resin with 3-(dimethylamino)-propylamine neat at 55°C for ~18 h and subsequently purified by reverse-phase high-performance liquid chromatography (HPLC).

Conjugation of polyamides to chlorambucil (4 eq) proceeded by activating the carboxylic acid with HBTU (4 eq) and DIEA (excess) for 30 min at r.t., after which the activation mixture was added to the polyamide. After allowing the reaction to continue for 1.5–2 h, products were purified by reverse-phase HPLC and immediately lyophilized.

The purity of all compounds were established by analytical HPLC and matrix-assisted laser desorption ionization-time-of-flight (MALDI-TOF) mass spectrometry.

**1R: ImImβIm-(R)<sup>H2N</sup>α-PyPyPyPy-βDp** Synthesized using Boc-D-Dab(Fmoc)-OH. MALDI-TOF C<sub>54</sub>H<sub>71</sub>N<sub>22</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1188.28, found: 1188.02

**1S: ImImβIm-(S)<sup>H2N</sup>α-PyPyPyPy-βDp** Synthesized using Boc-Dab(Fmoc)-OH. MALDI-TOF C<sub>54</sub>H<sub>71</sub>N<sub>22</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1188.28, found: 1187.99

**2R: ImImβIm-(R)<sup>H2N</sup>γ-PyPyPyPy-βDp** Synthesized using Fmoc-D-Dab(Boc)-OH. MALDI-TOF C<sub>54</sub>H<sub>71</sub>N<sub>22</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1188.28, found: 1187.95

**2S: ImImβIm-(S)<sup>H2N</sup>γ-PyPyPyPy-βDp** Synthesized using Fmoc-Dab(Boc)-OH. MALDI-TOF C<sub>54</sub>H<sub>71</sub>N<sub>22</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1188.28, found: 1187.77

**1R-Chl: ImImβIm-(R)<sup>Chl</sup>α-PyPyPyPy-βDp** Synthesized from **1R**. MALDI-TOF C<sub>68</sub>H<sub>88</sub>Cl<sub>2</sub>N<sub>23</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1474.48, found: 1474.98

**1S-Chl: ImImβIm-(S)<sup>Chl</sup>α-PyPyPyPy-βDp** Synthesized from **1S**. MALDI-TOF C<sub>68</sub>H<sub>88</sub>Cl<sub>2</sub>N<sub>23</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1474.48, found: 1475.03

**2R-Chl: ImImβIm-(R)<sup>Chl</sup>γ-PyPyPyPy-βDp** Synthesized from **2R**. MALDI-TOF C<sub>68</sub>H<sub>88</sub>Cl<sub>2</sub>N<sub>23</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1474.48, found: 1474.92

**2S-Chl: ImImβIm-(S)<sup>Chl</sup>γ-PyPyPyPy-βDp** Synthesized from **2S**. MALDI-TOF C<sub>68</sub>H<sub>88</sub>Cl<sub>2</sub>N<sub>23</sub>O<sub>10</sub><sup>+</sup> calculated [M + H]<sup>+</sup>: 1474.48, found: 1474.73

### Construction of plasmid pMFST2

Oligonucleotides were purchased from Integrated DNA Technologies. The plasmid pMFST2 was constructed by annealing the two oligonucleotides 5'-AGCTGCGCCCTTTAGGTGT-TACGTAGTGGTGGCGTAGGTCTTAGCCGTAGCTGTT-GCCGTAAGGGCGAATTCTGC-3' and 5'-GATCGCAGAA-TTCGCCCTTACGGCAACAGCTACGGCTAAGACCTAC-GGCACCACTACGTAACACCTAAAGGGCGC-3', followed by ligation into the BamHI/HindIII restriction fragment of pUC19 using T4 DNA ligase. The plasmid was then transformed into *Escherichia coli* JM109 competent cells. Ampicillin-resistant white colonies were selected from 25 ml Luria-Bertani (LB) agar plates containing 50 mg/ml ampicillin treated with XGAL and isopropyl-β-D-thiogalactopyranoside (IPTG) solutions and grown overnight at 37°C. Cells were harvested the following day and purification of the plasmid was performed with a Wizard Plus Midiprep DNA purification kit (Promega). DNA sequencing of the plasmid insert was

performed by the sequence analysis facility at the California Institute of Technology.

### Preparation of 5' <sup>32</sup>P-end-labeled DNA

The primer 5'-GAATTCGAGCTCGGTACCCGGG-3' was labeled at the 5' end and subsequently used with the primer 3'-CAGCCCTTTGGACAGCACGGTC-5' to amplify plasmid pMFST2 as described previously (24).

### DNase I footprint titrations

Polyamide equilibrations and DNase I footprint titrations were conducted on the 5' end-labeled PCR product of pMFST2 according to standard protocols (24). DNA was incubated with polyamide conjugates or water (control) for 12 h at r.t. prior to reaction with DNase.

### Thermal cleavage assays

Thermal cleavage assay experiments were conducted on the 5' end-labeled PCR product of pMFST2 as described previously (19). DNA was incubated with polyamide conjugates or water (control) for 24 h at 37°C prior to work-up, unless otherwise noted.

### Cell culture

The cells used in this work were derived from the human colon adenocarcinoma cell line SW620 (purchased from ATCC) and were maintained in Leibovitz's L-15 medium (Gibco) supplemented with 10% heat-inactivated FBS (Cambrex), 10 mM HEPES buffer (Gibco), 1 mM sodium pyruvate (Gibco) and 1% antibiotic-antimycotic (Invitrogen). Cells were grown in a humidified 5% CO<sub>2</sub> atmosphere at 37°C. For polyamide treatment experiments, cells were plated in 25 cm<sup>2</sup> cell culture flasks (Corning) at 600 000 cells/flask in 10 ml media. Flasks were incubated at 37°C for 2.5 h before addition of polyamide stock solution or water (control). Three flasks were treated per condition. Cells were incubated for 3 days prior to analysis. Cell morphology was recorded by phase contrast microscopy with 40× magnification (Nikon Diaphot with attached Nikon D100 camera). Cells were then trypsinized for 7 min at 37°C and recombined with their media. An aliquot was removed to determine cell viability and proliferation via trypan blue exclusion and cell count using a Vi-Cell XR cell viability analyzer (Beckman-Coulter). One hundred images were obtained per flask (three flasks/condition). The remaining cells from each treatment condition were then combined, pelleted, washed with Dulbecco's phosphate-buffered saline (PBS) (Gibco) and fixed at -20°C with 70% EtOH for 35 min. The fixed cells were then pelleted, washed with Dulbecco's PBS, re-pelleted and suspended in a solution containing 0.1% Triton X-100, 0.2 mg/ml RNase A (Sigma) and 0.02 mg/ml propidium iodide (Sigma) in Dulbecco's PBS. Cells were incubated overnight at 4°C in the dark prior to FACS analysis (FACSCalibur; Becton Dickinson). Parameters for FL2 were adjusted to give histograms with G<sub>0</sub>/G<sub>1</sub> peaks centered at 200 for each sample to compensate for differences in cell size.



## Animal experiments

Female BALB/c mice were purchased from The Scripps Research Institute Division of Animal Resources. Mice 8–12 weeks of age were given injections of PBS (control) or 100 nmol polyamide/injection via the tail vein on days 0, 2 and 4, with the exception of **2R-Chl**. Doses of **2R-Chl** were given only on days 0 and 2 due to lethality by day 4. The mice were weighed and their behavior monitored and recorded daily. Experimental protocols were approved by The Scripps Institutional Animal Welfare Committee and were conducted in conformity with institutional guidelines, which are in compliance with national and international laws and policies.

## RESULTS AND DISCUSSION

### Polyamide synthesis

Four hairpin polyamides were prepared by solid phase synthesis (23). The hairpin polyamides differed in the turn substituents utilized: Boc-D-Dab(Fmoc)-OH (**1R**, **1R-Chl**), Boc-Dab(Fmoc)-OH (**1S**, **1S-Chl**), Fmoc-D-Dab(Boc)-OH (**2R**, **2R-Chl**) and Fmoc-Dab(Boc)-OH (**2S**, **2S-Chl**) (Figure 1B). Conjugates were generated by coupling the HBTU-activated carboxylic acid of chlorambucil to the parent polyamide following reverse-phase HPLC of the free amine and purified similarly.

### Plasmid design

The core polyamide structure was constructed to bind the sequence 5'-WWGGWGW-3', where W = A or T. In order to probe the affinity and sequence specificity of the parent polyamides, plasmid pMFST2 was designed with an insert containing a single match site ('M' = 5'-TAGGTGT-3') and two single base pair mismatch sites (Figure 2A). The mismatch sites consist of single G•C to C•G replacements at different locations within each binding site ('A' = 5'-TAGGTCT-3', 'B' = 5'-TAGCTGT-3'). For the purpose of thermal cleavage assays, each binding site was designed with proximal adenines which the chlorambucil moiety would be expected to alkylate (19). In addition to the designed sites, inherent in the plasmid structure is a single base pair mismatch site ('C' = 5'-AGTGGTG-3') for a polyamide binding in reverse (Supplementary Figure S1).

### DNA affinity and sequence specificity

Quantitative DNase I footprint titrations to determine binding site affinities and specificities were conducted for parent polyamides **1R**, **1S**, **2R** and **2S** on the 5'-labeled 280 bp PCR product of plasmid pMFST2 (Figure 2 and Table 1). Previous studies have indicated that attachment of the chlorambucil moiety does not significantly alter the DNA-binding affinity of the parent polyamide (19).

The  $\alpha$ -DABA polyamide **1R** bound the match site, M, with  $K_a = 6.7 \times 10^9 \text{ M}^{-1}$  and it had a 94-fold specificity over single mismatch site A and 26-fold specificity over single mismatch site B (Figure 2B). The enantiomer **1S** bound the match site M with  $K_a = 4.8 \times 10^8 \text{ M}^{-1}$ , a 14-fold decrease relative to **1R**. Binding affinities at the other sites were too

low to be measured and the polyamide was found to coat DNA at high concentrations (Figure 2C).

The  $\gamma$ -DABA polyamides **2R** and **2S** bound DNA in accordance with the expected orientation of the free amine (14). **2R** was shown to bind the match and both single base pair mismatch binding sites (Figure 2D). The affinity of **2R** for site M was  $K_a = 1.6 \times 10^{10} \text{ M}^{-1}$ , with 15- and 6-fold preferences over mismatch sites A and B, respectively. The enantiomer, **2S**, bound the single base pair mismatch reverse site C with  $K_a = 1.2 \times 10^9 \text{ M}^{-1}$ , a 21-fold enhancement over the forward match site M (Figure 2E).

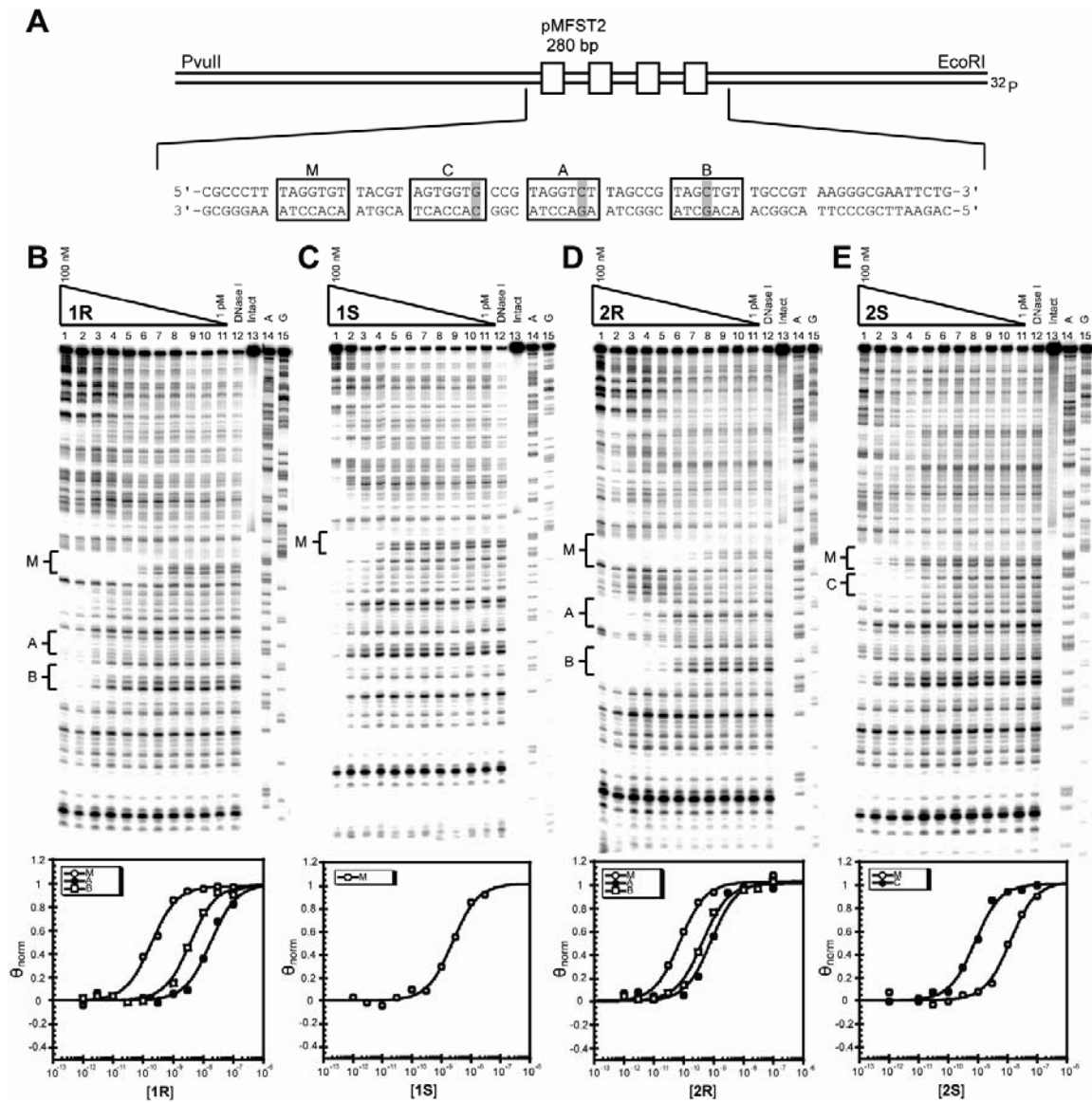
The binding isotherms for all four molecules were best-fit to a 1:1 binding model. Previous studies have found a glycine amino acid-linked polyamide is able to bind DNA in an extended dimeric conformation that can be observed via a larger binding site in a DNase I footprinting gel. We do not observe evidence of an extended conformation for either **1R** or **1S**, as the binding sites of both molecules are the same size as those of **2R** and **2S**. The substituted glycine favors a turn conformation.

The DNase I footprinting experiments reveal that the  $\alpha$ -DABA polyamide **1R** has a 2-fold reduced binding affinity relative to the  $\gamma$ -DABA polyamide **2R**. A contributing factor to the decreased binding observed for **1R** and **1S** could be a reduced ability of the molecules to fit into the minor groove. The shorter turn linkage between the two subunits in an  $\alpha$ -DABA polyamide may reduce its flexibility in conforming to the shape of the groove (Supplementary Figure S2). The energetics of  $\alpha$ -DABA hairpin polyamides appear less sensitive than the  $\gamma$ -DABA series to the *R* and *S* stereochemistry at the turn.

### DNA alkylation specificity and time-dependence

To determine the alkylation properties of the polyamide-chlorambucil conjugates, thermal cleavage assays were performed on the 5'  $^{32}\text{P}$ -labeled 280 bp PCR product of pMFST2 (Figure 3). The alkylation profiles of **1R-Chl** and **2R-Chl** are dramatically different: **1R-Chl** alkylates DNA more specifically than **2R-Chl** and appears less reactive. For the designed sequence, **1R-Chl** appears to specifically alkylate the adenine proximal to the turn at the match site M at concentrations up to 30 nM (Figure 3A). It only begins to alkylate at the designed mismatch binding sites at 100 nM concentration and alkylation can be observed at concentrations as low as 1 nM. In contrast, **2R-Chl** is more promiscuous, alkylating at site M and both mismatch sites, A and B, at concentrations as low as 3 nM (Figure 3B). Only at 1 nM concentration was **2R-Chl** observed to specifically alkylate at site M over sites A and B.

The *S* enantiomers of the conjugates are similar to their *R* analogs in their alkylation specificities, but they are muted in activity. In contrast to **1R-Chl**, **1S-Chl** appears to specifically alkylate the guanine between sites M and C up to a concentration of 30 nM (Figure 3C). Alkylation at N3 of guanine has been observed previously (18,21,25). Based on the DNase I footprinting data for **1S**, the observed alkylation is probably due to binding of **1S-Chl** at the match site M. The **2S-Chl** alkylation profile at 100 nM is similar to that of **2R-Chl** at 10 nM, with the polyamide alkylating at the match site and both single base pair mismatch sites



**Figure 2.** Plasmid design and DNA-binding properties of the parent polyamides. (A) Illustration of the designed portion of the EcoRI/PvuII restriction fragment derived from plasmid pMFST2. The four polyamide binding sites are indicated by boxes. Single base pair mismatches are indicated by shaded regions. Site C is a mismatch site for a polyamide binding in reverse. (B–E) Quantitative DNase I footprint titration experiments for polyamides **1R** (B), **1S** (C), **2R** (D), **2S** (E) on the 280 bp, 5' end-labeled PCR product of plasmid pMFST2: lanes 1–11, 100, 30, 10, 3, 1 nM; 300, 100, 30, 10, 3 and 1 ( $M^{-1}$ ) polyamide, respectively; lane 12, DNase I standard; lane 13, intact DNA; lane 14, A reaction; lane 15, G reaction. Each footprinting gel is accompanied by its respective binding isotherms for the indicated sites (below).

**Table 1.** Binding affinities ( $M^{-1}$ ) for parent polyamides

Polyamide	5'-TAGGTGT-3'	5'-TAGGTCT-3'	5'-TAGCTGT-3'	5'-AGTGGTG-3'
1R	$6.7 (\pm 0.1) \times 10^9$	$7.1 (\pm 1.8) \times 10^7$	$2.6 (\pm 0.2) \times 10^8$	—
1S	$4.8 (\pm 0.9) \times 10^8$	—	—	—
2R	$1.6 (\pm 0.3) \times 10^{10}$	$1.1 (\pm 0.1) \times 10^9$	$2.9 (\pm 1.0) \times 10^9$	—
2S	$5.8 (\pm 2.1) \times 10^7$	—	—	$1.2 (\pm 0.2) \times 10^9$

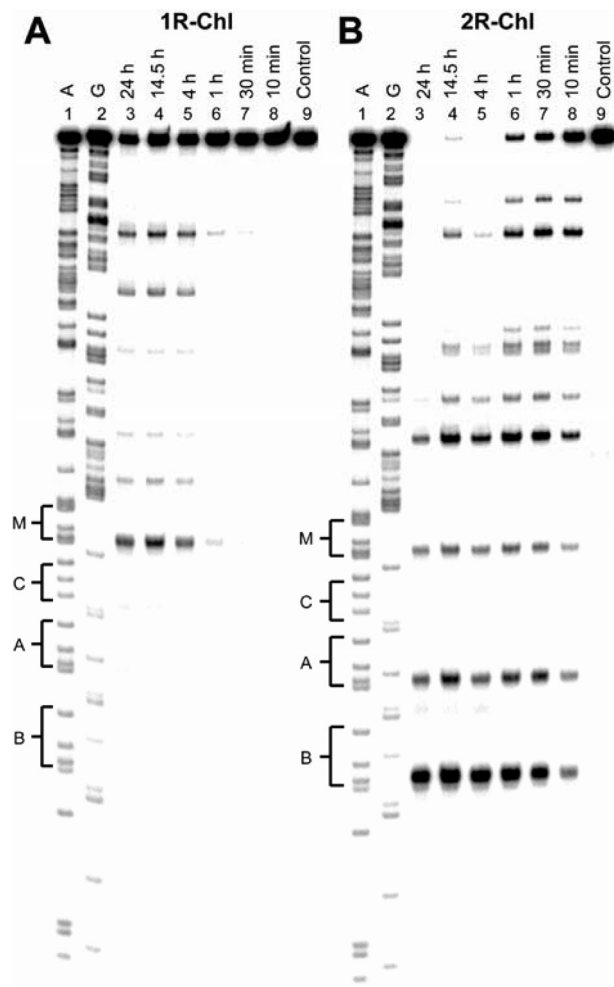
Equilibrium association constants reported are mean values from three DNase I footprint titration experiments; SDs are shown in parentheses.

(Figure 3D). However, at 30 nM, **2S-Chl** is specific for the match site.

Outside of the designed pMFST2 insert, **1R-Chl** and **1S-Chl** alkylate fewer sites than **2R-Chl** and **2S-Chl**,

respectively, further demonstrating their enhanced alkylation specificity. There are a number of forward-binding and reverse-binding single base pair mismatch sites, as well as one reverse-binding match site, inherent in the labeled





**Figure 4.** Time-dependence of alkylation for polyamide conjugates. Thermal cleavage assay experiments with 500 nM **1R-Chl** (A) and 500 nM **2R-Chl** (B) on the 280 bp, 5' end-labeled PCR product of pMFST2: lane 1, A reaction; lane 2, G reaction; lanes 3–8, equilibrations for 24, 14.5, 4, 1 h; 30 and 10 min, respectively; lane 9, intact DNA.

fragment of pMFST2 used in these studies. The major sites of alkylation observed for **1R-Chl** and **2R-Chl** outside of the designed plasmid insert can generally be attributed to forward-binding single base pair mismatch sites, although the gel resolution is often too poor to definitively identify specific alkylation sites. The additional major alkylation sites observed for **2S-Chl** can be attributed to a combination of forward-binding and reverse-binding single base pair mismatch sites, while the additional major alkylation site observed for **1S-Chl** may correspond to a reverse-binding single base pair mismatch site.

We next studied the time-dependence of alkylation for **1R-Chl** and **2R-Chl** to quantitate their kinetic parameters (Figure 4). At 500 nM polyamide and 37°C, the half-life of the labeled DNA (i.e. disappearance of full-length DNA) is 20 h for **1R-Chl** and 10 min for **2R-Chl**. Thus, **2R-Chl** alkylates DNA 120 times faster than **1R-Chl** under these conditions. Given the minimal structural differences between **1R-Chl** and **2R-Chl** and the similar binding affinities of their parent molecules, the dramatic difference in alkylation

specificity and kinetics between hairpins with a common alkylator moiety would not have been predicted.

### Effects of polyamides on SW620 cells

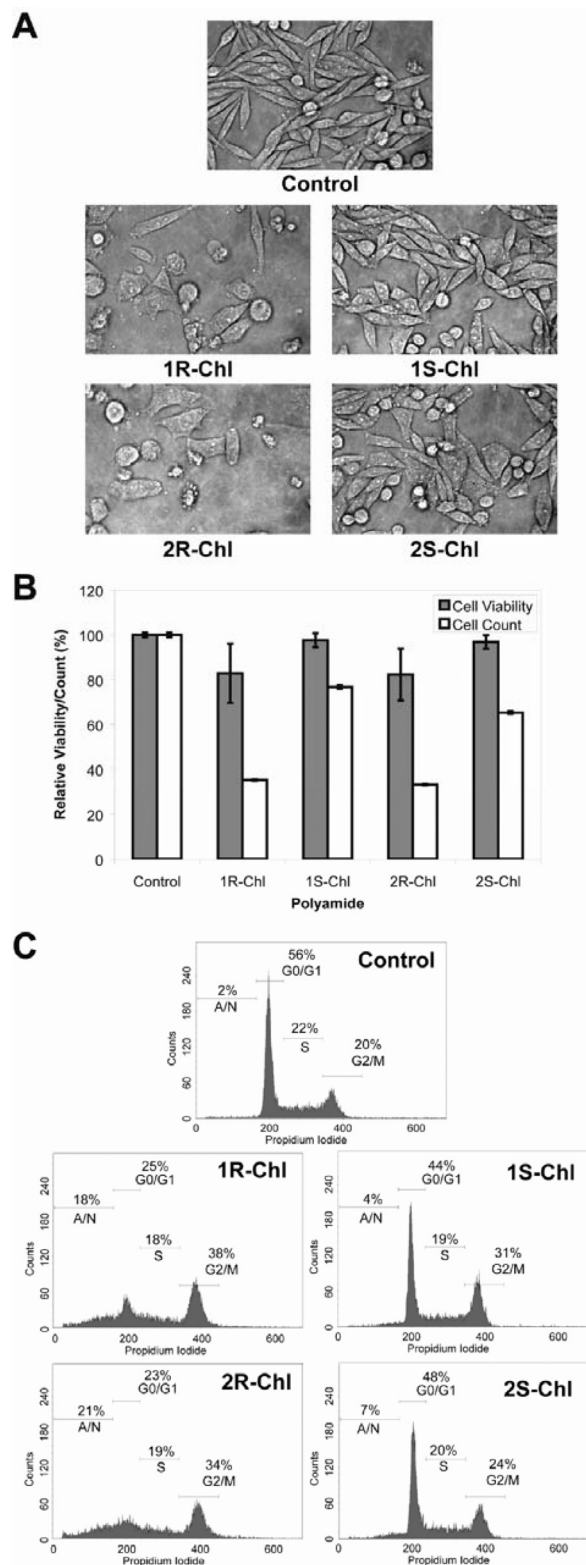
**1R-Chl** treatment of cultured SW620 cells has been shown to induce a morphology change and arrest cell growth in the G<sub>2</sub>/M phase. In addition, the polyamide was found to be a cytostatic agent, decreasing cell proliferation without significantly affecting cell viability, as determined by trypan blue staining (21). A two-hit mechanism for growth arrest by **1R-Chl** was suggested by experiments where a different polyamide-chlorambucil conjugate, which by itself had no effect on SW620 cell proliferation, was found to cause growth arrest after treatment of these cells with an siRNA directed against H4c mRNA (26). Our model for the action of **1R-Chl** involves direct alkylation of the H4c gene, leading to a block in transcription and eventual depletion of H4 protein, opening up the genome for massive alkylation by **1R-Chl** and G<sub>2</sub>/M arrest.

We studied the effects of our polyamide library on SW620 cultured cells to determine whether the differences between the polyamides observed *in vitro* would translate to disparities in cell morphology, proliferation, viability and cell cycle profile. As expected, no effects were observed for the treatment of cells with the four parent polyamides (Supplementary Figure S3). Cells treated for 3 days with **1R-Chl** or **2R-Chl** show the same morphological change as established previously (Figure 5A). Each conjugate also decreases cell proliferation to ~30% relative to control with only a small decrease in cell viability, as measured with trypan blue exclusion staining (Figure 5B). FACS analysis using propidium iodide staining indicates both molecules arrest cells in the G<sub>2</sub>/M phase of growth (38% for **1R-Chl** and 34% for **2R-Chl**, compared to 20% for control; Figure 5C).

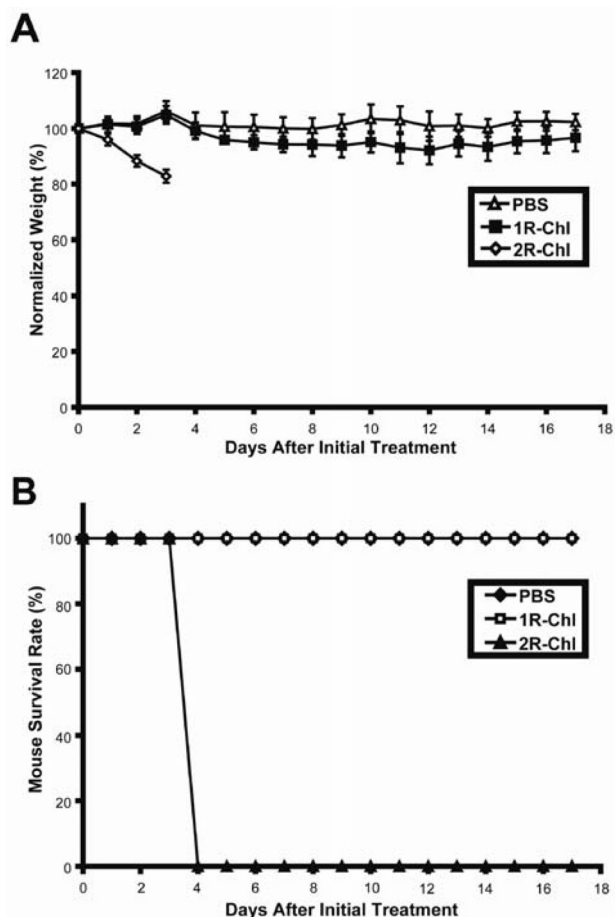
Interestingly, a less pronounced cellular effect is observed for cells treated with **1S-Chl** and **2S-Chl**. Although most cells appear identical to control, some of the cells treated with each of the *S* enantiomers display a morphology change (Figure 5A). Cell viability is not significantly reduced, but proliferation decreases slightly (77% for **1S-Chl** and 65% for **2S-Chl** relative to control; Figure 5B), indicating a slight cytostatic effect for these conjugates. This is also apparent in the FACS data for **1S-Chl**, where 31% of the cells are found to be in the G<sub>2</sub>/M phase of growth. The greater population of apoptotic/necrotic cells following **2S-Chl** treatment may explain why significant growth arrest is not observed with these cells. We find that the trend of similar but muted effects of *S* relative to *R* enantiomers observed in the alkylation data appears to correlate to biological effects on SW620 cells. Previous studies have correlated the DNA interstrand crosslinking ability of alkylating agents with biological activity (27,28). Since the *S* isomers are poor DNA alkylators and have minimal biological activity in SW620 cells, we chose not to investigate them further.

Additional experiments were performed with longer incubation times to determine if cells treated with **1R-Chl** or **2R-Chl** die by apoptosis (Supplementary Figure S4). SW620 cells were treated with 200 nM polyamide for either 3 or 6 days, with the 6-day experiment involving replacement on the third day with fresh media containing 200 nM





**Figure 5.** Effects of polyamide conjugates on cultured cells. Cultured SW620 cells were treated with 200 nM of the indicated polyamide for 3 days prior to analysis. (A) Representative phase microscopy images. (B) Normalized cell viability and proliferation values. Data shown are the averages of three samples; error bars indicate standard deviations. (C) Fluorescence-activated cell-sorting analysis of cells. Plots indicate cell numbers versus propidium iodide staining. Percentages of cells in G<sub>0</sub>/G<sub>1</sub>, S, G<sub>2</sub>/M and apoptotic/necrotic cells (A/N) are indicated.



**Figure 6.** Effects of polyamide conjugates on BALB/c mice. Female BALB/c mice were treated with PBS (control), 100 nmols **1R-Chl**, or 100 nmols **2R-Chl** at 0, 2 and 4 (PBS and **1R-Chl** only) days. Five mice were treated for each experimental condition. (A) Average mouse weights, normalized to weight at day 0, over time. Error bars indicate standard deviations. (B) Survival rates for treated mice.

polyamide. Comparison of the FACS analyses of these cells using annexin V-fluorescein isothiocyanate (annexin V-FITC) and propidium iodide staining shows that after 6 days, **2R-Chl** treatment leads to greater cell death than **1R-Chl** treatment (33.7 versus 20.2%, respectively). However, a significant difference in apoptosis (i.e. cells that are annexin V positive and propidium iodide negative) between treatments with **1R-Chl** and **2R-Chl** is not observed (5.28 versus 6.73%, respectively).

#### *In vivo* effects of polyamides on BALB/c mice

**1R-Chl** has been shown to arrest cancer growth in a SW620 xenograft nude mouse tumor model, whereas **2R-Chl** treatment was toxic to the mice (21). The effects of **1R-Chl** and **2R-Chl** on a normal mouse strain were tested. Groups of five BALB/c mice were injected with PBS (control), 100 nmols **1R-Chl**, or 100 nmols **2R-Chl** every other day for up to 5 days. The mice were observed and weighed each day for up to 17 days (Figure 6A). Mice treated with **1R-Chl** do not differ significantly from the control group. However, **2R-Chl** treatment of mice results in an average weight loss



of 17% after 3 days and is found to be lethal to 100% of the mice after 4 days (Figure 6B).

## CONCLUSION

We have shown that changing the turn unit of a hairpin pyrrole-imidazole polyamide-chlorambucil conjugate from the standard  $\gamma$ -DABA to an  $\alpha$ -DABA can result in dramatically different chemical and biological properties. Although unconjugated parent hairpin polyamides **1R** and **2R** bind DNA with comparable binding affinities, their corresponding chlorambucil conjugates have different alkylation profiles. The polyamide conjugate **1R-Chl**, which contains the (*R*)- $\alpha$ -DABA turn unit, alkylates DNA much more specifically and at a decreased rate relative to **2R-Chl**, which contains the (*R*)- $\gamma$ -DABA turn. Strikingly, treatment of a common mouse strain with **1R-Chl** did not appear to have adverse effects on the mice, whereas treatment with **2R-Chl** was lethal. Although it is possible the difference in toxicities between the two molecules is due to variance in absorption, distribution, metabolism or excretion, given their structural similarities, it is not unreasonable to speculate the origin is the dramatic difference in alkylation specificity and reactivity. These results would not have been predicted a priori and indicate that hairpin polyamides containing the  $\alpha$ -DABA turn unit may be an important class of DNA-binding small molecules with interesting biological properties.

## SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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