

Coarse-Grained Modelling of DNA and DNA self-assembly



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Abstract

In this thesis I present a novel coarse-grained model of deoxyribonucleic acid (DNA). The model represents single-stranded DNA as a chain of rigid nucleotides, and includes potentials to represent chain connectivity, excluded volume, hydrogen-bonding and base stacking interactions. The parameterization of these interactions is justified by comparing the model's representation of a range of physical phenomena to experimental data. In particular, the geometrical structure and elastic moduli of duplex DNA, and the flexibility of single-stranded DNA, are shown to be physically reasonable. Additionally, the thermodynamics of single-stranded stacking, duplex hybridization, hairpin formation and more complex motifs are shown to agree well with experimental data.

The model is optimized for capturing the thermodynamic and mechanical changes associated with duplex formation from single strands. Considerable attention is therefore given to ensuring that single-stranded DNA behaves physically, an approach which differs from previous attempts to model DNA. As a result, the model is the first in which an explicit stacking transition is present in single strands, and also the only coarse-grained model to date to capture both hairpin formation within a single strand and duplex formation between strands.

The scope of the model is demonstrated by simulating DNA tweezers, an iconic nanodevice – the first time that coarse-grained modelling has been applied to dynamic DNA nanotechnology. The simulations suggest that branch migration during toehold-mediated strand displacement – a central feature of many nanomachines – does not have a flat free-energy profile, as is generally assumed.

This finding may help to explain the observed dependence of displacement rate on toehold length.

Finally, the operation of a two-footed DNA walker on a single-stranded DNA track is considered. The model suggests that several aspects of the walker will reduce its efficiency, including a tendency to bind to an undesired site on the track. Several design modifications are suggested to improve the operation of the walker.

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Chapter 1

Introduction

In this thesis I introduce a coarse-grained model of deoxyribonucleic acid (DNA) which is optimized for reproducing the thermodynamic and mechanical changes accompanying the formation of B-DNA duplexes from single strands. This process, known as hybridization, is a vital component of the fast-growing field of DNA nanotechnology, as well as being relevant to a wide range of biological systems.

The layout of this thesis is as follows: in Chapter 1 I will first introduce the DNA molecule, and discuss its relevance in biology and nanotechnology. Then I will consider modelling of DNA, and highlight the need for a new coarse-grained approach. My novel model is presented in Chapter 2, and the techniques used to simulate it are outlined in Chapter 3 (a technical issue with simulation is discussed in Chapter 4). In Chapters 5 and 6, the model is fitted and validated by comparison to extensive experimental data on thermodynamic and mechanical properties of DNA. Finally, to demonstrate the utility of the model, it is applied to two nanodevices (DNA tweezers in Chapter 7 and a two-footed DNA walker in Chapter 8) – in both cases, non-trivial results are observed.

1.1 DNA chemistry and structure

The discovery of the structure of (DNA) and its role in biology was one of the triumphs of 20th century science, revealing the molecular basis of genetics. The existence of DNA was first revealed in 1868/9 by Meischer [1], who discovered a novel substance common to all nuclei that contained large amounts of phosphorus and no sulphur. Levene later

proposed that DNA consisted of nucleotides (base, sugar and phosphate moieties – see Figure 1.1 (a)) linked by covalent bonds between the sugar and phosphate groups [4, 5], but dismissed its potential as an information carrier due to a belief that the bases formed small or repetitive chains. The hereditary significance of DNA was revealed in 1944 when Avery *et al.*, expanding on earlier work by Griffith [6], showed that DNA was responsible for the transfer of traits observed when dead bacteria are mixed with a live population [7].

To understand the mechanism of inheritance, however, it was necessary to find the structure of DNA. Although X-ray diffraction patterns of DNA existed prior to 1950, the elucidation of DNA structure was initially hindered by the existence of two allomorphs of DNA [8], the ‘A’ and ‘B’ forms. This dichotomy was realized by Rosalind Franklin [9], and X-ray data from both forms were combined with chemical knowledge¹ by Watson and Crick, who concluded that DNA was a right handed double helix of nucleotides (Figure 1.1) (b). The two strands are held together by specific hydrogen bonds between adenine (A) and thymine (T), and guanine (G) and cytosine (C) bases, and these base pairs (bp) are stacked on their neighbours. Thus, DNA forms a double helix stabilized by bases in the centre, and with sugar and phosphate groups connecting the bases along the outer edge. It is this complementary pairing of bases (AT and CG) that allows DNA to act as the mechanism of inheritance, as will be discussed in Section 1.2.

Since the work of Watson and Crick, our understanding of DNA structure has grown significantly, but the essential principle of DNA as a double helix of complementary base pairs remains valid. The most common A and B forms are now well characterized and example structures are shown in Figure 1.1 (b). Both are right-handed double helices, but whereas in B-DNA the base pairs lie astride and almost perpendicular to the helix axis, A-DNA base pairs are offset and significantly tilted with respect to the helix axis [8]. Specific repetitive sequences can also form alternative structures, such as the left-handed Z-DNA (Figure 1.1 (b)). B-DNA is the most common in physiological conditions, but A-DNA can

¹Chargaff *et al.* had shown that of the four base types in DNA, the pair adenine and thymine always occur in equal amounts, as do guanine and cytosine [10]. Gulland and coworkers had also suggested that bases were linked by hydrogen-bonding [11].

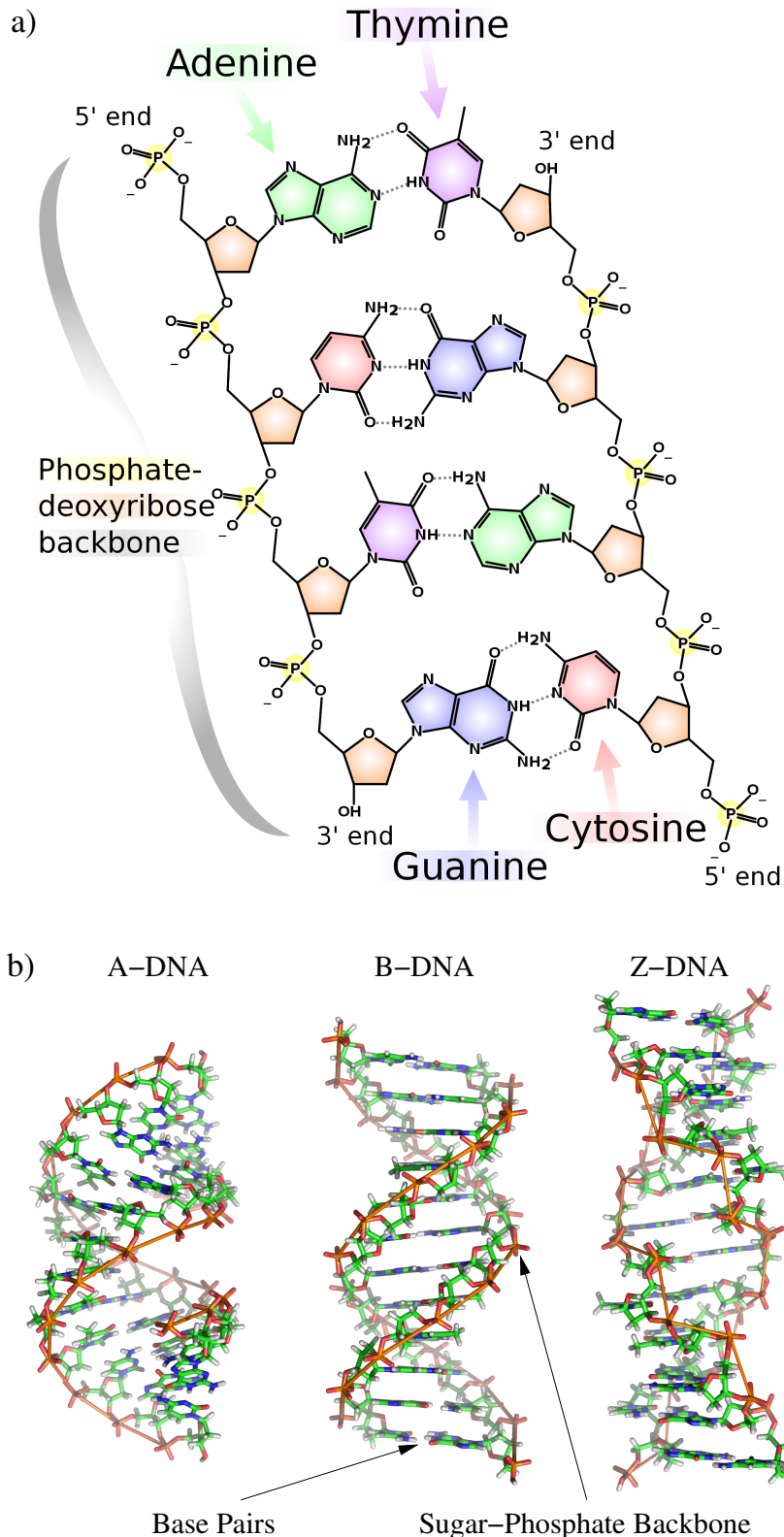


Figure 1.1: DNA chemistry and structure. (a) The chemical composition of DNA: two strands of two nucleotides each are shown in a schematic view (taken from Reference [2]), with covalent bonds shown as solid lines and hydrogen bonds as dashed lines. (b) Helical structures of duplex DNA: the A, B and Z conformers (taken from Reference [3]).

be favoured by the lower humidity in X-ray scattering experiments, and RNA-DNA hybrids also form A-type helices (as do RNA duplexes themselves) [12].

More exotic structures can also be formed through alternative binding mechanisms for certain DNA sequences. Hydrogen bonding with a different side of the base (Hoogsteen hydrogen bonding) allows the formation of ‘G-quadruplex’ structures [8], which occur in nature as the telomeres at the end of a chromosome.

1.2 The role of DNA in biology

Complementary base-pairing is the key property that allows DNA to function as the mechanism of information storage and inheritance. Firstly, the sequence of bases in DNA stores the information required to self-assemble and maintain an organism – the most obvious example is the use of sequence to specify the proteins to be constructed in a cell [13]. The specificity of DNA binding (A-T and G-C base pairs are the most favourable) means that each strand in a double helix carries a negative image of the information on the other strand. As a consequence, it is possible to copy DNA by separating the two strands and using the information on both to construct copies of the original, as is done by DNA polymerase [13].

In order for the information stored in DNA sequences to be useful, the bases must be accessible to enzymes such as RNA polymerase. These enzymes need to interpret the base sequence in order to function – in the case of RNA polymerase, sequences are used to generate messenger RNA molecules, which can trigger the assembly of specific proteins. It is therefore necessary that base-pairing is only marginally stable, so that the helix can be opened and the sequence read.

1.3 DNA nanotechnology

1.3.1 DNA nanostructures

With the advent of the ability to make short DNA sequences to order has come the realization that DNA has ideal properties for use in nanotechnology. A set of single strands can be designed with a pattern of complementarity that specifies a certain 2- or 3-dimensional

structure (usually formed from branched double-helices) as the global free energy minimum of the system. Strands can then be mixed and self-assemble, provided the sequences are well designed. The structural properties of DNA make it ideal for this purpose – double-stranded DNA (dsDNA) is stiff on the nanoscale, with helices having a persistence length of around 50 nm or 150 bp [14]. By contrast, single-stranded DNA (ssDNA) has the flexibility to act as hinges between duplex sections.

The idea of using DNA crystals to facilitate protein crystallography was the original spark that led Nadrian Seeman to found the field of DNA nanotechnology. The self-assembly of short strands (oligonucleotides) was first demonstrated by the Seeman lab, who created a four-armed junction [15]. Junctions of this type, and more complex motifs [16, 17], have been used to create lattices [18, 19, 20] and ribbons [17]. Recently, proteins have been attached to a two-dimensional DNA crystal to facilitate electron cryomicroscopy studies [21]. 3-dimensional structures have also been realized: initially, the Seeman group constructed a cube [22] and a truncated octahedron [23] in several discrete stages. Polyhedral cages that rapidly form as solutions of oligonucleotides are cooled have recently been developed [24, 25, 26, 27, 28]. Self-interactions within a single strand have also been used to create a tetrahedron [29].

An alternative approach to self-assembly, DNA origami, was recently developed by Rothemund [30]. In this case, a long single strand is folded into a desired structure by short “staple” strands, allowing the assembly of an enormous range of 2-dimensional structures. This approach has recently been extended to three dimensions, either by linking together 2-dimensional sheets [31, 32], or by using the twist of DNA to form inherently 3-dimensional folded strands [33]. By designing the staples to form links between helices that are not commensurate with DNA periodicity, strain can be incorporated into origami structures, allowing curved and twisted structures to be created [34]. Recently, complex 3-dimensional curved structures such as spheres and bottles have also been realized [35]

Origami has already been shown to have useful applications. It has become a biophysical breadboard, allowing nanoscale placement of the components of interest, such as the binding sites of a DNA walker [37]. The Dietz lab have also developed origami tools for specific

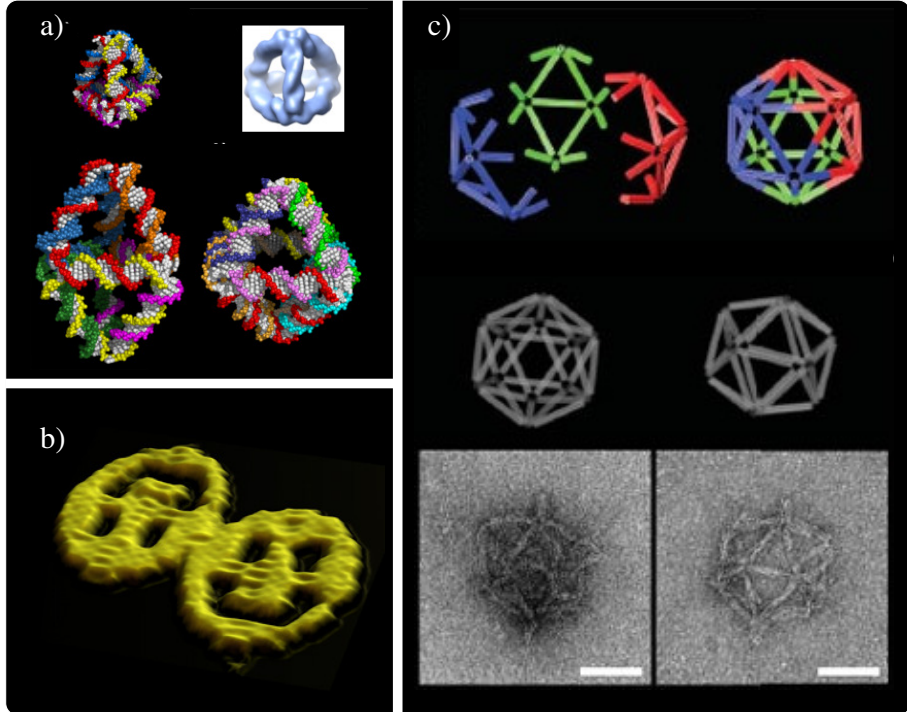


Figure 1.2: Examples of DNA nanostructures. a) Illustrations and a reconstruction from cryoEM data of polyhedra constructed by the Turberfield group [24, 25, 36]. b) AFM images from Rothemund’s original DNA origami work [30]. c) Illustrations and AFM images of icosahedra constructed from origami subunits by the Shih group, taken from Reference [33].³

uses in the laboratory, such as DNA ‘calipers’ for measuring biomolecular dimensions and fluctuations [38]. Liquid crystals of stiff origami rods have also been used to partially align membrane proteins to allow NMR structure determination [39].

The use of origami within more sophisticated structures has also recently been pioneered. Origami tiles have been connected by ‘sticky ends’ (extra bases which can bind to bases on other tiles) to form tubes, 1-dimensional arrays and cages [33, 40, 41]. Some of the edges of an origami consist of blunt-ended helices, which can undergo coaxial stacking interactions with other origami edges to form extended structures, as is evident from the images in Rothemund’s original paper [30]. Recent work has explored the possibility of making such

³Some of the images in this figure were reproduced with permission from the following sources. CryoEM image in a): T. Kato *et al.*. High-resolution structural analysis of a DNA nanostructure by cryoEM. *Nano Letters*, 9(7):2747-2750, 2009. Copyright 2009 American Chemical Society. Bipyramid image in a): C. M. Erben *et al.*. A self-assembled DNA bipyramid. *J. Am. Chem. Soc.*, 129(22):6992-6993, 2007. Copyright 2007 American Chemical Society. c): S. M. Douglas *et al.*. *Nature*, 459:414-418, 2009. Copyright 2009 Macmillan Publishers Ltd:Nature.

interactions selective by introducing patterning to the origami edges [42]. Liedl *et al.* [43] have linked origami (in the form of bundles of DNA double helices) with ssDNA to engineer ‘tensegrity’ structures. In these systems, the three-dimensional conformation of the system is maintained by a balance of tension within the ssDNA sections and compression of the origami bundles.

DNA has also been combined with other materials to create pre-assembled components for self-assembly. Small organic molecules have been used as vertices in structures held together by DNA [44, 45], and colloidal crystallization has been achieved by functionalizing nanoparticles with DNA [46]. Another possibility is to use DNA in combination with ribonucleic acid (RNA) [47], itself a promising material for use in nanotechnology [48].

1.3.2 DNA nanodevices - switches

Marginal stability is thought to be useful for a wide class of assembly processes, as it allows malformed structures to rearrange themselves into the desired configuration [49, 50, 51]. Thermal fluctuations in DNA binding, however, have been explicitly put to use in designing dynamic nanodevices [52]. Two principles are central to much of the work on DNA nanodevices:

- DNA binding can introduce mechanical change to a system, as binding causes strands to be held (reasonably rigidly) in close proximity, and unbinding causes this restriction to be released.
- A strand in a partially-formed duplex with a substrate can be replaced by a strand with a greater degree of complementarity with the substrate [53]. This process is known as *toehold-mediated strand displacement* – see Figure 1.3. Displacement relies on the fluctuational opening of base pairs, so that strands can compete for binding.

The potential for creating nanodevices using these principles was demonstrated by Yurke, Turberfield and coworkers, who constructed the iconic ‘DNA tweezers’ [54], a nanodevice which is the topic of Chapter 7. The tweezers, shown in Figure 1.3, consist of three strands that form two rigid arms with a flexible ssDNA joint. The arms possess overhanging ssDNA

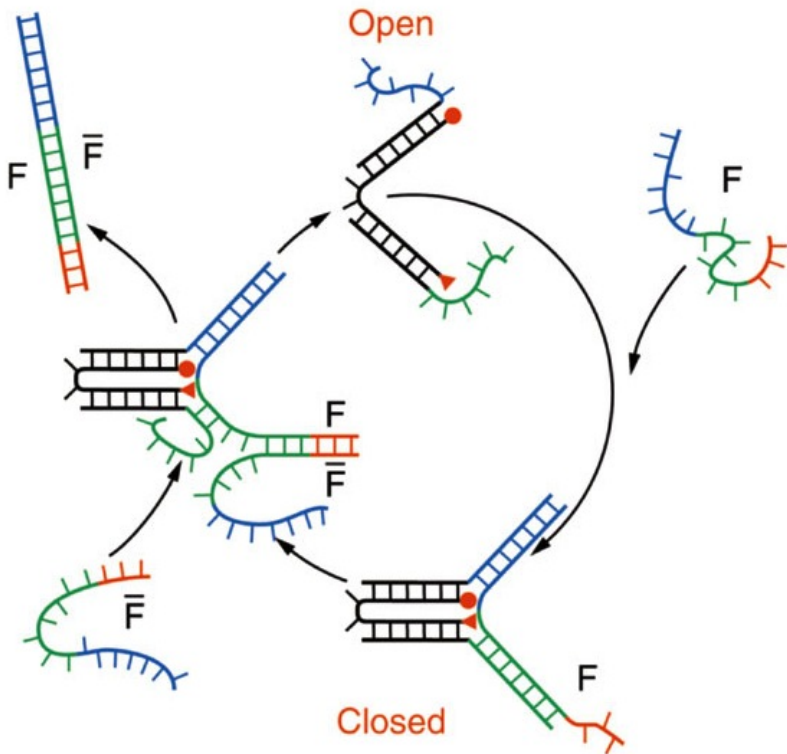


Figure 1.3: DNA tweezers, an example of a DNA nanodevice which relies upon strand displacement. The tweezer unit is initially open. When the fuel (F) is added, it binds to the tweezer arms, bringing them together and closing the tweezers. The antifuel (\bar{F}) is then added and binds to the fuel toehold. The antifuel competes with the tweezers for bonding to the fuel. Finally, the tweezers are displaced by the antifuel, allowing them to open again. Reproduced by permission from Macmillan Publishers Ltd:*Nature*. B. Yurke *et al.*. A DNA-fueled molecular machine made of DNA. *Nature*, 406:605-608, 2000. Copyright 2000.

sections, and a strand of the correct sequence (known as the fuel) can bind to the two arms and pull the tweezers shut. The fuel, however, also possesses an overhanging toehold. Introducing a further strand (the antifuel) which is complementary to the entirety of the fuel leads to the eventual displacement of the tweezers, returning them to their original open state and producing a waste duplex. The tweezers can be thought of as a switch that changes state in response to changes in the environment.

The tweezers themselves do not have an obvious purpose, except perhaps as a method to sense the presence of the fuel strand, but the basic mechanism is at the heart of work to develop potentially useful devices. DNA hybridization and strand displacement have been

used to create boxes [31] and cages [55] that can be opened or closed in response to the presence of certain species of ssDNA in solution (these containers have a size comparable to a small virus). Strand-displacement triggered release of gold nanoparticles from within wires constructed from DNA and small organic molecules has also been demonstrated [56]. Displacement on a much grander scale has also been used to reconfigure a Möbius strip constructed from DNA origami [57], offering a new assembly technique which may assist the construction of topologically complex structures.

DNA strands are not the only signal to which switches can be designed to respond. Douglas *et al.* [58] have designed a DNA origami cage that is locked by two duplexes. One of the strands in these duplexes, however, is an aptamer for a cancer marker, which can displace the other locking strand and lead to the opening of the cage only when bound to a cancerous cell. The system shows promise as a mechanism for the targeted drug delivery.

1.3.3 DNA nanodevices - walkers

An attractive idea is to couple the mechanical changes to directional motion, creating ‘walkers’ inspired by the molecular motors of biology. Typically, walkers have one or two feet which attach to binding sites on a track. The earliest designs used sequential addition of strands to generate coordinated, unidirectional motion (via displacement) [59, 60].

Autonomous, unidirectional motion, which does not rely on external control of the environment (in this case through manipulating the concentrations of fuel strands as a function of time), must catalyze the release of free energy from a fuel source [52]. The free energy release upon the hydrolysis of the phosphodiester backbone of nucleic acids can be used for this purpose [37, 61, 62]. In these systems, the presence of a walker catalyzes the hydrolysis of the single-stranded binding site to which it is attached, thereby encouraging the walker to step to the next site. Unidirectional motion arises as the track behind the walker is modified, making it unfavourable to step backwards.

An alternative source of free energy is in catalyzing DNA hybridization itself [63]. If the fuel strands are designed to form self-complementary hairpins, whose opening is catalyzed by the walker operation, the need for sequential addition of strands can be overcome. This

idea has been used to create a two-footed walker that catalyzes the association of fuel with its track [64].

By coordinating the interaction of the feet of a two-footed walker with its track, the Turberfield group have also demonstrated the possibility of autonomous motion on a track that can be reused. Motors have been designed that catalyze both the hydrolysis and hybridization of fuel strands [65, 66] – the former of which is the subject of Chapter 8.

As an alternative to conventional strand-displacement, the migration of the branch point of a four-armed (Holliday) junction has also been used in the design of walkers [67, 68]. The principle here is very similar to displacement, except that instead of one strand taking base pairs from another, base pairs are transferred between the helices at the branch point. By including an additional toehold, the reaction can be biased in the desired direction.

Walking devices have a clear potential to act as active agents in a molecular assembly line. Initial studies have demonstrated the possibility of using DNA hybridization to accelerate chemical processes through bringing reagents into close proximity [69], and the possibility of using a walker to selectively pick up gold nanoparticle cargo [70].

1.3.4 DNA computation

The ability of DNA to carry information and undergo reactions based on that information has lead to the suggestion that it can be used for computation. In 1994, Adleman demonstrated that a Hamiltonian path problem could be encoded into DNA strands, which could then solve the problem upon being mixed [71]. The novel aspect of DNA computation is the high parallelization – the presence of a thermodynamically large number of strands means that the system can attempt many solutions to a combinatorial problem simultaneously. This potential for high parallelization has lead to a great deal of work aimed at creating architectures for DNA computation that have the potential to solve useful problems [72]. Considerable effort has also been devoted to developing logic gates based on DNA displacement [73].

Although DNA possesses the advantages of parallelism and miniaturization over conventional computing technology, it also possesses several obvious disadvantages. Firstly, the

creation of the strands and analysis of the results are complicated and time-consuming. This problem is made worse by the fact that DNA computers are currently ‘one-shot’ devices: you have to recreate your system every time you want to run a new calculation, and they cannot be programmed easily [72].

Perhaps the best hope for applications of DNA-based decision-making is to use it *in vivo*, where its ability to interface directly with biological matter becomes an advantage, and the difficulty in extracting human-readable output may not be relevant. For example, one might consider a system that could perform a logical calculation based on the environment in a cell, and respond in a potentially therapeutic manner (such as by releasing drugs from containers like those discussed in Section 1.3.2). In this case the massively parallel nature of DNA computation would be used to treat many cells at once, and there would be no need to convert the results into readable output. Indeed, RNA computation has recently been used to trigger selective cell death in response to the presence of a cancer marker [74].

1.4 Modelling DNA self-assembly

1.4.1 Why model DNA self-assembly?

As discussed in Section 1.3, DNA nanotechnology is a rapidly growing field of great potential. Much of DNA nanotechnology relies either largely or entirely upon the formation of B-DNA duplexes from single strands (although other transitions can be exploited, such as the formation of ‘i-motif’ structures [75]).

Currently there is only a limited theoretical understanding of the processes involved in DNA self-assembly, which hampers efforts to design ever more sophisticated systems. In particular, information about the intermediate states in assembly processes, which are often difficult to resolve in experiment yet crucial to the processes as a whole, would aid the design of nanotechnology. Computer modelling, provided it can capture the transition between single- and double-stranded DNA, has the potential to offer significant insight into these systems. For example, a simulation might be able to explain why some systems are more successful than others, or provide an efficient way to test novel ideas.

A model of DNA that captures the transition from ssDNA to dsDNA would therefore be of great use to the DNA nanotechnology community. Furthermore, many systems of biological relevance (such as the opening of transient ‘bubbles’ (stretches of broken bps) within helices and the extrusion of cruciform structures in negatively supercoiled DNA [76]) are governed by the properties of single and double strands, and the competition between the two. A reliable model would also deepen our understanding of such systems.

1.4.2 Atomistic and continuum models of DNA

At the most detailed level, atomistic simulations using force fields such as AMBER or CHARMM offer an intimate representation of DNA [77]. A large-scale systematic study of the structural properties of short sequences as represented by AMBER has been carried out by the Ascona B-DNA Consortium [78]. Unfortunately, the number of degrees of freedom (including those of the solvating H₂O molecules) prohibits the simulation of large molecules for long periods of time. For example, simulations of double helices (on the scale of 10–20 base pairs) have only recently been extended to time scales of $\sim 1 \mu\text{s}$ [79, 80]. The use of enhanced sampling techniques has given atomistic simulations some access to hybridization transitions in the smallest duplexes [81] and hairpins [82, 83], although larger systems remain prohibitively expensive to model.

At the other end of the spectrum, continuum models of DNA [84] treat the double helix as a uniform medium. Whilst these approaches can provide important insight into DNA behaviour on long length-scales, they are by definition unable to deal directly with processes involving duplex hybridization or melting.

It is also worth noting models that have been introduced for the explicit purpose of modelling DNA origami. Sherman and Seeman have presented a geometrical scheme for minimizing strain in origami structures [85], and a finite element method (which treats dsDNA as an elastic rod) has been developed by Castro *et al.* [86] to predict the structure of stressed origami. Although these tools are useful in the nanotechnology design process, they are also inherently inapplicable to the assembly process itself.

1.4.3 Coarse-grained models of DNA

To gain further insight into hybridization, coarse-grained models, which represent DNA through a reduced set of degrees of freedom with effective interactions, are required. Models of DNA with approximately 10 coarse-grained units per nucleotide have been successfully used to study the interaction of DNA with lipids [87, 88], but in order to explore assembly transitions simpler models are required. In particular, models whose coarse-grained scale is approximately that of the nucleotide may provide the ideal compromise between resolution and computational speed for assembly transitions.

Statistical models of DNA

The simplest available coarse-grained models are statistical, neglecting structural and dynamical detail. These models use sequence-dependent parameters that describe the free-energy gain per base pair relative to the denatured state, with extra parameters used for initialization of duplex regions and to describe unpaired sections within the structure. Among the most popular are the Poland-Scheraga [89] and nearest-neighbour models [90, 91], generally used in the context of polynucleotide and oligonucleotide melting, respectively. A particularly important version of the nearest-neighbour model, which has been shown to reproduce experimental melting temperatures of duplexes ranging from 4–16 bp in length with a standard deviation of 2.3 K, was introduced by SantaLucia and Hicks [90, 91]. In this model, the concentrations of oligonucleotides A and B , and their duplex AB , are given by:

$$\frac{[AB]}{[A][B]} = \exp \left(-\beta(\Delta H_{AB} - T\Delta S_{AB}) \right), \quad (1.1)$$

where the constants ΔH_{AB} and ΔS_{AB} are computed by summing contributions from each nearest-neighbour set of two base pairs, together with terms for helix initiation and various structural features, all of which are assumed to be temperature independent. Such a description, in which ΔH_{AB} and ΔS_{AB} are temperature independent, constitutes a ‘two-state’ model. A two-state model essentially neglects the variation in energy within the bound and unbound ensembles, and is equivalent to approximating each as a single state.

Statistical models, although extremely useful, are unable to describe dynamics of systems or the effects that arise from the geometry and topology of DNA, and hence are not complex enough to study many of the processes involved in DNA nanotechnology.

Models of DNA with reduced dimensionality

Alternatives to these purely statistical models have also been proposed. Everaers *et al.* [92] have suggested a lattice model of DNA explicitly designed to unify nearest-neighbour and Poland-Scheraga models, with the added advantage that some structural information is also preserved. Peyrard-Bishop-Dauxois (PBD) models [93] represent base pairs through a continuous 1-dimensional coordinate, allowing dynamical simulations of denaturation bubbles in polynucleotide DNA. An extension of the PBD model to include twist has also made the investigation of torque induced denaturation possible [94, 95]. None of the models discussed, however, provide a sufficiently sophisticated representation of the 3-dimensional structure of DNA to allow the detailed study of the transitions involved in nanotechnology.

Rigid base-pair models

Rigid base-pair models, in which undeformable base pairs are the fundamental unit, have been used to study perturbations to DNA such as those induced by enzymes [96]. By definition, such models cannot represent the transition from single strands to duplexes, and hence are inappropriate for the study of assembly processes. Lankas *et al.* [97] directly compared rigid base-pair and rigid base models that were parameterized to reproduce positional time-series that were generated from atomistic simulations of B-DNA. Interestingly, the authors found that the rigid base models, in which the base pairs are deformable and nucleotides are the essential unit of simulation, generated a more local representation of the interactions than rigid base-pair models did, suggesting that the individual bases are a more appropriate level of description for structural and mechanical properties of B-DNA.

Rigid and stiff base models

To study the processes involved in nucleic acid structure formation, a fully 3-dimensional coarse-grained model, in which individual bases are able to move separately, is required.

Several models in which the base is either represented as a rigid unit, or with stiff internal degrees of freedom, have been proposed in the last decade. These models represent nucleotides by one or more interaction sites, and can be divided into two kinds. Firstly, some modellers parameterize their effective force fields by direct comparison with either atomistic simulations or data from crystal structures. An alternative is to take a more heuristic approach, designing force fields to provide a reasonable description of a range of large-scale properties (such as melting temperatures of helices) when compared to experiment: these two approaches could be described as ‘bottom-up’ and ‘top-down’, respectively.

Bottom-up approaches have been used to study RNA nanostructures [98], the response of DNA minicircles to supercoiling [99, 100, 101], the behaviour of B-DNA over a range of conditions [102], and the properties of the resultant DNA model as a function of parameterization [103]. These models have one [99, 100, 101], three [98, 103] or six [102] sites per nucleotide.

Typically, adjacent sites within a strand are connected by ‘bonded interactions’, which involve bond stretching, angular and dihedral potentials and provide much of the structure of the model. Additional, ‘non-bonded’ interactions represent base-pairing, stacking, excluded volume and in some cases electrostatic interactions (either treated with explicit ions [101] or implicit linearized Poisson-Boltzmann methods [102, 103]). Additional structural information, enabling the specificity of double helix structures, is encoded in potentials which represent hydrogen-bonding. This is done either through terms which depend on the orientation of individual bases [103], by having bases with internal structure [102] or by having hydrogen-bonding interactions depend on the location of several sites neighbouring the bases in question [98, 99, 100, 101]. In some cases, the interactions are specified by the ‘native state’ of a certain system, so that in a given simulation bases can only bind in one way [98, 99, 100, 101].

Many of these models are parameterized using all-atom simulations of DNA by extracting the distribution functions of various degrees of freedom from small simulations, and then fitting CG potentials to reproduce these distributions. Often this is done using ‘Boltzmann inversion’ (whereby potentials of a variable q are taken as $V(q) = -1/\beta \ln W(q)$, with $W(q)$

being the distribution function of q in the original simulation) as an initial approximation [98, 100, 103]. This procedure has also been performed using X-ray crystal structures of DNA as the source of $W(q)$ [99]. Alternatively, Savalyev and Papoian have pioneered the ‘molecular renormalization group’ technique, which is a systematic method for reproducing correlations in the CG model [101].

It is also worth mentioning the application of a similar methodology to the study of DNA binding to the nucleosome [104]. This model involves one interaction site for each amino acid C_α atom and one for each phosphate of DNA, and interactions are parameterized by Boltzmann inversion of atomistic simulations to reproduce fluctuations around the native state.

Although systematically coarse-graining removes some of the arbitrary choices in designing a minimal model, there are drawbacks. Firstly, the resultant force-field will be biased towards the structures with which it was parameterized: in particular, equilibrium duplex structures are often the primary source of information, and hence single-stranded behaviour is not necessarily well reproduced. In some cases, the potential is actually designed only to reproduce fluctuations about a certain structure of interest [98, 104], and in many cases the bonding pattern of the ‘native state’ is required as an input [98, 99, 100, 101, 104], reducing the general applicability of such models. It is worth noting that there has been little use of these models to date to rigorously study systems and effects other than those with which they were parameterized.

Secondly, the transition between ssDNA and dsDNA may be poorly represented: indeed, none of the bottom-up approaches described above have been used to investigate melting transitions in a rigorous way, with the focus being largely on structural properties. Thirdly, ‘representability problems’ [105] mean that careful fitting to distribution functions will not necessarily reproduce thermodynamic properties in a reliable fashion [106]. Finally, it is not yet known how accurate atomistic simulations are in reproducing the duplex hybridization transition – indeed, some authors have commented that their CG potentials give incorrect structural properties due to issues with the atomistic potentials from which they were parameterized [101].

All coarse-grained models represent a compromise, and an appropriate model must be chosen for the investigation at hand. Current examples of bottom-up approaches seem well-suited to studying fluctuations in the vicinity of the equilibrium structure in question. By contrast, top-down approaches appear to lend themselves to the study of larger changes, particularly assembly transitions.

Top down approaches have been used to study RNA folding and unfolding [107, 108, 109, 110, 111]. These methods have variable levels of detail – several have multiple interaction sites per nucleotide, with complex interactions designed to mimic specific effects like stacking and hydrogen-bonding [108, 109, 110]. The model of Ding *et al.* [109] appears to be particularly promising. It has been used to predict with some success a number of structures formed from a single folded RNA with no input except sequence, including systems as large as tRNA (almost 80 nucleotides in length). It should be noted, however, that this description includes an additional (arbitrary) multi-body loop formation term, with the need to parameterize this term effectively reducing the predictive power of the model. A more simple, one site-per-nucleotide approach has also been used to study the folding and unfolding of large RNA motifs [108]. In this case, attractive interactions are introduced between neighbouring nucleotides specifically to reproduce a certain native state, reducing the general applicability of the approach.

Top-down models of DNA have also been suggested, all with multiple interaction sites per nucleotide, and physically motivated potentials such as stacking and hydrogen-bonding [112, 113, 114, 115, 116, 117, 118, 119, 120, 121, 122, 123, 124, 125, 126, 127]. Drukker *et al.* [112] suggested the first fully 3-dimensional, helical, dynamical⁴ model of DNA, and used it to observe denaturation. Several simpler models of DNA in which helicity is neglected were then used to study the thermodynamics of duplex hybridization [125], duplex mediated gelation of colloids [114] and self-complementary hairpin formation [113, 115, 116]. Using an alternative helical model, Niewieczorzał and Cieplak studied the affects of applying tension to a duplex [117].

⁴In this context a dynamical model is one that makes predictions for the kinetics of processes, as opposed to a statistical model which does not directly give kinetic information.

A large body of work has also been based around a model initially proposed by Knotts *et al.* [118]. Similarly to many of the bottom-up approaches, this model involves three sites-per-nucleotide, one each for base, sugar and phosphate groups. The potential includes extensional, angular and dihedral terms associated with the covalent links within each strand, and hydrogen-bonding, intrastrand stacking, excluded volume and a Debye-Hückel electrostatic term between phosphates. In the original work, very little rigorous analysis of the model was performed, but it was used to study the conformations of a ring nanostructure [119]. Later work has studied the melting transition in some detail [120, 121] and hybridization when one of the strands is tethered [126, 127]. It should be noted that the methodology neglects significant issues with inferring bulk properties from small simulations, as discussed in Chapter 4. In this updated work, the model was re-parameterized and a medium-range attractive potential, the origin of which is unclear, was introduced to facilitate duplex formation. A variant of the model with sequence-dependent structural properties has been applied to nucleosome binding [124]. Other authors have attempted to augment the potential with explicit electrostatics [122] and even an explicit representation of the solvent [123]. It is clear from these approaches that replacing a given implicit, effective interaction with a more explicit form is not a trivial process.

For the purposes of this work, the study of complexes involving B-DNA and their formation, a good representation of the structural, mechanical and thermodynamic properties of single strands and B-DNA is required. Previous models have not been optimized for this purpose. In many cases, models can only represent the duplex state [101], or are strongly biased towards a representation of a single native confirmation [98, 99, 100, 102, 104]. Of the models which are designed to capture duplex formation, the majority represent the helicity of DNA in a somewhat unphysical fashion. Physically, the helicity of DNA derives from the tendency of consecutive bases to form coplanar stacks, with an average separation of around 3.4 Å [128], shorter than the equilibrium separation of phosphates of approximately 6.5 Å [129]. As a result, single strands undergo a transition from a largely ordered, helical structure at low temperature to a disordered one at high temperature [12]. This transition has been largely neglected in the past (helicity is usually either absent [113, 114, 115, 116, 125] or

enforced largely through dihedral and angular potentials imposed on the backbone of a single strand [112, 120, 121, 117, 122, 123, 124, 126, 127]), but it has important consequences. In particular, unstacked strands are extremely flexible relative to duplexes, permitting the formation of DNA structures which involve sharply bent single-stranded regions, such as hairpins. It is worth noting that none of the models of duplex formation have been used to study even simple hairpin-forming systems. Furthermore, it has significant consequences for the thermodynamics and kinetics of assembly (the role of stacking in the thermodynamics of duplex formation is discussed in Chapter 6).

The work of Morriss-Andrews *et al.* [103] and Ding *et al.* [109] are exceptions to the previous comments, in that they capture helicity of duplexes whilst permitting single strands to be unstacked and flexible. The key aspect of these models is that stacking and hydrogen-bonding interactions have orientational dependence, meaning that the potentials which maintain the backbone structure need to be less specific for right-handed double helices to form. The model presented in this work will pursue a similar approach.

As well as the structure and flexibility of duplexes and single strands, the thermodynamics of hybridization is an important aspect to capture. Rigorous thermodynamic simulations, in which melting temperatures are compared to experiment, have not been performed on the majority of models. Of those models for which such comparisons have been made, it has either been exclusively for duplexes [120] or hairpins [113, 115]. An additional concern is the temperature ranges over which transitions occur. For complex assembly processes involving several interactions, it is important that the widths of transitions (and not just the melting temperatures) are similar to experiment, so that certain features such as hierarchical assembly are preserved. More generally, transition widths determine the response of melting temperatures to concentration changes and the addition of stabilizing/destabilizing motifs. Where it was considered, the melting transition in previous models was generally significantly wider than experimentally reported [113, 115, 120].

Chapter 2

A Novel DNA Model

2.1 A feasibility study

When this project was started in 2007, very few models were available in the literature. In particular, no thermodynamic simulations of systems involving branched duplexes had been performed. It was therefore necessary to establish whether the aim of simulating nanotechnology with a coarse-grained model was a feasible one.

At the time, the model which had been studied most rigorously and used to simulate the largest systems (involving many particles undergoing DNA-mediated aggregation) was that of Starr and Sciortino [114]. This model represents DNA as an essentially linear molecule which has the potential to form ladder-like duplexes with its complementary strand. I adapted this model and used it to investigate the formation of ‘Holliday Junctions’, branched four-armed junctions involving four strands of DNA [51].

The junctions considered were based on those used by Malo *et al.* [19] to construct a two-dimensional DNA crystal. These junctions, as shown in Figure 2.1, had two 13-bp long arms and two 7-bp long arms. Due to the extra bonding in the longer arms, these were predicted to be stable at higher temperature, and indeed UV absorbance did suggest that the junction forming process proceeded in two stages as the system was cooled.

Rigorous model thermodynamics were obtained for the entire assembly process, demonstrating the possibility of using coarse-grained models to simulate DNA nanotechnology. The model reproduced the greater stability of 13-bp duplexes, and also suggested the possibility of hierarchical assembly at constant temperature (having formed the longer arms,

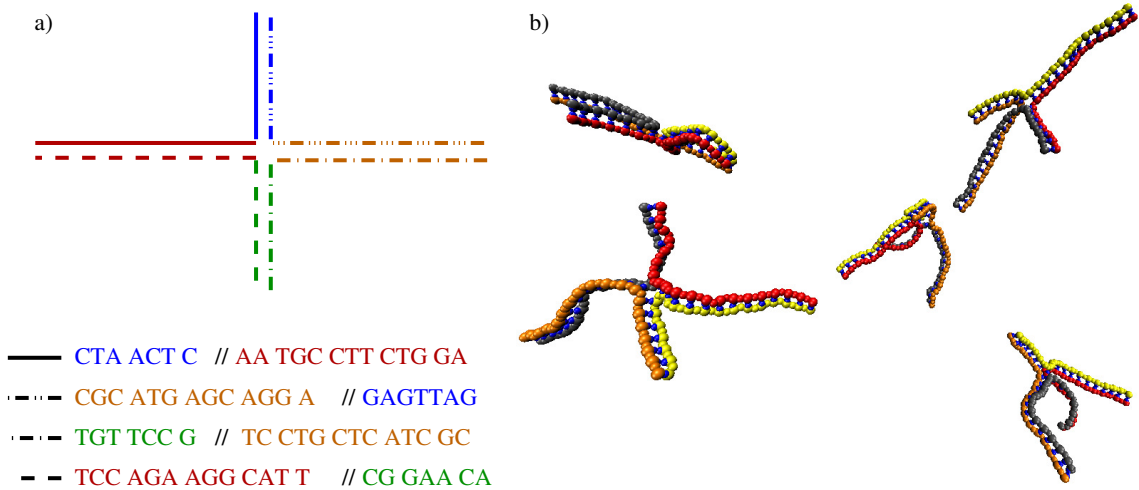


Figure 2.1: a) Strand sequences and schematic Holliday Junction structure from the system simulated in Reference [51]. b) Five identical Holliday junctions as represented by the model of Reference [51].

the two shorter arms of the junction could form cooperatively at temperatures above their individual melting points). Another pleasing result was that displacement of strands from partially bound duplexes was observed, an important process in DNA nanotechnology. As is evident from Figure 2.1 (b), however, there were significant issues with this model. Due to the ladder-like nature of the model and absence of an explicit stacking interaction, the geometrical and mechanical properties of duplex DNA were far from realistic. For example, the duplex arms in Figure 2.1 (b) are unrealistically flexible perpendicular to the plane of bonding.

2.2 The philosophy of the model

Having established that modelling DNA nanotechnology was feasible, it was important to decide which aspects of DNA to emphasize in the new model. For the purposes of simulating much of DNA nanotechnology, I have aimed to embed the thermodynamics of transitions involving ssDNA and dsDNA (in the most common B-form) into a 3-dimensional, dynamical, coarse-grained representation that provides a reasonable description of the structural and mechanical features of the molecule. This ambition naturally coincides with a top-down approach. I have not been primarily concerned with the chemical details of interactions, but rather their net effect with regard to the properties of DNA.

Thermodynamically, the most important transitions to represent are the stacking of single strands, the formation of single-stranded hairpins and the hybridization of two separate strands to form duplexes. In terms of structure, it is vital that a model captures the ability of single strands to be both helically ordered and disordered. The helicity of dsDNA is also crucial, as it has a potentially large role in the kinetics of assembly, in particular leading to frustration of bonding when strands are topologically constrained [130]. A reasonable representation of the mechanical properties of DNA is also necessary. Single strands should be flexible, and duplexes comparatively stiff to represent their roles in nanotechnology. Further, quantities like the torsional and extensional moduli of dsDNA are important if the model is to be used to study systems involving DNA under stress, such as minicircles [131].

I have attempted to capture these properties by using only physically motivated interactions. Pairwise potentials representing excluded volume, backbone connectivity, hydrogen-bonding, stacking, coaxial stacking and cross-stacking have been included (with no terms that possess explicitly length or loop size dependence [109, 120]). Analogously to the work of Morriss-Andrews *et al.* [103] and Ding *et al.* [109], the structure of dsDNA in the model is largely enforced by orientational dependencies of the hydrogen-bonding and stacking potentials.

An additional consideration in model design is the need for computational efficiency (if assembly transitions of complex structures are to be simulated). In the model, all interactions are pairwise (i.e., only involve two nucleotides, which are taken as rigid bodies). This pairwise character allows me to make efficient use of cluster-move Monte Carlo (MC) algorithms [132], which facilitate relaxation on all length-scales in a bound structure, and allow a much larger typical step size than possible in explicitly dynamical simulations (for more information, refer to Chapter 3). Designing the interactions to be truncated at short distances also improves simulation efficiency.

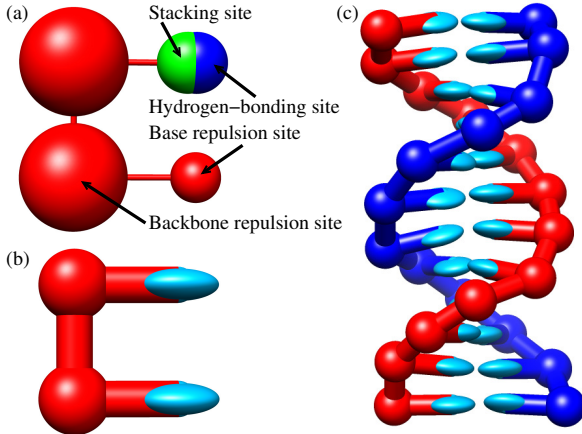


Figure 2.2: (a) Model interaction sites. For clarity, the stacking/hydrogen-bonding sites are shown on one nucleotide and the base excluded volume on the other. The sizes of the spheres correspond to interaction ranges: two repulsive sites interact with a Lennard-Jones σ (see main text) equal to the sum of the radii shown (note that the truncation and smoothing procedure extends the repulsion slightly beyond this distance). The distance at which hydrogen-bonding and stacking interactions are at their most negative is given by the diameter of the spheres. Visualization was found to be clearer with nucleotides depicted as in (b), with the subfigures (a) and (b) representing identical nucleotides on the same scale. The ellipsoidal bases allow a representation of the planarity inherent in the model, with the shortest axis corresponding to the base normal. (c) A 12 bp duplex as represented by the model.

2.3 Degrees of freedom in the model

The model consists of rigid nucleotides with three interaction sites, illustrated in Figure 2.2. The three interaction sites lie in a line, with the base stacking and hydrogen-bonding/base excluded volume sites separated from the backbone excluded volume site by 6.3 Å and 6.8 Å respectively. The orientation of bases is specified by a normal vector, which gives the notional plane of the base: the relative angle of base planes is used to modulate interactions (rather than through the use of off-axis sites).

It must be emphasized that the (often fairly complex) details of the interactions should not be over-interpreted – they are a means to an end. They are an attempt to mathematically quantify the tendency of nucleotides to interact favourably when in the geometry of duplex DNA, through the positions and orientations of the rigid model nucleotides. The widths and well depths of these potentials have been optimized to fit the properties of DNA, under the condition that the formation of unphysical structures was limited.

For the rest of this work, the model will be described in terms of reduced units, where one unit of length corresponds to $l = 8.518\text{ \AA}$ (this value was chosen to give a rise per bp of

approximately 3.4 \AA) and one unit of energy is equal to $E = 4.142 \times 10^{-20} \text{ J}$ (or equivalently, kT at $T = 300 \text{ K}$ corresponds to $0.1E$).

2.4 The potential

2.4.1 Functional forms

The potential used to model DNA consists of a sum of terms designed to represent physical interactions, such as excluded volume, base-stacking and hydrogen bonding. These terms are constructed from the functions given below:

- FENE spring (used to connect backbones):

$$V_{\text{FENE}}(r, \epsilon, r^0, \Delta) = -\frac{\epsilon}{2} \ln \left(1 - \frac{(r - r^0)^2}{\Delta^2} \right). \quad (2.1)$$

- Morse potential (used for stacking and H-bonding):

$$V_{\text{Morse}}(r, \epsilon, r^0, a) = \epsilon \left(1 - \exp(-(r - r^0)/a) \right)^2. \quad (2.2)$$

- Harmonic potential (used for cross-stacking and coaxial stacking):

$$V_{\text{harm}}(r, k, r^0) = \frac{k}{2} (r - r^0)^2. \quad (2.3)$$

- Lennard - Jones potential (used for soft repulsion):

$$V_{\text{LJ}}(r, \epsilon, \sigma) = 4\epsilon \left(\left(\frac{\sigma}{r} \right)^{12} - \left(\frac{\sigma}{r} \right)^6 \right). \quad (2.4)$$

- Quadratic terms (used for modulation):

$$V_{\text{mod}}(\theta, a, \theta^0) = 1 - a(\theta - \theta^0)^2. \quad (2.5)$$

- Quadratic smoothing terms for truncation:

$$V_{\text{smooth}}(x, b, x^c) = b(x^c - x)^2. \quad (2.6)$$

These functional forms are combined to give the following truncated, smooth and differentiable functions:

- The radial part of the stacking and hydrogen-bonding potentials:

$$f_1(r) = \begin{cases} V_{\text{Morse}}(r, \epsilon, r^0, a) - V_{\text{Morse}}(r^c, \epsilon, r^0, a)) & \text{if } r^{low} < r < r^{high}, \\ \epsilon V_{\text{smooth}}(r, b^{low}, r^{c,low}) & \text{if } r^{c,low} < r < r^{low}, \\ \epsilon V_{\text{smooth}}(r, b^{high}, r^{c,high}) & \text{if } r^{high} < r < r^{c,high}, \\ 0 & \text{otherwise.} \end{cases} \quad (2.7)$$

- The radial part of the cross-stacking and coaxial stacking potentials:

$$f_2(r) = \begin{cases} V_{\text{harm}}(r, k, r^0) - V_{\text{harm}}(r^c, k, r^0) & \text{if } r^{low} < r < r^{high}, \\ k V_{\text{smooth}}(r, b^{low}, r^{c,low}) & \text{if } r^{c,low} < r < r^{low}, \\ k V_{\text{smooth}}(r, b^{high}, r^{c,high}) & \text{if } r^{high} < r < r^{c,high}, \\ 0 & \text{otherwise.} \end{cases} \quad (2.8)$$

- The radial part of the excluded volume potential:

$$f_3(r) = \begin{cases} V_{\text{LJ}}(r, \epsilon, \sigma) & \text{if } r < r^*, \\ \epsilon V_{\text{smooth}}(r, b, r^c) & \text{if } r^* < r < r^c, \\ 0 & \text{otherwise.} \end{cases} \quad (2.9)$$

- The angular modulation factor used in stacking, hydrogen-bonding, cross-stacking and coaxial stacking:

$$f_4(\theta) = \begin{cases} V_{\text{mod}}(\theta, a, \theta^0) & \text{if } \theta^0 - \Delta\theta^* < \theta < \theta^0 + \Delta\theta^*, \\ V_{\text{smooth}}(\theta, b, \theta^0 - \Delta\theta^c) & \text{if } \theta^0 - \Delta\theta^c < \theta < \theta^0 - \Delta\theta^*, \\ V_{\text{smooth}}(\theta, b, \theta^0 + \Delta\theta^c) & \text{if } \theta^0 + \Delta\theta^* < \theta < \theta^0 + \Delta\theta^c, \\ 0 & \text{otherwise.} \end{cases} \quad (2.10)$$

- Another modulating term which is used to impose right-handedness (effectively a one-sided modulation):

$$f_5(\phi) = \begin{cases} 1 & \text{if } x > 0, \\ V_{\text{mod}}(x, a, 0) & \text{if } x^* < x < 0, \\ V_{\text{smooth}}(x, b, x^c) & \text{if } x^c < x < x^*, \\ 0 & \text{otherwise.} \end{cases} \quad (2.11)$$

2.4.2 Interactions

The functional forms of Section 2.4.1 are combined to produce a potential for model DNA:

$$\begin{aligned} V = & \sum_{\text{nn}} (V_{\text{backbone}} + V_{\text{stack}} + V'_{\text{exc}}) \\ & + \sum_{\text{other pairs}} (V_{\text{HB}} + V_{\text{cross_stack}} + V_{\text{coaxial_stack}} + V_{\text{exc}}). \end{aligned} \quad (2.12)$$

Here the sum over nn runs over consecutive bases within strands. For illustrations of the degrees of freedom to which the potential is applied, refer to Figure 2.3.

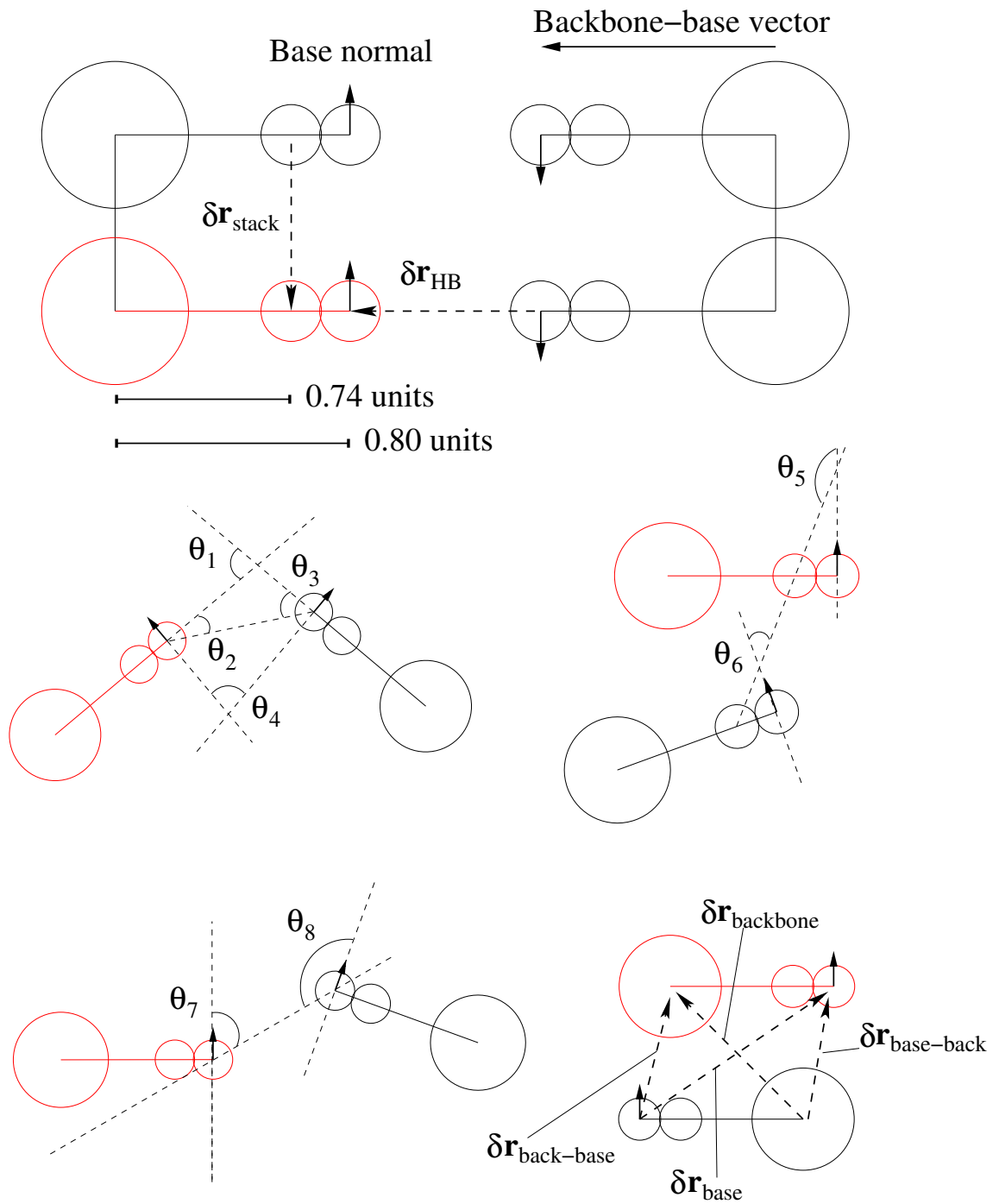


Figure 2.3: Illustration of the variables used to parameterize the potential. Not shown in this diagram are the chirality inducing terms $\cos(\phi_1)$, $\cos(\phi_2)$, $\cos(\phi_3)$ and $\cos(\phi_4)$, which are discussed in detail in Sections 2.4.2 and 2.4.2. It is convenient to define each pairwise interaction as if one nucleotide (coloured red in this picture) is being influenced by the other (coloured black): this allows each angle to be well-defined when calculating forces and torques. When calculating the energy, of course, the final result does not depend on the labelling of nucleotides. I define θ_2 , θ_5 and θ_7 as being measured with respect to the orientation of the red nucleotide, and θ_3 , θ_6 and θ_8 with respect to the orientation of the other nucleotide.

Backbone springs

Consecutive backbone sites on the same strand are connected by finitely-extensible nonlinear elastic (FENE) springs, which maintain backbone connectivity and specify the equilibrium separation of bases along the backbone ($\delta r_{\text{backbone}}^0$). As bases are linked by several covalent bonds in physical DNA, the extensibility of the backbone represents the possibility of rotating these covalent bonds with respect to each other, as well as the extension of the bonds themselves. Here, δr_X is the separation of interaction sites X on two nucleotides.

$$V_{\text{backbone}} = V_{\text{FENE}}(\delta r_{\text{backbone}}, \epsilon_{\text{backbone}}, \delta r_{\text{backbone}}^0, \Delta_{\text{backbone}}). \quad (2.13)$$

Excluded volume

In order to prevent the collapse of model DNA into a dense, strongly-interacting cluster it is necessary to include excluded volume terms (which also prevent the backbones of two strands crossing during dynamical simulations). Excluded volume interactions occur between all combinations of backbone and base excluded volume sites on any two nucleotides (see Figure 2.2). The only exception is for nearest neighbours, for which the backbone excluded volume sites do not interact (as their separation is controlled by the backbone FENE spring).

$$\begin{aligned} V_{\text{exc}} &= f_3(\delta r_{\text{backbone}}, \epsilon_{\text{exc}}, \sigma_{\text{backbone}}, \delta r_{\text{backbone}}^*) + f_3(\delta r_{\text{base}}, \epsilon_{\text{exc}}, \sigma_{\text{base}}, \delta r_{\text{base}}^*) \\ &\quad + f_3(\delta r_{\text{back-back}}, \epsilon_{\text{exc}}, \sigma_{\text{back-back}}, \delta r_{\text{back-back}}^*) \\ &\quad + f_3(\delta r_{\text{base-back}}, \epsilon_{\text{exc}}, \sigma_{\text{back-back}}, \delta r_{\text{back-back}}^*). \end{aligned} \quad (2.14)$$

$$\begin{aligned} V'_{\text{exc}} &= f_3(\delta r_{\text{base}}, \epsilon_{\text{exc}}, \sigma_{\text{base}}, \delta r_{\text{base}}^*) \\ &\quad + f_3(\delta r_{\text{back-back}}, \epsilon_{\text{exc}}, \sigma_{\text{back-back}}, \delta r_{\text{back-back}}^*) \\ &\quad + f_3(\delta r_{\text{base-back}}, \epsilon_{\text{exc}}, \sigma_{\text{back-back}}, \delta r_{\text{back-back}}^*). \end{aligned} \quad (2.15)$$

Stacking

Bases have a tendency to form coplanar stacks, which drives the formation of helices in single- and double-stranded DNA due to the difference in length between the separation of bases along the backbone, and the optimal distance of stacking. The model reproduces this tendency with a stacking interaction between nearest neighbours on the same strand. The potential contains a radial term on the distance between stacking sites (r_{stack}), modulated

by angular terms that favour the alignment of base normals with each other (θ_4) and with the separation between stacking sites ($\theta_{5'}$, $\theta_{6'}$). Right-handed helices are imposed through additional modulating factors which reduce the interaction to zero for increasing amounts of left-handed twist (measured by the quantities $\cos(\phi_1)$ and $\cos(\phi_2)$).

$$\begin{aligned} V_{\text{stack}} = & f_1(\delta r_{\text{stack}}, \epsilon_{\text{stack}}, a_{\text{stack}}, \delta r_{\text{stack}}^0, \delta r_{\text{stack}}^{c,\text{low}}, \delta r_{\text{stack}}^{c,\text{high}}, \delta r_{\text{stack}}^{\text{low}}, \delta r_{\text{stack}}^{\text{high}}) \\ & \times f_4(\theta_4, a_{\text{stack},4}, \theta_{\text{stack},4}^0, \Delta\theta_{\text{stack},4}^*) \\ & \times f_4(\theta_{5'}, a_{\text{stack},5}, \theta_{\text{stack},5}^0, \Delta\theta_{\text{stack},5}^*) f_4(\theta_{6'}, a_{\text{stack},6}, \theta_{\text{stack},6}^0, \Delta\theta_{\text{stack},6}^*) \\ & \times f_5(\cos(\phi_1), a_{\text{stack},1}, \cos(\phi_1)_{\text{stack}}^*) f_5(\cos(\phi_2), a_{\text{stack},2}, \cos(\phi_2)_{\text{stack}}^*). \end{aligned} \quad (2.16)$$

See errata at end of document.

When calculating an interaction, it is convenient to arbitrarily label one of the nucleotides as being influenced by the other, as the angles can then all be defined with respect to one or other of the nucleotides (see Figure 2.3). This is helpful when calculating the torques on a nucleotide, as in this situation one must consider the forces and torques on one of the nucleotides due to the other (and vice versa). If the labelling is swapped, there is of course no difference in the overall energy, and in most cases the interaction is symmetric under exchange of nucleotides so the calculation is also unaffected.

The stacking term is unusual, in that it is not symmetric under the exchange of the two nucleotides in question. $\theta_{5'}$ and $\theta_{6'}$ are calculated differently depending on which nucleotide is chosen to be influenced by the other, allowing the interaction to distinguish between the 3' and 5' directions.

- If the labelled nucleotide is in the 5' direction, $\theta_{5'} = \pi - \theta_5$ and $\theta_{6'} = \theta_6$.
- If the labelled nucleotide is in the 3' direction, $\theta_{5'} = \theta_5$ and $\theta_{6'} = \pi - \theta_6$.

By defining $\theta_{5'}$ and $\theta_{6'}$ in this way, it is possible to require that stacking can only occur when both normals point in the 5' direction. The base normals then define an axis about which the handedness of twist can be measured, and allow anti-parallel base-pairing to be enforced.

The chirality of DNA is introduced into the model via the terms $\cos(\phi_1)$ and $\cos(\phi_2)$ in the stacking interaction, which also distinguish between the 3' and 5' directions. Assume one of the nucleotides has been chosen to be labelled – the vector $\delta\hat{\mathbf{r}}_{\text{backbone}}$ is then the normalized vector **to** this nucleotide **from** the other one. Let \mathbf{y} and $\tilde{\mathbf{y}}$ by the unit vectors

defined by the cross product of the normal and backbone-base vectors of the labelled and unlabelled nucleotides respectively.

- Labelled nucleotide in the 5' direction: $\cos(\phi_1) = \mathbf{y} \cdot \delta\hat{\mathbf{r}}_{\text{backbone}}$, $\cos(\phi_2) = \tilde{\mathbf{y}} \cdot \delta\hat{\mathbf{r}}_{\text{backbone}}$.
- Labelled nucleotide in the 3' direction, $\cos(\phi_1) = -\mathbf{y} \cdot \delta\hat{\mathbf{r}}_{\text{backbone}}$, $\cos(\phi_2) = -\tilde{\mathbf{y}} \cdot \delta\hat{\mathbf{r}}_{\text{backbone}}$.

When the stack forms in a right-handed fashion, $\cos(\phi_1)$ and $\cos(\phi_2)$ will be negative (this result relies on the fact that stacking normals must point in the 3' to 5' direction).

Hydrogen bonding

DNA bases can undergo hydrogen bonding with each other, most commonly along the ‘Watson-Crick’ faces (the edge of the base furthest from the sugar group), with the planes of the bases approximately antiparallel. When taken in conjunction with stacking, the result is the famous B-DNA double helix. The model reproduces base-pairing through the V_{HB} term, which incorporates a radial term dependent on the separation of hydrogen-bonding sites, δr_{HB} . This interaction is modulated by terms that encourage the co-linear alignment of all four backbone and hydrogen-bonding sites (quantified by the angles θ_1 , θ_2 , and θ_3). A further factor is included to encourage the planes of bases to be antiparallel (measured by θ_4). Finally, terms are included that penalize pairs in which the separation of bonding sites is far from orthogonal with the base normals (these angles are θ_7 and θ_8). This final term tends to have only a small role for correctly formed base pairs, but is important in minimizing hydrogen bonding between bases that are not opposite each other in the helix.

$$\begin{aligned} V_{\text{HB}} = & f_1(\delta r_{\text{HB}}, \epsilon_{\text{NB}}, a_{\text{HB}}, \delta r_{\text{HB}}^0, \delta r_{\text{HB}}^{c,\text{low}}, \delta r_{\text{HB}}^{c,\text{high}}, \delta r_{\text{HB}}^{\text{low}}, \delta r_{\text{HB}}^{\text{high}}) \\ & \times f_4(\theta_1, a_{\text{HB},1}, \theta_{\text{HB},1}^0, \Delta\theta_{\text{HB},1}^*) f_4(\theta_2, a_{\text{HB},2}, \theta_{\text{HB},2}^0, \Delta\theta_{\text{HB},2}^*) \\ & \times f_4(\theta_3, a_{\text{HB},3}, \theta_{\text{HB},3}^0, \Delta\theta_{\text{HB},3}^*) f_4(\theta_4, a_{\text{HB},4}, \theta_{\text{HB},4}^0, \Delta\theta_{\text{HB},4}^*) \\ & \times f_4(\theta_7, a_{\text{HB},7}, \theta_{\text{HB},7}^0, \Delta\theta_{\text{HB},7}^*) f_4(\theta_8, a_{\text{HB},8}, \theta_{\text{HB},8}^0, \Delta\theta_{\text{HB},8}^*). \end{aligned} \quad (2.17)$$

Cross-stacking

$V_{\text{cross_stack}}$ represents cross-stacking interactions between a base in a base pair and nearest-neighbour bases on the opposite strand, providing additional stabilization of the duplex [133, 134]. Here it is incorporated through a potential which is a function of the distance between

hydrogen-bonding sites r_{HB} , modulated by the alignment of base normals and backbone-base vectors with the separation vector (the same angles that appear in the definition of the hydrogen bonding potential) in such a way that its minimum is approximately consistent with the structure of model duplexes.

$$\begin{aligned}
 V_{\text{cross_stack}} = & f_2(\delta r_{\text{HB}}, k_{\text{cross}}, \delta r_{\text{cross}}^0, \delta r_{\text{cross}}^{c,low}, \delta r_{\text{cross}}^{c,high}, \delta r_{\text{cross}}^{low}, \delta r_{\text{cross}}^{high}) f_4(\theta_1, a_{\text{cross},1}, \theta_{\text{cross},1}^0, \Delta\theta_{\text{cross},1}^*) \\
 & \times f_4(\theta_2, a_{\text{cross},2}, \theta_{\text{cross},2}^0, \Delta\theta_{\text{cross},2}^*) f_4(\theta_3, a_{\text{cross},3}, \theta_{\text{cross},3}^0, \Delta\theta_{\text{cross},3}^*) \\
 & \times (f_4(\theta_4, a_{\text{cross},4}, \theta_{\text{cross},4}^0, \Delta\theta_{\text{cross},4}^*) + f_4(\pi - \theta_4, a_{\text{cross},4}, \theta_{\text{cross},4}^0, \Delta\theta_{\text{cross},4}^*)) \\
 & \times (f_4(\theta_7, a_{\text{cross},7}, \theta_{\text{cross},7}^0, \Delta\theta_{\text{cross},7}^*) + f_4(\pi - \theta_7, a_{\text{cross},7}, \theta_{\text{cross},7}^0, \Delta\theta_{\text{cross},7}^*)) \\
 & \times (f_4(\theta_8, a_{\text{cross},8}, \theta_{\text{cross},8}^0, \Delta\theta_{\text{cross},8}^*) + f_4(\pi - \theta_8, a_{\text{cross},8}, \theta_{\text{cross},8}^0, \Delta\theta_{\text{cross},8}^*)).
 \end{aligned} \tag{2.18}$$

Coaxial stacking

The final term, $V_{\text{coax_stack}}$, is introduced to capture the tendency of stacking across nicked backbones to stabilize the binding of oligomers to duplexes with overhanging single-stranded tails [91, 135, 136, 137, 138, 139] (coaxial stacking between blunt helix ends is also known to cause origami structures to associate [42]). The interaction is designed to be very similar in form to conventional stacking, with minor differences. Firstly, the radial component of the potential is taken to be of a more truncated, quadratic form – this was to prevent interactions between two bases which were stacked above and below their mutual neighbour. Secondly, it is impossible to define a 3' – 5' axis with two non-neighbouring helices: hence it was necessary to make the potential symmetric with respect to the alignment of base normals with their separation (θ_5 and θ_6). Similarly, without a 3' to 5' axis, the modulating terms which impose right-handedness also have to be designed differently – the new quantities $\cos(\phi_3)$ and $\cos(\phi_4)$ are defined below. Finally, in the case of nearest-neighbour stacking, the restriction of the backbone link between nucleotides means that the modulations described in Section 2.4.2 are enough to describe the geometry of stacking. For non-neighbour nucleotides, however, it was found that configurations in which the nucleotides were poorly stacked also gave significant interactions. To overcome this, an additional modulation in the angle

between the two nucleotides' backbone–base vectors (θ_1) was also included.

$$\begin{aligned}
V_{\text{coax_stack}} = & f_2(\delta r_{\text{stack}}, k, \delta r_{\text{coax}}^0, \delta r_{\text{coax}}^{c,\text{low}}, \delta r_{\text{coax}}^{c,\text{high}}, \delta r_{\text{coax}}^{\text{low}}, \delta r_{\text{coax}}^{\text{high}}) f_4(\theta_4, a_{\text{coax},4}, \theta_{\text{coax},4}^0, \Delta\theta_{\text{coax},4}^*) \\
\times & (f_4(\theta_1, a_{\text{coax},1}, \theta_{\text{coax},1}^0, \Delta\theta_{\text{coax},1}^*) + f_4(2\pi - \theta_1, a_{\text{coax},1}, \theta_{\text{coax},1}^0, \Delta\theta_{\text{coax},1}^*)) \\
\times & (f_4(\theta_5, a_{\text{coax},5}, \theta_{\text{coax},5}^0, \Delta\theta_{\text{coax},5}^*) + f_4(\pi - \theta_5, a_{\text{coax},5}, \theta_{\text{coax},5}^0, \Delta\theta_{\text{coax},5}^*)) \\
\times & (f_4(\theta_6, a_{\text{coax},6}, \theta_{\text{coax},6}^0, \Delta\theta_{\text{coax},6}^*) + f_4(\pi - \theta_6, a_{\text{coax},6}, \theta_{\text{coax},6}^0, \Delta\theta_{\text{coax},6}^*)) \\
\times & f_5(\cos(\phi_3), a_{\text{coax},3'}, \cos(\phi_3)_{\text{coax}}^*) f_5(\cos(\phi_4), a_{\text{coax},4'}, \cos(\phi_4)_{\text{coax}}^*).
\end{aligned} \tag{2.19}$$

Chirality is also imposed for the coaxial stacking term. Here there is no natural 3' to 5' direction, and so the vector between stacking sites rather than base normals must be used. I define the vector $\delta\hat{\mathbf{r}}_{\text{stack}}$ to be the normalized vector **to** the labelled nucleotide's stacking site **from** the other nucleotide, with a similar definition for $\delta\hat{\mathbf{r}}_{\text{backbone}}$. Taking \mathbf{b} and $\tilde{\mathbf{b}}$ to be the backbone-base vectors for the labelled and unlabelled nucleotides respectively:

- $\cos(\phi_3) = \delta\hat{\mathbf{r}}_{\text{stack}} \cdot (\delta\hat{\mathbf{r}}_{\text{backbone}} \times \mathbf{b})$.
- $\cos(\phi_4) = \delta\hat{\mathbf{r}}_{\text{stack}} \cdot (\delta\hat{\mathbf{r}}_{\text{backbone}} \times \tilde{\mathbf{b}})$.

In this case, $\cos(\phi_3)$ and $\cos(\phi_4)$ are both positive for a right-handed stack.¹

2.4.3 Parameterization

The previous section is summarized in Tables 2.1 and 2.2, where the values of parameters of the model are also given. Note that the hydrogen bonding interaction is taken as non-zero only for complementary base pairs A–T and G–C, but beyond this there is no sequence dependence in the model. Also note that the factors b and x^c used in the quadratic smoothing parts of each interaction are not explicitly given. For each function, these quantities are specified by demanding differentiability and continuity at the point where the form of the potential changes. For example, consider an angular modulation $f_4(\theta)$. To be continuous at $\theta^0 \pm \Delta\theta^*$, we require:

$$1 - a(\Delta\theta^*)^2 = b(\theta^c - (\theta^0 \pm \Delta\theta^*))^2, \tag{2.20}$$

and

$$2a(\Delta\theta^*) = 2b(\theta^c - (\theta^0 \pm \Delta\theta^*)). \tag{2.21}$$

¹In fact, after the model was implemented it was found that $\cos(\phi_3) = \cos(\phi_4)$, and so the modulation could be simply calculated as $\cos^2(\phi_3)$.

Interaction		Parameters		
V_{backbone}	$V_{\text{FENE}}(\delta r_{\text{backbone}})$	$\epsilon_{\text{backbone}} = 2$	$\Delta_{\text{backbone}} = 0.25$	$\delta r_{\text{backbone}}^0 = 0.7525$
V_{HB}	$f_1(\delta r_{\text{HB}})$	$\epsilon_{\text{HB}} = 1.077$ $\delta r_{\text{HB}}^c = 0.75$	$a_{\text{HB}} = 8$ $\delta r_{\text{HB}}^{low} = 0.34$	$\delta r_{\text{HB}}^0 = 0.4$ $\delta r_{\text{HB}}^{high} = 0.70$
	$f_4(\theta_1)$	$a_{\text{HB},1} = 1.50$	$\theta_{\text{HB},1}^0 = 0$	$\Delta\theta_{\text{HB},1}^* = 0.70$
	$f_4(\theta_2)$	$a_{\text{HB},2} = 1.50$	$\theta_{\text{HB},2}^0 = 0$	$\Delta\theta_{\text{HB},2}^* = 0.70$
	$f_4(\theta_3)$	$a_{\text{HB},3} = 1.50$	$\theta_{\text{HB},3}^0 = 0$	$\Delta\theta_{\text{HB},3}^* = 0.70$
	$f_4(\theta_4)$	$a_{\text{HB},4} = 0.46$	$\theta_{\text{HB},4}^0 = \pi$	$\Delta\theta_{\text{HB},4}^* = 0.70$
	$f_4(\theta_7)$	$a_{\text{HB},7} = 4.00$	$\theta_{\text{HB},7}^0 = \pi/2$	$\Delta\theta_{\text{HB},7}^* = 0.45$
	$f_4(\theta_8)$	$a_{\text{HB},8} = 4.00$	$\theta_{\text{HB},8}^0 = \pi/2$	$\Delta\theta_{\text{HB},8}^* = 0.45$
V_{stack}	$f_1(\delta r_{\text{stack}})$	$\epsilon_{\text{stack}} = 1.3448$ $+ 2.6568 kT$ $\delta r_{\text{stack}}^c = 0.9$	$a_{\text{stack}} = 6$ $\delta r_{\text{stack}}^{low} = 0.32$	$\delta r_{\text{stack}}^0 = 0.4$ $\delta r_{\text{stack}}^{high} = 0.75$
	$f_4(\theta_4)$	$a_{\text{stack},4} = 1.30$	$\theta_{\text{stack},4}^0 = 0$	$\Delta\theta_{\text{stack},4}^* = 0.8$
	$f_4(\theta_5')$	$a_{\text{stack},5} = 0.90$	$\theta_{\text{stack},5}^0 = 0$	$\Delta\theta_{\text{stack},5}^* = 0.95$
	$f_4(\theta_6')$	$a_{\text{stack},6} = 0.90$	$\theta_{\text{stack},6}^0 = 0$	$\Delta\theta_{\text{stack},6}^* = 0.95$
	$f_5(\cos(\phi_1))$	$a_{\text{stack},1} = 2.00$	$\cos(\phi_1)_{\text{stack}}^* = 0.65$	
	$f_5(\cos(\phi_2))$	$a_{\text{stack},2} = 2.00$	$\cos(\phi_2)_{\text{stack}}^* = 0.65$	
V_{exc}	$f_3(\delta r_{\text{backbone}})$ $+ f_3(\delta r_{\text{base}})$ $+ f_3(\delta r_{\text{back-base}})$ $+ f_3(\delta r_{\text{base-back}})$	$\epsilon_{\text{exc}} = 2.00$ $\epsilon_{\text{exc}} = 2.00$ $\epsilon_{\text{exc}} = 2.00$ $\epsilon_{\text{exc}} = 2.00$	$\sigma_{\text{backbone}} = 0.70$ $\sigma_{\text{base}} = 0.33$ $\sigma_{\text{back-base}} = 0.515$ $\sigma_{\text{back-base}} = 0.515$	$\delta r_{\text{backbone}}^* = 0.675$ $\delta r_{\text{base}}^* = 0.32$ $\delta r_{\text{back-base}}^* = 0.50$ $\delta r_{\text{base-back}}^* = 0.50$

Table 2.1: Parameter values in the model. In this table, all energies and lengths are in terms of the simulation units E and l . When more than one function is listed for an interaction, the total interaction is a product of all the terms. Given the parameters of the main part of the interaction (for example, ϵ , r_0 , a and r_c for the V_{Morse} part of $f_1(r)$), the parameters of the smoothed cutoff regions are uniquely determined by ensuring continuity and differentiability at the boundaries (r^{low} and r^{high} for $f_1(r)$).

Given a , θ^0 and $\Delta\theta^*$, b and θ^c can be extracted by solving Equations 2.20 and 2.21 simultaneously. An equivalent procedure can be performed for any of the product functions which constitute a given interaction.

There are many free parameters in this model. This is, however, somewhat deceptive. The parameters relating to smoothing the truncation of potentials are generally of minor importance (at least for the thermodynamics of the system), as the effect of a potential well

Interaction		Parameters		
$V_{\text{cross_stack}}$	$f_2(\delta r_{\text{HB}})$	$k = 47.5$ $\delta r_{\text{cross}}^{\text{low}} = 0.495$	$r_{\text{cross}}^0 = 0.575$ $\delta r_{\text{cross}}^{\text{high}} = 0.655$	$\delta r_{\text{cross}}^c = 0.675$
	$f_4(\theta_1)$	$a_{\text{cross},1} = 2.25$	$\theta_{\text{cross},1}^0 = \pi - 2.35$	$\Delta\theta_{\text{cross},1}^* = 0.58$
	$f_4(\theta_2)$	$a_{\text{cross},2} = 1.70$	$\theta_{\text{cross},2}^0 = 1.00$	$\Delta\theta_{\text{cross},2}^* = 0.68$
	$f_4(\theta_3)$	$a_{\text{cross},3} = 1.70$	$\theta_{\text{cross},3}^0 = 1.00$	$\Delta\theta_{\text{cross},3}^* = 0.68$
	$f_4(\theta_4) + f_4(\pi - \theta_4)$	$a_{\text{cross},4} = 1.50$	$\theta_{\text{cross},4}^0 = 0$	$\Delta\theta_{\text{cross},4}^* = 0.65$
	$f_4(\theta_7) + f_4(\pi - \theta_7)$	$a_{\text{cross},7} = 1.70$	$\theta_{\text{cross},7}^0 = 0.875$	$\Delta\theta_{\text{cross},7}^* = 0.68$
	$f_4(\theta_8) + f_4(\pi - \theta_8)$	$a_{\text{cross},8} = 1.70$	$\theta_{\text{cross},8}^0 = 0.875$	$\Delta\theta_{\text{cross},8}^* = 0.68$
$V_{\text{coax_stack}}$	$f_2(\delta r_{\text{coax}})$	$k_{\text{coax}} = 46$ $\delta r_{\text{coax}}^{\text{low}} = 0.22$	$\delta r_{\text{coax}}^0 = 0.4$ $\delta r_{\text{coax}}^{\text{high}} = 0.58$	$\delta r_{\text{coax}}^c = 0.6$
	$f_4(\theta_1) + f_4(2\pi - \theta_1)$	$a_{\text{coax},1} = 2.00$	$\theta_{\text{coax},1}^0 = \pi - 0.60$	$\Delta\theta_{\text{coax},1}^* = 0.65$
	$f_4(\theta_4)$	$a_{\text{coax},4} = 1.30$	$\theta_{\text{coax},4}^0 = 0$	$\Delta\theta_{\text{coax},4}^* = 0.8$
	$f_4(\theta_5) + f_4(\pi - \theta_5)$	$a_{\text{coax},5} = 0.90$	$\theta_{\text{coax},5}^0 = 0$	$\Delta\theta_{\text{coax},5}^* = 0.95$
	$f_4(\theta_6) + f_4(\pi - \theta_6)$	$a_{\text{coax},6} = 0.90$	$\theta_{\text{coax},6}^0 = 0$	$\Delta\theta_{\text{coax},6}^* = 0.95$
	$f_5(\cos(\phi_3))$	$a_{\text{coax},3'} = 2.00$	$\cos(\phi_3)_{\text{coax}}^* = -0.65$	
	$f_5(\cos(\phi_4))$	$a_{\text{coax},4'} = 2.00$	$\cos(\phi_4)_{\text{coax}}^* = -0.65$	

Table 2.2: Further model parameters.

is largely determined by its width and depth rather than the details of its edge. Furthermore, a number of parameters are essentially dictated by DNA geometry, and of those that are unconstrained, many are necessarily equal to others by symmetry. The parameters are also constrained by the requirement that the model behaves ‘like DNA’. In other words, if stacking and pairwise bonding are to drive the formation of double helices, rather than just cause collapse, the potentials need to have fairly narrow widths. Despite these caveats, there are still a large number of parameters to fit, especially via a top-down approach. This fitting is made more complex as it involves a compromise between the representation of various aspects of DNA. In particular, a given parameter may influence a wide range of properties and it is difficult to design a simple metric to compare the reproduction of thermodynamic and mechanical DNA behavior. In this case, lengths and potential minima were initially chosen to give model duplexes approximate B-DNA geometry. The stacking interaction strength and stiffness were then altered by hand to be consistent with the experimental thermodynamics reported for 14-base oligomers by Holbrook *et al.* [140]. Hydrogen-bonding

and cross-stacking potentials were then added, and adjusted to give duplex and hairpin formation thermodynamics consistent with the SantaLucia parameterization of the nearest-neighbour model [91], which can be viewed as an accurate empirical fit to experimental data. For comparison with Reference [91], I considered an ‘average base pair step’ – details are provided in Chapter 6 – as the model contains limited sequence dependence. Mechanical properties, such as persistence lengths, were then compared to experiment and interaction stiffnesses adjusted by hand to provide improved agreement. This process was then iterated until the current set of parameters was found.²

In general, the interaction energy in a coarse-grained model should be interpreted as a free energy, as it incorporates a number of implicit degrees of freedom [92], and thus it is plausible that interaction strengths could be temperature dependent. To reduce free parameters, I have avoided this temperature dependence except with regard to the stacking interaction. It was found that it was difficult to generate a stacking transition with an entropy as small as required (for details, refer to Chapter 6) whilst maintaining an appropriate stiffness for dsDNA. I therefore took the stacking strength parameter to be linearly dependent on temperature: over the range 270-370 K, the stacking strength increases by $\sim 6\%$, in effect reducing the entropy cost of the transition.

There are several possible causes of this underestimation of the transition width. A contribution may be that in order to replicate the flexibility of single strands, the conformation of bases is unrestricted except by excluded volume, certainly a significant simplification. For physical DNA, a range of different conformations are accessible (allowing the large flexibility of unstacked single strands), but the available fraction of configuration space is restricted by specific steric clashes, which would be exceedingly difficult to reproduce in a bead-spring model such as mine. This overestimate of available configurations is the compromise necessary to allow hairpins and nanostructures to form. An alternative cause may be the lack of stacking heterogeneity. In reality, each pair of bases will stack with a different strength, resulting in a different stacking probability from other neighbouring bases. When

²as the coaxial stacking term has an almost negligible effect on the properties discussed above, it was separately fitted to thermodynamic data on coaxial stacking [91, 141, 137, 138, 139, 136, 142, 143].

looking at the average properties of a mixed sequence, however, an observer would see a combination of a number of transitions which would appear broader than the individual transitions themselves. This broadening effect is absent in my average base model. Finally, the stacking interaction itself is thought to rely partially on hydrophobic effects [12, 144], and hence would be temperature dependent in any model without explicit water.

2.4.4 Neglected features of DNA

The model currently neglects some features of DNA. Although it incorporates sequence specificity (only A–T and G–C hydrogen bonds are possible), there is no other sequence dependence of interactions. I have made the simplifying assumption that noncomplementary base pairs have zero attraction, and also neglected the possibility of alternative base-pair geometries (such as Hoogsteen [12]). Due to this simplification, this version of the model cannot probe many sequence-dependent effects, such as the preferred sites of bubble formation (internal melting of base pairs) in DNA.

There are no explicit electrostatic interactions in the model, which may be expected to be important as bare ssDNA has a charge of $-e$ per base associated with the phosphate groups. For this reason, I fit to experimental data (where possible) at $[Na^+] = 500\text{ mM}$, where electrostatic properties are strongly screened. Indeed, at these ionic concentrations, the Debye screening length is approximately 4.3 \AA , smaller than the excluded volume diameter for backbone-backbone interactions in our model ($\sim 6\text{ \AA}$). At the shortest distances allowed by the steric interactions, charges would have an energy of $\sim 2kT$ in a Debye-Hückel approximation. Other authors have attempted to explicitly include a Debye-Hückel term [120], but also included a salt-dependent, medium-range attraction between strands in monovalent salt to facilitate hybridization, the physical origin of which is unclear. Due to this simplification, the model is unable to probe low salt regimes, and any result which relies upon DNA backbones coming into close proximity should be treated with caution.

The model also simplifies the geometry of DNA. Although the pitch, rise and diameter of helices are approximately consistent with experimental data, the grooves in between the backbones are of equal size. For physical DNA, by contrast, the major groove is significantly

larger than the minor groove [12]. This is important for protein binding, and is also likely to have consequences for the strain inherent in certain origami designs [59].

Finally, it should be noted again that the description of the behaviour of the backbone of single-strands is very simplistic. In particular, bases are fully able to rotate and bend about the backbone bond with no restriction except for steric clashes. Whilst this seems to give a reasonable description of large-scale single-stranded mechanical properties (as discussed in Chapter 5), one should be wary of any predictions that rely heavily on one or two nucleotides adopting a particular configuration.

The simplifications in the model were made partly to reduce the number of possible parameters. For example, sequence dependence would give 16 combinations of stacking pairs, each pair requiring several parameters to describe their interaction. It was also felt that, as an initial step in modelling, it was important to obtain a good physical representation of the underlying properties of DNA assembly (such as the generic dependence of melting temperature on length), before incorporating sequence specific or low salt effects. Furthermore, some generic effects may be obscured by sequence-specific terms (for instance, free-energy profiles such as those in Chapter 6 would have sequence-dependent fluctuations overlying the general trend).

2.4.5 Additional parameters required for dynamical simulations

The parameters presented so far specify the thermodynamic properties of the model. To use explicitly dynamical algorithms, one must introduce masses, moments of inertia and drag coefficients. For simplicity, I assume that each nucleotide is a uniform density sphere centred on the backbone-base axis, $0.24l$ from the backbone site (this assumption is discussed in Section 3.2). Taking the unit of mass in simulations to be $M = 100$ AMU, an average mass of a nucleotide is around $3.1575M$, and the moment of inertia is 0.43512 in simulation units. Using Langevin dynamics to simulate the model is discussed further in Chapter 3 and Appendix B, where the damping coefficients are defined and given values. Note that this definition of M , when combined with the reduced length and energy scales L and E , defines the reduced unit of time in simulations $T = (E/M/L^2)^{-0.5} = 1.706$ ps.

Chapter 3

Methods

Due to the number of nucleotides in a typical system of interest, and the complexity of the interactions between them, it is impossible to find exact analytical expressions for the behaviour of a general coarse-grained model of DNA. Computer simulations can provide numerical information in the absence of any exact solutions, and guide the application of simpler, analytic descriptions.

For an atomistic (classical) model of DNA with explicit solvent, the obvious way to extract information from the model would be to integrate Newton's equations of motion from a variety of initial states. The trajectories generated could then be used to extract statistical and kinetic properties of the system.

For efficiency of simulation, solvent effects are usually treated implicitly in coarse-grained models. Thus solvent-mediated interactions must be directly included as interactions between solute particles. Even if, however, the statistical consequences of these interactions are perfectly captured by the implicit solvent model, the question of how to generate dynamical trajectories remains. Simply integrating Newton's equations would generate unphysical ballistic motion of solute particles between collisions, and would also result in simulations at constant total energy (whereas in reality the solvent acts as a thermal reservoir for the solute).

In this chapter, the two basic algorithms used in this work (*Langevin dynamics* and *cluster-move Monte Carlo*) are discussed. Techniques for enhancing sampling of thermodynamics (umbrella sampling) and kinetics (forward flux sampling) are also introduced.

3.1 Monte Carlo simulation

Technically, Monte Carlo (MC) simulation evades the question of dynamics altogether, and simply randomly generates microstates¹ with a probability distribution appropriate to a given ensemble. For example, consider a system in the canonical ensemble (i.e., a system at constant volume but in contact with a thermal reservoir at temperature T). Elementary classical statistical mechanics dictates that the microstates of a system of N bodies should be occupied with a density:

$$\rho(\mathbf{r}^N, \mathbf{p}^N, \boldsymbol{\Omega}^N, \mathbf{L}^N) \propto \exp(-\beta \mathcal{H}(\mathbf{r}^N, \mathbf{p}^N, \boldsymbol{\Omega}^N, \mathbf{L}^N)), \quad (3.1)$$

where \mathbf{r}^N , \mathbf{p}^N , $\boldsymbol{\Omega}^N$ and \mathbf{L}^N are the centre of mass position, centre of mass momentum, angular orientation and angular momentum of N bodies, $\beta = 1/k_B T$ and \mathcal{H} is the Hamiltonian of the system [145].

Generally, \mathbf{p}^N and \mathbf{L}^N enter the Hamiltonian in the form:

$$\mathcal{H} = \sum_n^N \left(\frac{(\mathbf{p}_n)^T (\mathbf{p}_n)}{2m_n} + \frac{1}{2} (\mathbf{L}_n)^T I_n^{-1} \mathbf{L}_n \right) + V(\mathbf{r}^N, \boldsymbol{\Omega}^N), \quad (3.2)$$

where I_n^{-1} is the inverse of the 3x3 moment of inertia tensor. The quadratic terms in the generalized momenta are trivially integrated over, giving a constant independent of \mathbf{r}^N and $\boldsymbol{\Omega}^N$. As a consequence, the occupation of microstates with a given configuration $(\mathbf{r}^N, \boldsymbol{\Omega}^N)$ is simply given by:

$$\rho(\mathbf{r}^N, \boldsymbol{\Omega}^N) \propto \exp(-\beta V(\mathbf{r}^N, \boldsymbol{\Omega}^N)). \quad (3.3)$$

Monte Carlo simulation (in the canonical ensemble) generates configurations sampled from this distribution. The role of the solvent as a thermal bath is therefore directly incorporated – generalizations to alternative ensembles can also be considered.

¹To avoid confusion, I shall take the term *microstate* to be a given set of positions, orientations, velocities and angular velocities for the system in question. The *configuration* is defined by the positions and orientation of entities in the system. Finally, I shall use the term *state* more broadly, to apply to a set of microstates or configurations which are in some way similar (for instance, the ‘duplex state’ consists of all microstates in which the two strands are bound to each other).

3.1.1 Metropolis Monte Carlo

An enduringly simple and elegant Monte Carlo method was developed by Metropolis *et al.* in 1953 [146]. In this algorithm, one devises an arbitrary (ergodic) set of trial moves with which to generate new configurations from existing ones. Typically, these moves involve translations (and rotations) of single particles. The simulation then steps stochastically through the states of the system. At each step, a single move is chosen randomly which converts an initial configuration μ into a new configuration ν , and the move is accepted with the probability:

$$P_{\text{acc}}(\mu \rightarrow \nu) = \min\{1, \exp(-\beta(E^\nu - E^\mu))\}, \quad (3.4)$$

where E^μ and E^ν are the configurational energies in the old and new configurations. Provided the probability of generating a trial move from state μ , $P_{\text{gen}}(\mu \rightarrow \nu)$, is equal to the probability of generating the reverse move when in state ν , $P_{\text{gen}}(\nu \rightarrow \mu)$, one can easily show that:

$$\frac{P_{\text{move}}(\mu \rightarrow \nu)}{P_{\text{move}}(\nu \rightarrow \mu)} = \frac{P_{\text{gen}}(\mu \rightarrow \nu)P_{\text{acc}}(\mu \rightarrow \nu)}{P_{\text{gen}}(\nu \rightarrow \mu)P_{\text{acc}}(\nu \rightarrow \mu)} = \exp(-\beta(E^\nu - E^\mu)). \quad (3.5)$$

This condition is known as *detailed balance*. If it holds for all pairs of states of the system, then the stationary distribution of the Monte Carlo process (the distribution that it samples from in the limit of infinite steps) is such that:

$$\frac{\rho(\mu)}{\rho(\nu)} = \frac{P_{\text{move}}(\mu \rightarrow \nu)}{P_{\text{move}}(\nu \rightarrow \mu)} = \exp(-\beta(E^\nu - E^\mu)). \quad (3.6)$$

Such a stationary distribution is exactly the one required by Equation 3.3.

Metropolis MC has been extremely widely used since its invention, and has been employed to study self-assembly processes [50, 147, 148] and even DNA [113]. One of its major advantages is that it does not involve integration of equations of motion, and hence there is no issue with convergence or stability if the typical moves are fairly large (one can even choose unphysical moves). Such large moves mean that systems simulated using Metropolis MC can be made to equilibrate extremely quickly (in terms of computer time).

It does, however, have its drawbacks. Firstly, as there are no equations of motion, one cannot rigorously study the dynamics of processes using Monte Carlo. It has been argued that Metropolis MC can give a qualitative picture of the ease with which a system explores its state-space [50, 147, 148], but it is difficult to make concrete statements beyond the fact that isolated particles will undergo diffusive dynamics [149].

One of the particular issues with interpreting a Metropolis MC dynamically is the tendency to suppress the diffusion of clusters of particles [149], as there is no opportunity for particles to move cooperatively. For an aggregate to move, each individual element must separately move in the desired direction. Furthermore, the vast majority of large moves of monomers will cause a significant increase in energy. Consequentially, these moves will tend to be rejected by the criterion of Equation 3.4. The result is that strongly bound clusters, which in reality move in coordination due to the forces between their constituents, diffuse unreasonably slowly compared to monomers.

The problem of suppressed cluster motion can become so severe that it hinders not only dynamical interpretation of Metropolis MC trajectories, but also the ability of these trajectories to reach states representative of equilibrium [51, 132, 149].

3.1.2 Cluster moves and Virtual Move Monte Carlo

In order to overcome the issue of the suppressed motion of aggregates in Monte Carlo simulations, a variety of ‘cluster algorithms’ have recently been introduced [132, 149, 150, 151]. In general, these algorithms aim to reproduce cooperative motion by proposing changes of state that involve several particles moving in unison, in a way that reflects the interactions within the system.

In particular, the Virtual Move Monte Carlo (VMMC) algorithm of Whitelam and Geissler [132, 149], illustrated in Figure 3.1, has been extensively used in this work.² The algorithm requires the random choice of a seed particle and a move (typically translation or rotation). Links are then attempted with all particles with which the seed interacts in the

²The algorithm used is actually the variant detailed in the appendix of Reference [132].

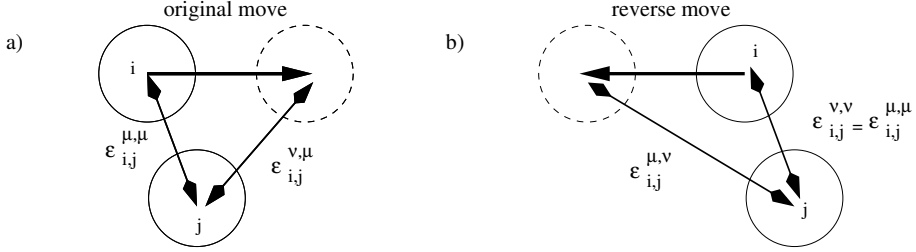


Figure 3.1: Cluster building in a VMMC simulation step. Particle i is randomly selected and a random move is chosen (a). Particle j interacts with i in the initial state, and the appropriate pairwise energies before and after the move are calculated. A partial link is formed between i and j with a probability given by equation 3.7. An equivalent procedure is performed in (b) for the reverse move. If a link would also be formed in this case, a full link is formed and j is added to the cluster. The procedure is repeated until all particles that interact with particles in the cluster have been tested.

initial state, and are formed with the probability:

$$P_{\text{link}}(i, j) = \max(0, 1 - \exp(-\beta(\epsilon_{i,j}^{\nu,\mu} - \epsilon_{i,j}^{\mu,\mu}))), \quad (3.7)$$

where $\epsilon_{i,j}^{\nu,\mu}$ is the pairwise energy between the seed particle i and a neighbor j after i has been moved but j has not, and $\epsilon_{i,j}^{\mu,\mu}$ is their initial energy. The reverse move, in which both particles begin in the new state and the seed is moved in the opposite direction, is then considered. If the linking is also successful in this case, a ‘full link’ is formed, and the new particle becomes part of the cluster and moves in conjunction with the seed, preserving their pairwise interaction energy. Otherwise, the link is termed a ‘partial link’ and the new particle is not included in the cluster. Links are attempted from all particles that are added to the cluster.

Due to the complexity of the cluster building, forwards and reverse moves are not generated with equal probability. Preserving the detailed balance condition of Equation 3.5 therefore requires an alternative criterion to that in Equation 3.4:

$$P_{\text{acc}}(\mu \rightarrow \nu | \mathcal{R}) = D_{\mathcal{R}}(\mu \rightarrow \nu) \min\{1, \Pi_{i,j}^{n/o} \exp(-\beta(\epsilon_{i,j}^{\nu} - \epsilon_{i,j}^{\mu}))\}. \quad (3.8)$$

In this case, $P_{\text{acc}}(\mu \rightarrow \nu | \mathcal{R})$ is the probability of accepting a move from state μ to state ν , given a realization of links and partial links \mathcal{R} . $D_{\mathcal{R}}$ is a factor which is zero if there are any partial links between a particle in the cluster and a particle outside it (and unity otherwise). Finally, the product $\Pi_{i,j}^{n/o}$ is taken over all pairs of particles which are non-interacting in

state μ and have positive energy in state ν , and vice versa. $\epsilon_{i,j}^\nu$ and $\epsilon_{i,j}^\mu$ are simply pairwise energies in the trial configuration and before the move respectively.

The algorithm and acceptance probability are more opaque than for Metropolis MC. The net result, however, is that collective moves are possible for aggregates (or subsections of aggregates) that at least partially reflect the gradients in potential energy of the system. The outcome, at least for the DNA model introduced in Chapter 2, is vastly improved sampling when compared to basic Metropolis MC, which more than compensates for the extra complexity of the algorithm.

The primary distinction between this algorithm, and others such as that suggested by Troisi and coworkers [151], is that this version considers the energy change due to the move when building a cluster, rather than simply the strength of interaction in the initial state. As a consequence, proposed moves are more sensitive to potential energy gradients than in other algorithms. A drawback is that the cluster-building process is more complex, and hence each step requires greater computational power. The relative advantages of these factors remains unclear at the time of writing.

By their very nature, cluster algorithms are most effective when the majority of the interactions in the system are pairwise, as clusters are built up by considering pairwise energies. Multi-body potentials can be included, but not within the cluster building framework, and they simply modify the final acceptance criterion:

$$P_{\text{acc}}(\mu \rightarrow \nu | \mathcal{R}) = D_{\mathcal{R}}(\mu \rightarrow \nu) \min \left\{ 1, \left(\prod_{i,j}^{n/o} e^{(-\beta(\epsilon_{i,j}^\nu - \epsilon_{i,j}^\mu))} \right) e^{-\beta(E_m^\nu - E_m^\mu)} \right\}, \quad (3.9)$$

where here E_m^μ and E_m^ν are the multi-body contributions to the total energy of the initial and trial-move states. The utility of pairwise interactions in this regard was a consideration in designing the model, and led to the use of rigid-body nucleotides.

As with Metropolis MC, cluster move algorithms do not technically provide a dynamical representation of the time evolution of the model system. If the moves used are ‘local’, involving small displacements and rotations of clusters of particles, one can interpret the sequence of states visited as a pseudo-dynamical description of the model. The rates of processes then give some idea of the ease with which the free-energy landscape of the system

is navigated. However, although cluster moves overcome the Metropolis issue of extremely slowly diffusing clusters, it is not easy to ensure that clusters of different sizes move at the appropriate relative rates. In addition, it is an even more challenging task to ensure that internal relaxation and diffusion of clusters happen at appropriate rates. Furthermore, due to the subtleties of the move generation and acceptance criteria, it is difficult to compare the rates of processes in simulations of slightly different systems.

3.2 Langevin dynamics

The Langevin formalism is a self-consistent way of incorporating an implicit source of noise and dissipative forces into deterministic equations of motion [152], such that the simulation will sample microstates with a weight given by Equation 3.1. For example, the model presented here can be described in terms of a Hamiltonian $\mathcal{H}(\mathbf{r}^N, \mathbf{p}^N, \mathbf{q}^N, \boldsymbol{\Pi}^N)$, which is a function of the positions, momenta, orientations (here represented through quaternions – refer to Appendices A and B for details) and generalized angular momenta of all objects in the system. Such a Hamiltonian would generate deterministic motion via the equations:

$$\dot{\mathbf{r}}_i = \frac{\partial}{\partial \mathbf{p}_i} \mathcal{H}, \quad \dot{\mathbf{p}}_i = -\frac{\partial}{\partial \mathbf{r}_i} \mathcal{H}, \quad \dot{\mathbf{q}}_i = \frac{\partial}{\partial \boldsymbol{\Pi}_i} \mathcal{H}, \quad \dot{\boldsymbol{\Pi}}_i = -\frac{\partial}{\partial \mathbf{q}_i} \mathcal{H}. \quad (3.10)$$

Following the Langevin formalism, as detailed by Davidchack *et al.* [153], one augments these equations with damping and noise terms so that:³

$$\dot{\mathbf{p}}_i = -\frac{\partial}{\partial \mathbf{r}_i} \mathcal{H} - \gamma \mathbf{p}_i + b \mathbf{w}_i(t), \quad \dot{\boldsymbol{\Pi}}_i = -\frac{\partial}{\partial \mathbf{q}_i} \mathcal{H} - \Gamma \mathbf{G}(\mathbf{q}_i, \boldsymbol{\Pi}_i) + B \mathbf{W}_i(t). \quad (3.11)$$

Here γ and Γ give the strength of linear and rotational damping, and $G(\mathbf{q}_i, \boldsymbol{\Pi}_i)$ is a four-dimensional vector that couples the rotational momenta. $\mathbf{w}_i(t)$ and $\mathbf{W}_i(t)$ are 3- and 4-vectors, each component being a gaussian distributed random force with zero mean and lag-covariance $\delta(t, 0)$. b and B are constants which give the magnitude of the random forces. In principle, γ , Γ , b and B are matrices, coupling the noise and friction applied to

³Here I consider only additive noise, and can therefore neglect the subtleties of the distinction between Ito and Stratonovich calculus [152].

different degrees of freedom. For simplicity, I have chosen to assume that all translational modes (and separately all rotational modes) experience the same strength of noise and damping.

By requiring that the stationary distribution of the Fokker-Planck process that corresponds to this Langevin equation is given by the Boltzmann distribution, one can relate the damping and noise terms by fluctuation-dissipation relations, as discussed in Appendix B:

$$\gamma = \frac{\beta b^2}{2m}, \quad \Gamma = \frac{\beta B^2}{2M}, \quad \text{and} \quad M = \frac{4}{\text{Tr}(I^{-1})}. \quad (3.12)$$

Here, I is taken as the moment of inertia tensor in the principal axis frame, and the form of Here γ and Γ give the strength of linear and rotational damping, and $G(\mathbf{q}_i, \mathbf{\Pi}_i)$ is given in Appendix B. These equations can be integrated numerically, using the methods outlined by Davidchack *et al.* [153] – in this work, I use the ‘Langevin A’ algorithm.

Unlike Monte Carlo, Langevin algorithms are explicitly dynamical, but issues remain with inferring kinetics from the simulations. Firstly, it is well known that coarse-graining has the effect of speeding up dynamics by smoothing free-energy landscapes [154]. Although this may seem undesirable, it is, in a sense, one of the advantages of coarse graining that allows long time scale processes to be accessed. Secondly, rare event kinetics are exponentially sensitive to the heights of free-energy barriers, which are difficult to fit in a coarse-grained model. Perhaps the most important problem, however, is the lack of hydrodynamic interactions in coarse-grained Langevin methods.

Hydrodynamic effects arise because the solvent flow induced by the motion of one solute particle affects the motion of nearby solute particles [155]. The affect can be approximately treated within an implicit solvent model, for example by use of Oseen tensor formalism [155], but such methods are extremely computationally expensive.

A typical consequence of neglecting hydrodynamics is that collective motion is suppressed. For example, the Rouse model of polymer diffusion (which neglects hydrodynamic interactions) predicts that the diffusion coefficient scales as $1/N$, where N is the molecular weight of the polymer [155]. By contrast, the Zimm model (which incorporates an approxi-

mate treatment of hydrodynamics) predicts a diffusion coefficient which scales as the inverse of the radius of gyration [155].

When implementing Langevin dynamics, one has a degree of choice of the form of the friction and damping terms, although they must always be related by a fluctuation-dissipation theorem if the equilibrium distribution is to be the correct one. Given the degree of simplification inherent in the simulation, it seems unlikely that being precise about the damping terms will prove significantly beneficial. This reasoning justified the simplifying assumptions about the form of the noise and damping terms, effectively treating each nucleotide as a sphere for dynamical purposes

If each nucleotide is taken as a sphere of radius $r_0 = 5 \text{ \AA}$, mass $m = 315.75 \text{ AMU}$ and uniform density, continuum hydrodynamics for an isolated nucleotide would then give

$$\gamma = \frac{6\pi\eta r_0}{m} \quad \text{and} \quad \Gamma = 8\pi\eta r_0^3 \text{Tr}(I^{-1}) = 10\gamma \quad (3.13)$$

In principle, one could take $1/\gamma = 0.03$ in reduced time units (giving $\gamma \approx 20 \text{ ps}^{-1}$), which would correspond to a nominal value of $\eta = 1.1 \times 10^{-3} \text{ kg m}^{-1} \text{ s}^{-1}$, similar to experimental values for water at around 300 K.

Due to hydrodynamic interactions, however, such a drag coefficient would seriously underestimate the diffusion of large aggregates, which are necessary for the majority of processes of interest in this work (for example, the association of two single strands to form a duplex). In particular, it was found that such large drag coefficients led to prohibitively slow dynamics for large systems, such as the walker discussed in Chapter 8. Given the number of simplifications already present in the Langevin model of dynamics, it was decided that there was little justification in maintaining high drag coefficients at the expense of efficient sampling. It was therefore decided to set $\gamma = 1$ and $\Gamma = 3$ in reduced units, values which maintain the diffusive behaviour of the system but accelerate the dynamics.⁴

⁴Despite this reduction in noise, dynamics are still highly damped. For example, a simulation of a 10- bp duplex at 300 K initiated with all 10 base pairs formed but far from an optimal configuration (with around 70% of the typical binding energy and no kinetic energy) will reach states typical of equilibrium in around 10 units of reduced time. For comparison, diffusion over its own length is around 10 – 100 times slower for a 10- bp duplex.

Despite the difficulty of directly inferring time scales from Langevin simulations, there are clear advantages over Monte Carlo for analyzing kinetics. Firstly, there are qualitatively different methods to generate trial MC moves (for example, the algorithms discussed in Section 3.1.2), with potentially large (and poorly understood) consequences for the kinetics of simulated processes. Secondly, for a given implementation of a cluster MC algorithm, it is extremely difficult to compare rates of processes for similar but distinct simulations. This difficulty arises because the probabilities with which moves are generated and accepted vary in a non-trivial manner with factors such as system size, temperature and external forces. For example, the comparison of the rate of binding of a DNA walker’s foot to a track with and without tension, as presented in Chapter 8, would have been much more problematic with MC simulations.

By contrast, applying Langevin dynamics to a given model involves significantly less arbitrary choice (and the choices are of a more quantitative, rather than a qualitative nature). Further, the kinetics of different systems can be directly compared, as the algorithm changes in a more intuitive and well-understood fashion as the parameters are changed. Nonetheless, all results should be interpreted with caution. In particular, due to the absence of hydrodynamic interactions between solute particles, small scale fluctuations will be excessively fast compared to large scale diffusion.

Given the caveats associated with Langevin dynamics, it is only sensible to compare relative rates for similar processes. Further, any relative difference that is observed should only be believed if a physical cause (which is not a peculiar result of the dynamics) can be identified. One possible consistency check is to compare the Langevin results with the pseudo-dynamics of VMMC – if the results are similar, despite the vast differences in algorithms, one can be confident that the result is a generic one.

In this work I use a time step of $h = 0.003$ units of simulation time (~ 5 ps). The accuracy of this time step, and of my implementation of both Langevin Dynamics and VMMC algorithms in general, is checked in Appendix C.

3.3 Advanced sampling techniques

Much of this thesis involves studying processes that require duplex hybridization and displacement. Such processes are rare-event dominated, in that they typically involve transitions between (meta)stable states such as duplexes and dissociated single strands, via unfavourable intermediate states.

Such processes can require long simulations, and it is particularly difficult to obtain enough statistics for reliable thermodynamic or kinetic data. As a consequence, a variety of techniques have been developed to improve the sampling of rare events – this work utilizes *umbrella sampling* and *forward flux sampling*.

3.3.1 Umbrella sampling

Umbrella sampling, introduced by Torrie and Valleau [156], uses unphysical biasing of configurations to encourage transitions between (meta)stable states. Firstly, an order parameter $Q(\mathbf{r}^N, \Omega^N)$, which identifies the extent of reaction, is defined. If the order parameter Q is well chosen, the free energy $F(Q)$ (defined in terms of the probability $P(Q)$ that the system is in state Q by $F(Q) = -k_B T \ln P(Q)$) will have minima at the metastable states and high values at the unfavourable states that cause the reaction bottleneck.

Umbrella sampling involves introducing an additional bias $w(Q)$ to the sampling distribution, so that configurations are sampled according to $w(Q)e^{-\beta V(\mathbf{r}^N, \Omega^N)}$. $w(Q)$ is generally chosen to increase the probability of occupying transition states, thereby increasing the flux between minima.

The thermodynamic average of any function $A(\mathbf{r}^N, \Omega^N)$ is given by:

$$\langle A \rangle = \frac{\int d\mathbf{r}^N d\Omega^N A(\mathbf{r}^N, \Omega^N) e^{-\beta V(\mathbf{r}^N, \Omega^N)}}{\int d\mathbf{r}^N d\Omega^N e^{-\beta V(\mathbf{r}^N, \Omega^N)}} = \frac{\int d\mathbf{r}^N d\Omega^N \frac{A(\mathbf{r}^N, \Omega^N)}{w(Q)} w(Q) e^{-\beta V(\mathbf{r}^N, \Omega^N)}}{\int d\mathbf{r}^N d\Omega^N \frac{1}{w(Q)} w(Q) e^{-\beta V(\mathbf{r}^N, \Omega^N)}}. \quad (3.14)$$

In order to unbias the results, therefore, one records $A/w(Q)$ and normalizes with respect to $1/w(Q)$.

In this work, umbrella sampling is used in conjunction with VMMC. $w(Q)$ is in effect an additional multi-body interaction which must be considered separately from the cluster gen-

eration, such that the final term in the braces of Equation 3.9 is modified by $w(Q^\nu)/w(Q^\mu)$, with Q^μ and Q^ν representing the initial and final values of the order parameter respectively.

For particularly difficult systems, it can be advantageous to split the simulation into several ‘windows’, each of which is strongly biased to remain within a certain range of Q . Equilibration within each window is easier than for the transition as a whole, and the separate windows can be combined using the WHAM algorithm of Kumar *et al.* [157].

3.3.2 Forward flux sampling

Although umbrella sampling is an extremely powerful technique for extracting thermodynamic averages, it technically provides no direct information on the pathways and kinetics of assembly processes. Forward flux sampling is an alternative which allows one to calculate the flux of systems between two local minima of free energy, and also sample from the trajectories that link the two minima [158, 159].

Once again, one considers an order parameter Q which measures the extent of the reaction, such that non-interacting interfaces, λ_{n-1}^n can be drawn between consecutive values of Q . Initially, simulations are performed that begin in the lowest value of Q (which I shall define as $Q = -2$), and the flux of trajectories crossing the surface λ_{-1}^0 (for the first time since leaving $Q = -2$) is measured.

The total flux of trajectories from $Q = -2$ to the alternative minima ($Q = N$) is then calculated as the flux across λ_{-1}^0 from $Q = -2$, multiplied by the conditional probability that these trajectories reach $Q = N$ before returning to $Q = -2$. This probability can be factorized into the product of the probabilities of trajectories starting from the interface λ_{n-1}^n reaching the interface λ_n^{n+1} before returning to $Q = -2$.

$$P(\lambda_{N-1}^N | \lambda_{-1}^0) = \Pi_n^N P(\lambda_{n-1}^n | \lambda_{n-2}^{n-1}). \quad (3.15)$$

The simulation then proceeds by randomly loading microstates which correspond to the crossing of λ_{-1}^0 , and using these as initial points from which to estimate $P(\lambda_0^1 | \lambda_{-1}^0)$. The process is then iterated for successive interfaces, allowing the estimation of flux and the construction of trajectories sampled from the distribution of transition pathways.

Chapter 4

Finite Size Effects

If the self-assembly of model DNA systems was easy to simulate, the best way to study such processes would be to model a large system where a number of the assembled structures can simultaneously form. This direct approach is not generally feasible, however, as the presence of significant free energy barriers to assembly makes equilibrium hard to achieve. Equilibration can be driven by rare-event techniques such as umbrella sampling (see Chapter 3), but these techniques are best suited to simulating a single target structure (I am unaware of any studies where biased sampling is used to simulate the formation of multiple targets). So if the assembly of a single structure is simulated, what issues arise with inferring the properties of a bulk system of the same strands?

Firstly, interactions between assembled structures are neglected. This is often a good approximation, because the interactions between them are likely to be mainly associated with excluded volume – any attractions are likely to be weak compared to the forces associated with the assembly itself – and assembly often occurs at relatively low concentrations. In addition to neglecting the interactions of assembled structures, such simulations will not capture states in which aggregates larger than the target structure have formed. Depending on the details of the system, these aggregates may constitute metastable states that can be significant in the dynamics.

The second potential source of error is due to neglected fluctuations of the local concentration of reactants, and this is the topic of this chapter. In particular, I will show that these finite-size errors in canonical ensemble simulations of dimerization can be significant,

but also how they can be corrected under the assumption that species behave ideally. I will also examine how the assembly yields converge towards the bulk values as the system size is increased, enabling the assumptions of the corrections to be checked. In Appendix D I discuss larger clusters and simulations in the grand canonical ensemble.

Although the extrapolation is not a complex procedure, it has been neglected in the past. In particular, simulations have been performed on DNA duplex formation [120, 121, 126, 127] without applying corrections. An attempt to correct for finite size effects during duplex formation was made by Prytkova *et al.* [122], but the method (which is not justified in the paper) involves an unclear renormalizaton of temperature and doesn't appear to correct for concentration fluctuations.

One should note that specific methods have been developed to calculate the equilibrium thermodynamics of heterodimer formation in the protein-ligand binding literature [160]. Such techniques aim to estimate the partition function of bound and separated molecules directly, and typical methods include calculating a potential of mean force for a certain pathway between bound and unbound structures, incorporating the effects of overall translational degrees of freedom separately. These techniques are optimized for problems of great computational difficulty and do not generalize well to multicomponent assembly. Furthermore, they lack the flexibility and simplicity of the approach analysed here in which nothing need be approximated or assumed about the nature of bonding, and no pathway need be imposed *a priori*.

4.1 Dimer formation in the canonical ensemble

4.1.1 Heterodimer formation

The physical cause of finite size statistical corrections can be seen by considering heterodimer formation – such a system may correspond to protein binding or DNA hybridization. Consider a simulation in a periodic cell of volume v , containing one monomer of type ‘A’ and one of type ‘B’. Assuming a criterion exists for defining a subset of states as ‘bound’,¹ a

¹The details of this criterion are not important, except for the fact that bound particles must be separated by a small distance compared to the simulation volume, and that the correlation of particles in the unbound

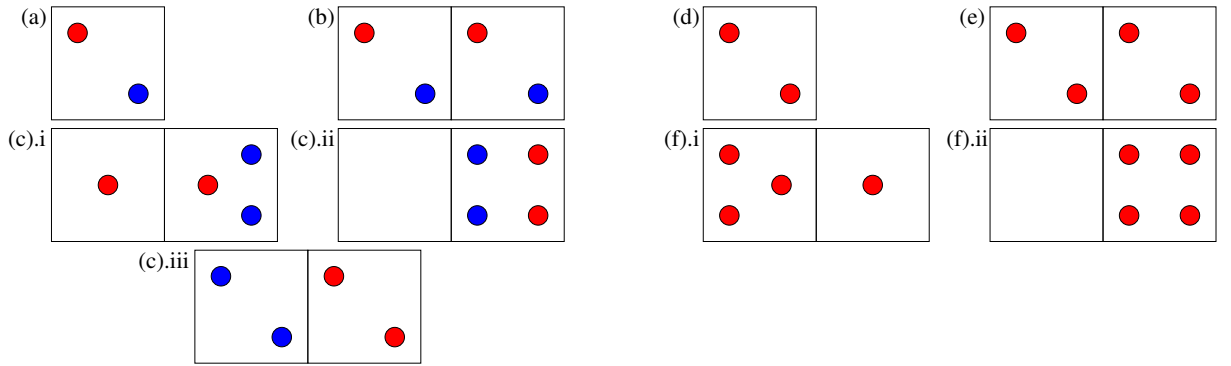


Figure 4.1: (a) Shows two different particles in a box of size v . One can imagine doubling the volume to $2v$ whilst doubling the number of particles, therefore maintaining the average concentration, as shown in (b). For the purposes of analysis, the new volume can be split in half with a line as indicated. All states with one of each of A and B on either side of the line, such as (b), will provide the same statistics as the original system. Macrostates such as those shown in (c), however, will have different statistics: for instance, (c).iii necessarily has a binding fraction of zero. (d)-(f) show an equivalent construction for homodimer formation.

simulation will estimate the relative probability with which bound (AB) and unbound (A,B) states are observed in such a system:

$$\Phi = \frac{\text{probability(AB)}}{\text{probability(A, B)}}, \quad (4.1)$$

with the fraction of bound pairs given by:

$$f_1 = \Phi / (1 + \Phi). \quad (4.2)$$

Naively, one might hope that $f_1 = f_\infty$, the bulk equilibrium bonding fraction at the same temperature and concentration. Unfortunately, this is not the case, because although the average concentration of a bulk system is matched, important concentration fluctuations are neglected (as shown in Figure 4.1).

One can apply corrections using simple thermodynamic arguments, if it is assumed that interactions between all particles that are not in a dimer state are negligible. Consider a periodic system of volume Dv , with D an integer,² with the same average concentration as the system with $D = 1$. I define:

- Z_{AB} and $Z_{A,B}$ as the partial partition functions of the $D = 1$ system when confined to the relevant subset of states. For future convenience, these quantities are defined

state must be small.

²In the limit $D \rightarrow \infty$, the details of boundary conditions should become irrelevant.

using *distinguishable statistics*, although it does not matter at this stage. Note that $\Phi = Z_{AB}/Z_{A,B}$.

- N as the total number of particles of type A or B (here $N = D$).
- N_i as the number of molecules of species i (in this case i is A, B or AB).
- q_i as the single particle partition function for species i , in the volume Dv , with the internal degrees of freedom treated using *indistinguishable statistics*.
- μ_i as the chemical potential of species i .

The μ_i are given by a standard result of statistical mechanics:

$$\mu_i = -k_B T \frac{\partial}{\partial N_i} \ln \left(\frac{q_i^{N_i}}{N_i!} \right) \approx -k_B T \ln \left(\frac{q_i}{N_i} \right), \quad (4.3)$$

where the approximation becomes an equality in the thermodynamic limit. In this limit, equilibrium thermodynamics gives $\sum_i \nu_i \mu_i = 0$, where ν_i are the stoichiometric coefficients of the species in the reaction, resulting in:

$$\frac{N_{AB}}{N_A N_B} = \frac{q_{AB}}{q_A q_B}. \quad (4.4)$$

As each q_i scales with the volume of the system:

$$q_{AB} = D Z_{AB}, \quad (4.5)$$

$$q_A q_B = D^2 Z_{A,B}, \quad (4.6)$$

which gives (using $D = N$):

$$\frac{[AB]}{[A][B]} = \frac{v Z_{AB}}{Z_{A,B}} = v \Phi = K_{A,B}^{\text{eq}}. \quad (4.7)$$

Note that the quantity $Z_{AB}/Z_{A,B}$ is that which is generally directly estimated in protein/ligand binding studies [160]; this is then multiplied by a reference concentration to give the equilibrium constant.

Substituting (4.1), (4.5) and (4.6) into (4.4) yields:

$$\frac{f_\infty}{(1 - f_\infty)^2} = \Phi \quad (4.8)$$

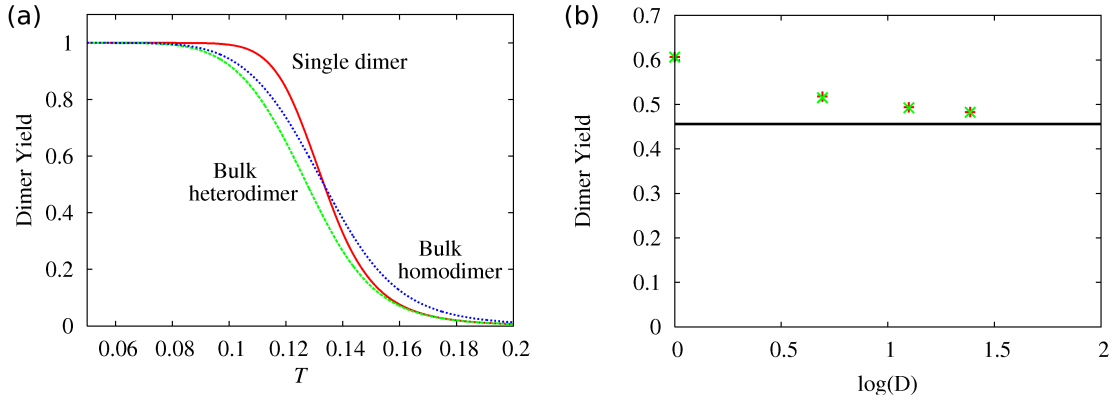


Figure 4.2: (a) Dimer yield for a system described by a two-state model $Z_2/Z_{1,1} = \exp(-\Delta E/T + \Delta S)$, with $\Delta E = 2$ and $\Delta S = 15$ in reduced units, with the values chosen for illustrative convenience. Plotted are the yield for a two-particle system and the bulk values at the same average concentration for homodimers and heterodimers. (b) Heterodimer yield as a function of system size D , with average concentration fixed. The ‘x’ symbols indicate results from simulations of my DNA model that were capable of forming D DNA duplexes of 5 bp, and the ‘+’ are the predictions of Equation (4.11), with Φ chosen to reproduce the $D = 1$ result. The solid line indicates f_∞ . Simulations were performed in a periodic cell of length $l = 10 \times 2^{(\log_2 D)} L$, where L is the reduced length scale of the model defined in Chapter 2, at a temperature of 303.03 K. At each D , a minimum of 1.5×10^{11} VMMC steps per strand were performed in total. No statistically significant deviations from the theory presented here were found.

$$\Rightarrow f_\infty = \left(1 + \frac{1}{2\Phi}\right) - \sqrt{\left(1 + \frac{1}{2\Phi}\right)^2 - 1}. \quad (4.9)$$

In this case, $f_\infty < f_1$ for all values of Φ , as is illustrated for a model dimer-forming system in Figure 4.2. It is also noticeable that the transition is wider for the bulk system. The physical causes of these two effects will be discussed at the end of Section 4.1.3.

4.1.2 Heterodimer convergence

It is useful to consider how the bonding fraction converges to the bulk result as the system size is increased from one cluster to the thermodynamic limit. Consider a system of volume Dv , and calculate the fraction of dimers (f_D) as a function of Φ , again neglecting interactions except dimer formation. Consider the macrostate with b dimers formed (out of a possible D). The partition functions of individual monomers and dimers scale with the size of the system (Dv), and the partition function of the system is the product of the individual partition functions together with combinatorial factors. Using Z_{AB} and $Z_{A,B}$ as defined before, the partition function of a macrostate with b dimers (using distinguishable statistics) is given

by:

$$Z_b(D) = \frac{(DZ_{AB})^b (D^2 Z_{A,B})^{D-b}}{b!} \left(\frac{D!}{(D-b)!} \right)^2, \quad (4.10)$$

in which the combinatorial factor is obtained from the total number of permutations of A and B ($D!^2$) divided by the permutations which exchange monomers for monomers ($(D-b)!^2$) or dimers for dimers ($b!$). The expression is divided by $D!^2$ to make the statistics indistinguishable. f_D is found in the usual way, using equation (4.1) to simplify:

$$f_D = \frac{\sum_{b=1}^D b \left(\frac{\Phi}{D}\right)^b \left(\frac{1}{(D-b)!}\right)^2 \frac{1}{b!}}{\sum_{b=0}^D D \left(\frac{\Phi}{D}\right)^b \left(\frac{1}{(D-b)!}\right)^2 \frac{1}{b!}} = \frac{\sum_{b=0}^D b Z'_b}{\sum_{b=0}^D D Z'_b}. \quad (4.11)$$

Plotting f_D against D for $\Phi = 1.54$ (Figure 4.2(b)) shows that the bonding fraction falls from 0.606 to a large D limit of 0.482. It is possible to formally find this limit by noting that for any value of Φ , Z'_b is sharply peaked about its maximum b_{mode} for large D . This allows the use of the saddle point approximation, whereby Z'_b is taken as Gaussian and therefore $f_\infty = b_{\text{mode}}/D$ by symmetry. Maximizing $\ln Z'_b$ with Stirling's approximation yields:

$$\frac{d \ln Z'_b}{db} \approx \ln \left(\frac{\Phi}{D} \right) + 2 \ln(D-b) - \ln(b), \quad (4.12)$$

$$\Rightarrow \frac{\Phi(D-b_{\text{mode}})^2}{D b_{\text{mode}}} = 1. \quad (4.13)$$

Using $b_{\text{mode}}/D = f_\infty$, Equations (4.13) and (4.8) are identical, as they should be.

The microscopic approach provides a simple mechanism for evaluating the accuracy of the correction scheme in certain cases. If it is possible to simulate the simultaneous formation of two or more targets, one can compare the change in dimer yield to the predictions of the microscopic approach, and then extend to the thermodynamic limit if the agreement is good. This is particularly useful if it is possible to consider an example with the relevant model where the self-assembly process is relatively simple. For example, for my DNA model, all interactions are truncated within distances much shorter than the typical separation of unbound strands, making the assumptions of ideality reasonable. Simulating duplex formation for short strands of about five bases in length is simple, and simulations forming several targets can be performed. The results are plotted in Figure 4.2 (b), showing perfect

agreement with Equation (4.11). Longer duplexes and complicated branched structures are much more challenging to simulate, meaning that only single target simulations are feasible. From the fact that the correction is successful for shorter duplexes, however, one can be confident that it will apply to longer strands when the concentration of DNA bases is similar.³ Throughout the rest of this work, I will use finite size corrections as detailed in this chapter to infer bulk binding properties of DNA systems.

4.1.3 Homodimer formation

It is instructive to consider the differences between homodimer and heterodimer corrections. For homodimers formed from two particles of type A, the partition functions of each particle species are given by:

$$q_{2A} = \frac{DZ_{2A}}{2}. \quad (4.14)$$

$$q_A q_A = D^2 Z_{A,A}, \quad (4.15)$$

where the factor of two compensates for the overcounting of indistinguishable states in Z_{2A} . Proceeding as in Section 4.1.1:

$$\frac{[2A]}{[A]^2} = \frac{vZ_{2A}}{2Z_{A,A}} = \frac{v\Phi}{2} = K_{2A}^{\text{eq}}. \quad (4.16)$$

The bound fraction in the thermodynamic limit follows as:

$$f_\infty = \left(1 + \frac{1}{4\Phi}\right) - \sqrt{\left(1 + \frac{1}{4\Phi}\right)^2 - 1}. \quad (4.17)$$

The behaviour of the correction is significantly different from that of heterodimers, as shown in Figure 4.2. In this case, the midpoint of the transition is unchanged, but the width is noticeably larger in bulk than for the two-particle system, i.e. $f_\infty > f_1$ for $f_1 < \frac{1}{2}$, and $f_\infty < f_1$ for $f_1 > \frac{1}{2}$.

³Due to the lack of non-specific attractions in my DNA model, it is unlikely that it will form large clusters unless the sequences are specifically designed for this purpose. This study shows that, neglecting the possibility of aggregation through base-pairing, the model's bulk behaviour is well described by the statistical extrapolation from a small system presented here. Bulk aggregation of DNA is possible, as discussed in Chapter 1, but in these examples the strands were specifically designed to form extended base-paired structures, and in other cases it can be neglected.

The physical mechanism for the broadening of the transition can be understood in terms of concentration fluctuations. Figure 4.1 (f) shows the states of a four-particle homodimer-forming system which cannot be sampled in a two-particle simulation. Of these, (f).i shows the smallest fluctuation in concentration, with three of the particles occupying half the volume and the remainder containing only one. In this case, it is impossible to have a binding fraction of unity. A binding fraction of zero is also less likely than in the two-particle case as the three monomers occupying the ‘right’ half of the system have a higher probability of forming one dimer than the two particles did in the original system. As a consequence, the fraction of dimers is pushed towards $\frac{1}{2}$ as the system grows in size, because larger concentration fluctuations are allowed which in turn favour the less probable configuration (whether dimer or monomer), leading to a broader transition in bulk.

The same argument can be applied to heterodimers, but with an important distinction. In this case, concentrations of individual species A and B can separately fluctuate. Unlike total concentration fluctuations, fluctuations in the relative concentrations of A and B will always reduce the probability of forming dimers, because of configurations like that in Figure 4.1 (c).iv where no dimers can be formed. Consequently the heterodimer yield is lower in bulk than for a two particle system, as well as having a broader transition.

4.2 Summary

In this chapter I have highlighted a necessity to correct equilibrium yields when extrapolating from small, canonical ensemble simulations of dimerization to bulk systems. I have provided a methodology to perform this correction and established a framework to explore its accuracy. I have tested the accuracy of the extrapolation for my model: agreement is essentially perfect, suggesting that, unless sequences are specifically designed to aggregate, small simulations are sufficient to extract the bulk thermodynamic properties of my model.

Dimerization in the canonical ensemble is of particular relevance to this thesis, especially in Chapter 6. In Appendix D, the discussion is extended to larger clusters, systems in the grand canonical ensemble and systems in which some of the reactants are localized.

Chapter 5

Structural and mechanical properties of model DNA

5.1 Basic structure

The model is specifically designed to allow an approximate representation of B-DNA in its double-stranded state. The relative sizes of the equilibrium backbone separation and ideal stacking distance lead to a pitch of 10.34 bp per turn at 296.15 K (23°C, approximately room temperature) similar to experimental estimates of 10–10.5 [12, 76]. The model length scale is chosen so that the average rise per bp at room temperature is 3.4 Å [128], which results in a helix with a radius (taken as the furthest extent of the excluded volume) of 11.5 Å, comparable to the experimental value of 11.5–12 Å [128, 161].

If strands are to form a double helix, it is not possible to optimize the stacking interaction, as consecutive stacking sites cannot sit directly above one another. Single strands, however, are not constrained in this way and hence form tighter helices, with a radius approximately 80% that of a duplex, similar to the 70–80% observed for a number of polynucleotide single helices [161]. A pleasing result is that, in order to alleviate the reduction in stacking, hydrogen-bonded bases in the model undergo ‘propellor twisting’ whereby bases in a pair twist in opposite directions in order to better align their stacking centres with adjacent bases in the same strand. Experimentally, propellor twist is seen to vary from around 5° to 15° in GC rich regions and from 15° to 25° in sections with large AT content [162]: my model has a slightly larger average propellor twist of 21.7° at 296.15 K.

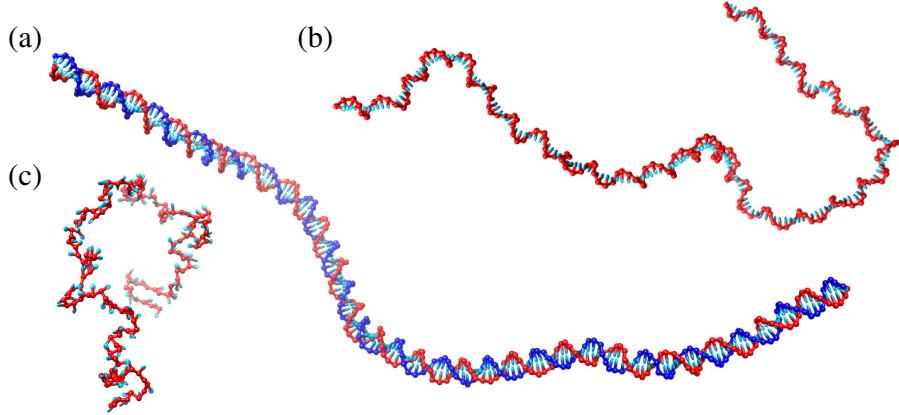


Figure 5.1: Typical configurations indicating relative flexibility of double-stranded, stacked single-stranded and unstacked single-stranded DNA. (a) 202 bp double helix at 296.15 K. (b) Stacked single strand of 202 bases at 277.15 K. (c) Unstacked single strand of 160 bases at 296.15 K.

5.2 Mechanical properties

It is important that the model captures the tendency of duplex DNA to be very stiff on the nanoscale, and the fact that ssDNA is comparatively flexible, if the model is to be used to study nanotechnology. As shown in Figure 5.1, this qualitative tendency is indeed reasonably well captured.

To make more quantitative statements, it is necessary to define metrics of rigidity. A commonly used measure of the large-scale properties of a polymer is its persistence length. A general definition is given, for example, in the textbook by Cantor and Schimmel [163]:

$$L_{ps} = \frac{\langle \mathbf{L} \cdot \mathbf{l}_0 \rangle}{\langle l_0 \rangle}, \quad (5.1)$$

with \mathbf{L} being the end to end vector of the polymer and \mathbf{l}_0 representing the vector between the first two monomers. For the case of an infinitely long, semi-flexible polymer in which the correlations in alignment decay exponentially with separation, Equation 5.1 is equivalent to the commonly used form:

$$\langle \mathbf{l}_n \cdot \mathbf{l}_0 \rangle = \exp(-n \langle l_0 \rangle / L_{ps}). \quad (5.2)$$

An alternative measure of polymer properties, the Kuhn length, is defined by [164]:

$$b_K = \langle L^2 \rangle / (L_{max}), \quad (5.3)$$

and gives the length of monomers for a freely-jointed chain (FJC) [164] with the same maximum end-to-end length L_{max} and $\langle L^2 \rangle$ as the polymer in question. For long semi-flexible chains $b_K = 2L_{ps}$ but for other models this equivalence may not hold.

5.2.1 Double-stranded DNA

Persistence-length

The persistence length of dsDNA is generally accepted to be approximately 450–500 nm at moderate to high $[Na^+]$, corresponding to around 130–150 base pairs [14, 165]. I performed four simulations of a duplex of length 202 bp at 296.15 K for 2×10^9 MC steps to extract the persistence length. It was found that base pairs at either end of single- or double-stranded DNA possess an increased relative flexibility. In order to obtain persistence length values that are valid for long strands where end effects are negligible, the behaviour of bases near the end of strands was ignored. The correlation of the helix axis (defined as the distance between consecutive base-pair midpoints) at two points was observed to decay exponentially with distance, allowing an estimate of L_{ps}^{duplex} through Equation 5.2. Figure 5.2 (a) indicates a model persistence length of around 123 base pairs, in reasonable agreement with experiment.

Torsional and extensional moduli

The stiffness of DNA duplexes is also manifested in a resistance to twisting. Torsional rigidity (in the linear regime) is quantified by an elastic modulus C , which relates applied torque G to resultant twist $\Delta\theta$ of a duplex of length l : $C = Gl/\Delta\theta$. Estimates for C have been made using cyclization kinetics and topoisomer distributions for minicircles [14, 166, 167], luminescence depolarization [168] and from twisting of DNA under tension [169], giving values in the range 170–440 fJ fm, with the effect of salt on C currently unclear from the literature [168].

Calculating the response to torsion is non-trivial, as the curvature of the DNA axis makes the twist between two ends hard to define. An approximate estimate of the torsional modulus can be obtained by considering the twisting of the central 10 base pairs of a 20 bp

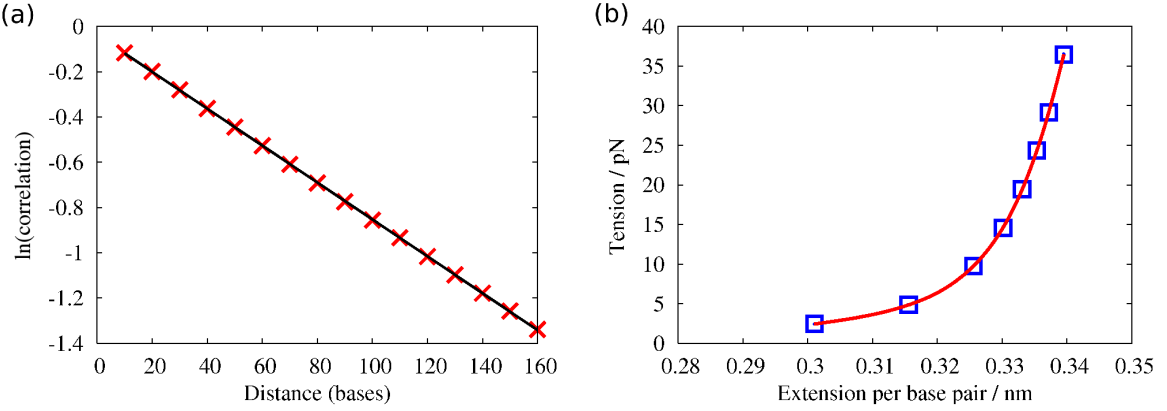


Figure 5.2: (a) Decay of the correlation of helix axis plotted against base pair separation for duplex at 296.15 K. The solid line is a fit of exponential behaviour. (b) Tension applied against extension for the central 100 bp of a 110-bp duplex at 296.15 K. The squares are simulation results, the solid line is a fit of Equation 5.4.

duplex, and the central 20 base pairs of a 30 bp duplex at 296.15 K. Such short sections are extremely stiff, minimizing the natural bending fluctuations. To provide an unambiguous definition of torsion and twist, Monte Carlo moves were chosen so that the base pairs at the end of the central section remained perpendicular to the vector between their midpoints, allowing the vector between the midpoints to define an axis about which torsion could be applied and twist measured.

Simulations were performed in which the torque applied to the end bases was varied between $\pm 8 \text{ pN nm}$, and the resultant twist used to infer C . A separate estimate was also obtained using the equipartition result for the variance in twist at zero torque: $\langle \Delta\theta_{\text{twist}}^2 \rangle = kTl/C$. Further simulations used the equipartition result to estimate C under a tension of 9 pN, to ensure that stretching the duplexes had no effect. All estimates (for both 10- and 20-bp regions of interest) gave $C \sim 450 - 475 \text{ fJ fm}$, suggesting that this is a reasonably robust estimate of the torsional stiffness of DNA duplexes in the model.

A long molecule of dsDNA under low tension responds as an extensible wormlike chain, with the behaviour initially dominated by the straightening of the chain, before stretching the base-pair rise itself becomes relevant as the chain extension approaches the contour length [170, 171]. At higher forces, the duplex undergoes an overstretching transition and the B-DNA structure breaks down [172]. Early experimental estimates for the extensional

modulus K , obtained from fitting force-extension curves to extensible wormlike chain models, give K in the region of 1050–1250 pN at high salt [170, 171], with smaller values at lower salt. A more recent study, at a monovalent salt concentration of 50 mM, claims $K \sim 1500$ pN [173].

The extensional modulus K was estimated by applying tension to a 100-bp region within a 110-bp double helix, and fitting the resultant force-extension curve to the result of Odijk for extensible wormlike chains [174]:

$$x = L_0 \left(1 + \frac{F}{K} - \frac{kT}{2F} [1 + y \coth y] \right), \quad (5.4)$$

where:

$$y = \left(\frac{FL_0^2}{L_{ps}kT} \right)^{1/2}, \quad (5.5)$$

in which x is the extension resulting from a force F applied to a duplex of contour length L_0 and persistence length L_{ps} . Performing an unconstrained three-parameter fit with the values of L_0 , L_{ps} and K gave an excellent agreement with the data, as shown in Figure 5.2 (b), with $K = 2166$ pN, $L_0 = 339.6$ Å and $L_{ps} = 431.0$ Å (126 bp). The value of L_0 is similar to that expected from the rise of a short duplex (exactly 3.4 Å per base pair would give $L_0 = 336.6$ Å), and L_{ps} is only slightly larger than the estimate from the decay of the correlation of the helix axis (123 bp). This agreement suggests that the extensible wormlike chain model provides a good description of the model's properties in this regime, and that the value of $K = 2165$ pN is a reasonably robust one for my model.

The model gives $C \approx 450 - 475$ fJ fm (slightly larger than the top of the experimental range of 170–440 fJ fm) and $K \approx 2165$ pN, (around ~ 1.5 to 2 times as large as typical experimental estimates). Although large, these values are not sufficiently different from measured values as to invalidate the majority of conclusions drawn from the model (although certain quantities, such as the critical twist density at which plectonemes are extruded, will be affected). It was found to be difficult to reparameterize the model to reduce these moduli without decreasing the persistence length, which is already slightly below experimental estimates. I feel that the current compromise, in which the persistence length is most

faithfully reproduced, is a reasonable one as it is easier to imagine that nanostructures and nanodevices would be more sensitive to bending than torsional or extensional stiffness.

It is worth noting that recent investigations have suggested that DNA initially overwinds when stretched [175]. The model does not reproduce this anti-intuitive behaviour, instead slightly untwisting as the stacking distance is extended. It is possible, therefore, that the model fails to capture the softness of a mode of deformation that leads to this behaviour – perhaps the sloping of base pairs with respect to the axis [176]. If this is the case, it is perhaps unsurprising that the estimated moduli are larger than experimental observations.

5.2.2 Single-stranded DNA

The mechanical properties of ssDNA are less well established than those of dsDNA. Furthermore, long single strands of DNA will generally be partially stacked, resulting in sections of substantially different flexibility [177, 178]. First, I shall explore persistence length of ssDNA when stacked.

Persistence length of stacked single strands

Mills *et al.* [177] have investigated the flexibility of gapped duplexes connected by poly(dA) (long single stands of DNA in which all the bases are adenine) at 4°C, when the bases are largely stacked into single helices. Although the interpretation depends on the probability of stacking, the intrinsic persistence length of the stacked regions was estimated to be around 100 Å, corresponding to approximately 30 bases. This value is noticeably larger than expected for unstacked strands (see Section 5.2.2), but smaller than for duplexes. For comparison, I performed four simulations of single strands of 202 identical bases at 4°C for 2×10^9 MC steps each (ignoring the data from the five bases at either end), requiring that all bases maintained a stacking interaction of $\geq -0.60 \text{ kcal mol}^{-1}$ with their neighbours (doubling this value had no discernible effect).

Taking the vectors between adjacent stacking sites as the axis of the polymer, Figure 5.3 shows that the correlation of these vectors decays exponentially with separation and hence that Equation 5.2 can be used to extract the persistence length of stacked ssDNA. Such a

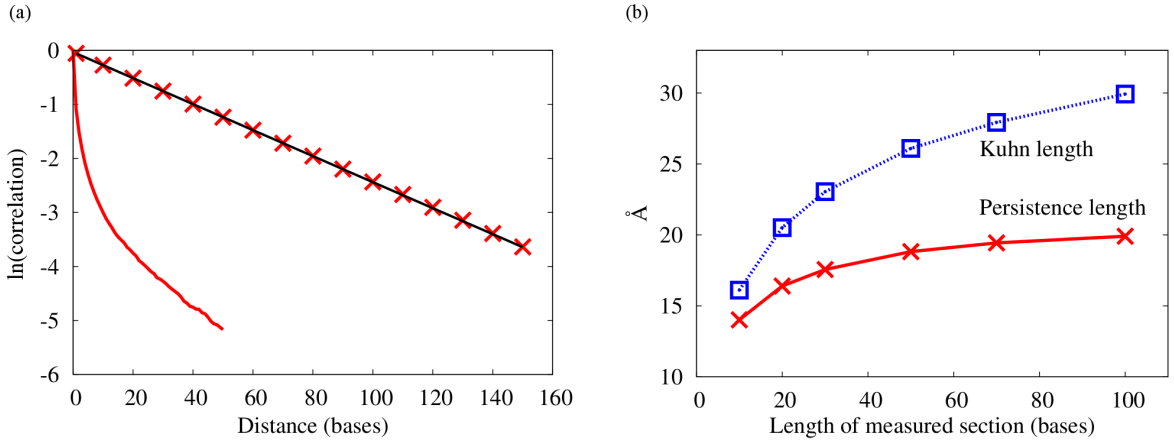


Figure 5.3: (a) Decay of the correlation of helix axis plotted against base separation for stacked single strands at 277.15 K (crosses), plotted logarithmically. The line is a fit of an exponential decay. Also shown (solid line, no symbols) is the decay of the correlation of backbone vectors for an unstacked single strand at 296.15 K. (b) Kuhn and persistence length inferred from simulations of unstacked single strands, as a function of the length of ssDNA over which they are measured.

fit gives $L_{ps}^{stack}/\langle l_0 \rangle = 41.6$ bases, somewhat larger than the value reported by Mills *et al.* [177], but not unreasonable.

Persistence length of unstacked single strands

Poly(dT) is generally assumed to be entirely unstacked at room temperature, and has little tendency to form secondary structure [12, 177]. As a consequence, it can be used to test the inherent flexibility of unstacked single strands. Gapped helices have been used by Mills *et al.* [177], who inferred a high salt persistence length of 20–30 Å from rotational decay rates, and Rivetti *et al.* [179], who studied length distributions with atomic force microscopy, finding ~ 16 Å for short sections (< 5 bases), growing to around 28 Å for longer regions. Fluorescence resonance energy transfer between donors and acceptors attached to either end of poly(dT) has also been used to fit polymer models to chain end-to-end distributions, with Murphy *et al.* finding a persistence length of around 19.4 Å at 500 mM $[\text{Na}^+]$ [129]. All of these results suggest persistence lengths on the scale of 2–5 bases.

In general, these results are obtained by first assuming a model of ssDNA (often a wormlike chain), and measuring a quantity which depends upon the conformational or dynamical properties of such a chain. The parameters of the model are then fitted to a

theory of the measured quantity which incorporates a polymer model. As such, the results are likely to be dependent on the choice of model for the polymer, and the accuracy of the theory used to interpret the data. As a consequence, one should treat these results with caution, but they do clearly indicate that unstacked ssDNA is very flexible.

When stacking and hydrogen-bonding interactions are removed from the model presented in this work, the conformation of ssDNA is exclusively determined by excluded volume interactions. Steric clashes limit backbone orientation, and this restriction on nearby nucleotides results in an effective stiffness of single strands. This description of ssDNA is undoubtedly a simplification, but it has the advantage of not biasing a particular structure and keeps all interactions as pairwise, which is ideal for the VMMC algorithm used in simulation.

To compare the model to experiment, I simulated single strands of one base type with stacking interactions set to zero to mimic poly(dT) (four simulations each for a range of strand lengths were performed at 296.15 K for 4×10^{10} VMMC steps. A typical configuration is shown in Figure 5.1(c)). As the stiffness of ssDNA in my model results from excluded volume interactions, meaning that interactions between non-neighbouring nucleotides play a role, it should not be a surprise that the correlation of backbone vectors does not decay exponentially with separation along the chain, as shown in Figure 5.3(a). It is therefore impossible to use Equation 5.2 to evaluate L_{ps} . If Equation 5.1 is used instead in order to compare to experimental results, the persistence length is observed to rise with the contour length considered (measured regions were embedded within strands with 30 extra bases at each end to avoid end effects), as shown in Figure 5.3(b). Similar behaviour is observed for the Kuhn length b_K , also shown in Figure 5.3(b). For strands of ~ 100 bases, the persistence length obtained using Equation 5.1 is similar to experimentally inferred values (19–30 Å), suggesting that the simplified description used in the model is not unreasonable.

Given that the experimentally inferred persistence lengths are so short, it seems plausible that non-neighbour interactions (which in physical DNA would include explicit electrostatic repulsion) may contribute towards single-stranded stiffness in physical DNA. Such interactions have been observed to dominate at lower salt concentrations in force-extension

experiments (see Section 5.2.2), when electrostatic interactions have a longer range, but they may also have a role at higher salt concentrations.

Force-extension properties

A number of groups have considered the stretching of ssDNA. In the work that first revealed the overstretching of dsDNA, Smith *et al.* [172] extended λ -phage ssDNA with optical tweezers. At moderately high salt (150 mM $[\text{Na}^+]$), the force-extension curve was well fitted by an extensible FJC model, with a Kuhn length of 15 Å and a stretch modulus of each Kuhn segment of 800 pN. At very low salt concentrations (and with formaldehyde present), the force-extension properties of the molecule were very similar to the higher salt results above 15 pN, but the DNA was much easier to extend at lower forces.

Even ignoring the failure of the FJC model to account for the behaviour in the second case, the use of the FJC in the first case is non-self-consistent. An assumption of the FJC model is that interactions between different segments can be neglected, but here the segment length (15 Å) is comparable both to the Debye screening length and the width of the polymer. In other words, if DNA really did want to behave like an FJC with segment length 15 Å, then at low forces it would find that it repeatedly bumped into itself and actually failed to behave like an FJC at all!

If the FJC model is not valid, one must explain why DNA in moderately high salt appears so hard to extend at low tension. Several authors have claimed that hairpins in the single strand, which must be unzipped if the molecule is to be extended, explain the original result of Smith *et al.* [172]. In the second case, formaldehyde and long-range electrostatic repulsion prevent the formation of hairpins, making ssDNA easier to extend. The long range repulsion, a result of low salt concentration, also favours extended conformations of a hairpin-free strand, making extension even easier.

To test the intrinsic force-extension behaviour of ssDNA, it is therefore necessary to prevent the formation of hairpins. Dessinges *et al.* considered stretching DNA in the presence of denaturants [180], and other authors have used homopolymers which are unlikely to form secondary structure [178, 181, 182, 183]. Both approaches, however, have their

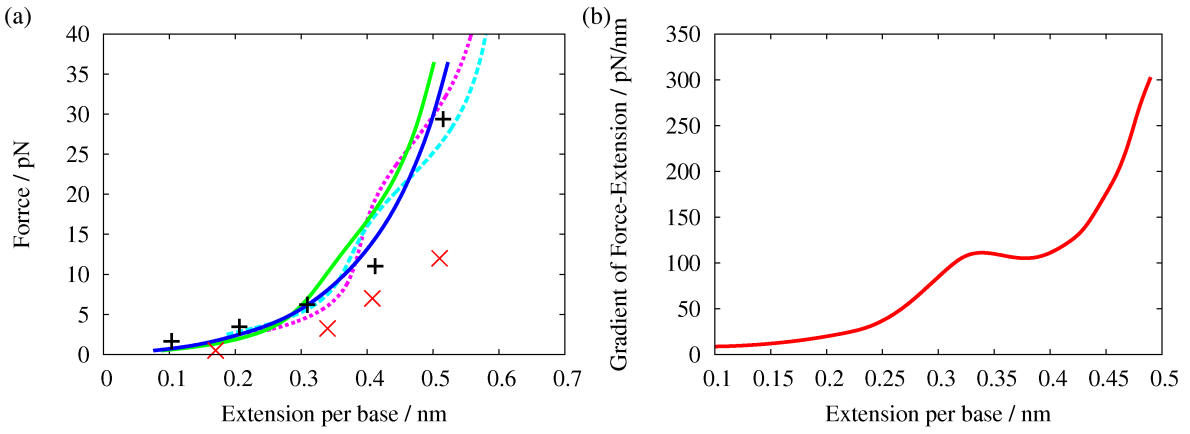


Figure 5.4: (a) Force - extension curves for model ssDNA compared to experiments. Solid curves represent simulations of model DNA: green for the normal parameterization, and blue without stacking. Dashed curves are inferred from the fits of Seol *et al.* [178]: magenta for poly(C) and cyan for poly(A). Black “+” are points from the force-extension curve of single-stranded λ -DNA in 150 mM $[\text{Na}^+]$, from Smith *et al.* [172]. Red “x” are points from the force-extension curve of charmoid DNA in low salt conditions and in the presence of denaturants, as measured by Dessinges *et al.* [180]. (b) Derivative of force-extension curve for model DNA when stacking is possible, calculated by differentiating a cubic spline fitted to the force-extension data. Note the plateau around at an extension of ~ 0.35 nm per base.

drawbacks: denaturants may have additional consequences beyond destabilizing the duplex, and homopolymers may behave in a manner which is not typical of a less ordered sequence.

Homopolymer stretching has revealed some non-trivial behaviour. When stretching ss-RNA (poly(A) and poly(C)), Seol *et al.* observed plateaus in the force-extension curve that were not seen for poly(U) [178, 181]. The authors were able to fit the curves using a statistical model in which the RNA was treated as a string of rigid, helically stacked sections and flexible, unstacked bases. Mishra *et al.* stretched homopolymeric ssDNA with an atomic force microscope, and found evidence of another plateau at higher forces (~ 110 pN) for poly(dA) that was absent for poly(dT) [182]. The existence of this plateau was confirmed by Chen *et al.*, who showed that it was associated with considerable hysteresis [183].

Testing the force-extension properties of model ssDNA is not difficult with the model – I simulated strands of 400 identical bases under constant tension, and extracted the average extension of the central 360 bases. Simulations were performed for 10^{10} VMMC steps and systems both with and without stacking interactions were considered. The results are plotted in Figure 5.4 (a), along with a number of experimental curves.

From Figure 5.4 (a) it is clear that the behaviour of model DNA is qualitatively rea-

sonable. It is difficult, however, to draw too many quantitative conclusions, as direct comparison to all curves is problematic. In particular, the curves of Seol *et al.* were obtained at the salt concentration appropriate to my model, but homopolymeric RNA may not give a good representation of generic DNA. Furthermore, the strand lengths in the experiment were unknown and only derived from fitting models to the data. The curve of Dessinges *et al.* was obtained in the presence of denaturants and at low salt, and hence the fact that ssDNA is far more extensible (particularly at low tension) is not surprising. The result of Smith *et al.*, as discussed above, shows the effect of hairpin formation (hence the resistance to extension at low force – the force at 0.1 nm/base is well above other curves). Above ~ 10 pN, however, hairpins should be of limited relevance and hence comparison may be worthwhile (although the salt concentration of 150 mM sodium is lower than that used to fit the model).

The effect of stacking

Seol *et al.* attributed plateaus in the force-extension curves of RNA to stacking in single strands [178]. They modelled ssRNA as a non-self-avoiding chain consisting of bases that could be stacked or random coil-like. Stacked regions had a shorter monomer length than unstacked sections, but a significantly larger persistence length. The assumptions of the statistical model used to fit the curves, at first sight, seems to apply fairly well to my system, and so it is worth considering my model in this light.

The first thing to note is that model strands are significantly easier to extend (above 10 pN) if stacking interactions are removed (Figure 5.4 (a)). This is indeed a result of the need to break stacking interactions in order to align more backbone vectors with the force. Furthermore, although it is not easy to see in Figure 5.4 (a), there is a plateau-like feature in the force-extension curve which is evident if the gradient is plotted (Figure 5.4 (b)). This feature becomes more obvious if the stacking strength is increased.

An obvious question is why my model shows a weaker plateau-like feature than the curves of Seol *et al.*, given that if anything the stacking interaction in my model is stronger than that used to fit the curves. The answer is that, in the model of Seol *et al.*, the separation

of stacked bases is assumed to lie along the helix axis, which tends to align with the force. This is a reasonable assumption for long helices, but for shorter sections with only a few bases, stacked regions will tend to rotate to align the longer vector between exterior sugars with the force. In my model, at high tension, single strands typically break up into a series of stacked pairs, which are capable of aligning their backbones with the applied force. By contrast, in the model of Seol *et al.*, the number of stacked bases tends to zero at high tension and this is why a relatively weak stacking interaction is able to provide such a strong signal.

Although my model's description of the backbone is simplistic, it is hard to see why the general principle that stacked bases can align their backbone separation, rather than their helix axis, with the applied force should be invalid. My model therefore suggests that the stacking parameters obtained from fitting the statistical model of Seol *et al.* to their data should be treated with some caution, although it supports the general principle that plateaus in force-extension curves can arise from the unstacking of helices.

5.3 Summary

Overall, the model gives a reasonably physical representation of the mechanical properties of DNA. Care must be used in systems when the specific values of moduli (rather than their order of magnitude) are of importance – for instance, the critical value of torque at which plectoneme extrusion occurs will be sensitive to the exact value of the torsional modulus. The model does, however, capture the tendency of dsDNA to be essentially rigid on the nanoscale, whilst single strands are extremely flexible – properties essential for much of DNA nanotechnology.

Chapter 6

Thermodynamic properties of model DNA

6.1 Single-stranded stacking transition

The attractive stacking interaction between adjacent bases causes single strands to form helical stacks at low temperature, with this order being disrupted as the temperature increases [12]. The literature is divided on both the nature of the attraction and the thermodynamics of the transition. The relative contributions of van der Waals, induced dipole, hydrophobic and permanent polar/electrostatic interactions remain unclear [144]. There has also been much debate on the cooperativity with which bases stack. Vesnaver and Bresslauer claim that a 13-base strand undergoes a completely cooperative transition between helical and random coil [184], whereas other authors have inferred essentially uncooperative transitions for the individual stacks in poly(C) and poly(A) [185, 186, 187, 188, 189]. Other groups claim weak to moderate cooperativity, with stacking probability affected by nearby base stacking [190, 191, 192]. It is clear, however, that stacking has a large influence on the thermodynamics of double helix formation, as the magnitude of the enthalpy and entropy changes of hybridization increase as the single-stranded state becomes more disordered [140, 184, 191, 193]. A limited number of short atomistic simulations of ssDNA have been performed [194, 195, 196, 197, 198]: all observe that the strands tend to adopt partially-stacked configurations, but there is insufficient data to draw firm conclusions about the thermodynamics of the process.

Given the uncertain nature of stacking behaviour it is difficult to constrain the model

in this regard. To introduce a large degree of cooperativity would, however, require adding internal degrees of freedom to the nucleotide or including next-nearest-neighbour interactions. For simplicity, therefore, I compare the model to reported uncooperative stacking. The study of Holbrook *et al.* [140] is most appropriate, as it deals with heterogeneous strands rather than homopolymers, and hence might be expected to provide a reasonable estimate of the average stacking strength.

To characterize the stacking properties of my model, I simulated oligonucleotides consisting of identical nucleotides (preventing the possibility of hydrogen bonding), and recorded the distribution of the number of neighbours with a stacking interaction stronger than a cut-off value¹ as a function of temperature and oligonucleotide length. For each strand length (5–9 and 14 bases), four simulations were performed at $T = 333\text{ K}$ for 4×10^9 VMMC simulation steps each, and I extrapolated the results to other temperatures using single histogram reweighting. For a 14-base nucleotide, around 50% of neighbours were found to be stacked at 338 K, with the transition being so broad that around 30% of neighbours remained stacked at 373 K, and 70% were stacked at around 306 K. Typical stacked and unstacked configurations are shown in Figure 6.1.

6.1.1 A statistical model of stacking

It is instructive to characterize the thermodynamics of stacking using a simpler, statistical model, as it highlights the causes of certain behaviour. I model the stacking transition using a statistical description based on that of Poland and Scheraga [199]. In this model, a given pair of neighbours can be either stacked or unstacked, and the list of stacked pairs specifies the system configuration.

If each stacking pair were independent and identical, the contribution to the partition function from a configuration (its relative probability of occurring) would be given by:

$$Z_{\text{config}} = z_0 u^{N_i} v^{N_j} \quad (6.1)$$

¹Bases were counted as stacked if their stacking interaction was less than -0.1 reduced units, or approximately $-0.60\text{ kcal mol}^{-1}$ (relative to a typical stacked interaction of -6 kcal mol^{-1}). Adjusting the cutoff to $-1.2\text{ kcal mol}^{-1}$ had a negligible effect.

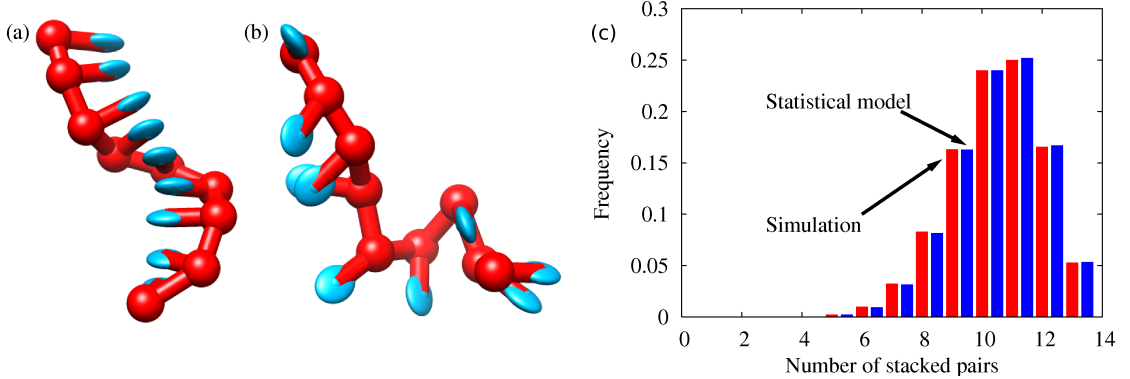


Figure 6.1: (a) and (b): two possible configurations of a 9-base strand at 333 K. (a) All neighbours stacked to form a right-handed helix. (b) Most neighbours unstacked, giving a flexible, disordered strand. (c) Frequency of total number of stacked bases in a 14-base single strand at 300 K from simulations of my model, and as predicted by the simpler statistical model with parameters as given in Equation 6.8.

where u and v represent the contributions to the partition function (statistical weight) of a stacked and an unstacked pair respectively, N_i and N_j are the number of stacked and unstacked pairs and z_0 denotes the trivial contribution from translation and orientation of the whole strand. To generalize to the case of non-independent neighbouring pairs, I introduce two new parameters. The statistical weight of a continuous section of n stacked pairs is now given by:

$$u(n) = \sigma u^n w^x, \quad (6.2)$$

with x being equal to the number of bases in the run of stacked pairs that lie at the end of the strand. n unstacked pairs contribute the same statistical weight as before:

$$v(n) = v^n. \quad (6.3)$$

If σ and w are unity, each neighbour pair is independent, and Equation 6.1 is recovered. σ takes the role of a cooperativity parameter: for $0 < \sigma < 1$, stacking is cooperative, in that configurations with multiple separate regions of stacking are disfavoured, and for $\sigma > 1$ stacking is anticooperative. w accounts for end effects: for $0 < w < 1$, end bases are less likely to stack: for $w > 1$ the opposite is true.

Using these definitions, the total partition function for a strand of length l becomes:

$$Z_l = \sum_{\{n_i, m_j; l\}} z_0 w^x \prod_i \sigma u^{n_i} \prod_j v^{m_j}. \quad (6.4)$$

Here, $\{n_i, r_j; l\}$ specifies a configuration, n_i being the number of stacked pairs in the i^{th} contiguous sequence of stacked neighbours, m_j being the number of unstacked pairs in the m^{th} sequence of unstacked bases and $x = \sum_i x_i$ is the total number of bases at the end of the strand involved in stacking.

Defining $t = u/v$, $n = \sum_i n_i$ and letting p be the total number of stacked regions:

$$Z_l = Z_l^u \sum_{\{n_i, m_j; l\}} w^x \sigma^p t^n. \quad (6.5)$$

with $Z_l^u = z_0 v^{l-1}$ being the partition function of a completely unstacked strand. To compare directly with simulations, the ratio of the probability of observing r stacked pairs to the probability of observing a completely unstacked strand is required:

$$\frac{Z_l(r)}{Z_l^u} = \sum_{\{n=r; l\}} w^x \sigma^p t^n = t^r \sum_{x=0}^2 w^x \sum_p \sigma^p \Omega_{\{x, r, p; l\}}, \quad (6.6)$$

with $\Omega_{\{x, r, p; l\}}$ defined as the number of distinct configurations of length l with r stacked pairs, of which x are at the end of the strand, divided between p contiguous regions of stacking. The advantage of this representation is that finding $\Omega_{\{x, r, p; l\}}$ is simply a matter of combinatorics.

$\Omega_{\{x, r, p; l\}}$ is given by the number of ways to split r stacked pairs into p sections, multiplied by the number of ways to split $l - r - 1$ unstacked pairs into $p + x - 1$ sections, with an additional factor of two for $x = 1$ as the order of stacked and unstacked regions can be swapped. This is equivalent to the number of ways of selecting $p - 1$ objects from a set of $r - 1$ objects (without replacement), multiplied by the number of ways to select $p + x - 2$ objects from a set of size $l - r - 2$. This is because there are $r - 1$ possible ways to divide the stacked pairs into non-empty sets, and we must do this division $p - 1$ times. Thus:

$$\Omega_{\{x, r, p; l\}} = \frac{(1 + \delta_{x,1})(r - 1)!(l - r - 2)!}{(r - p)!(p - 1)!(l - r - 2 - p + x)!(p - x)!}, \quad (6.7)$$

for all possible values of x , r and p for a strand of length l , with the exception that $\Omega_{\{0, 0, 0; l\}} = \Omega_{\{2, l-1, 1; l\}} = 1$.

I assume that the temperature dependence of stacking is manifested in the parameter t , which is defined as $t = \exp(-\Delta h^{st}/RT + \Delta s^{st}/R)$, with Δh^{st} and Δs^{st} representing the

(assumed constant) enthalpy and entropy changes associated with stack formation.² I take w and σ to be entropic (this will be justified later) and hence temperature independent.

The four parameter model detailed above can be fitted to simulation data, yielding:

$$\begin{aligned}\Delta h^{st} &= -5.47 \text{ kcal mol}^{-1}, \\ \Delta s^{st} &= -15.77 \text{ cal mol}^{-1} \text{ K}^{-1}, \\ \sigma &= 0.755, \\ w &= 0.789.\end{aligned}\tag{6.8}$$

As σ and w are close to unity, the model shows only weak cooperative and end effects. The entropy and enthalpy parameters are similar to those found by Holbrook *et al.* [140], to which the model's stacking behaviour was compared during fitting. The authors estimated $\Delta h^{st} = -5.7$ and $-5.3 \text{ kcal mol}^{-1}$ and $\Delta s^{st} = -16.0$ and $-15.0 \text{ cal mol}^{-1} \text{ K}^{-1}$ for two different strands at $[\text{Na}^+] = 120 \text{ mM}$. Similar results at $[\text{Na}^+] = 50 \text{ mM}$ suggest weak salt dependence in this regime [140].

Simulations performed in which the repulsive steric interactions were set to zero gave a slightly higher Δs^{st} and values of σ and w consistent with unity. Thus I conclude that the small cooperative effects in the model result from excluded volume. To understand the cause of the cooperativity, consider a chain of bases A , B , and C , and without loss of generality, consider B fixed whilst A and C move relative to it. Due to the requirement that base normals must point in the 3' to 5' direction to stack, the regions of space in which A and C stack with B do not overlap. Therefore, if A and B are stacked, the excluded volume that A represents to C only prevents C adopting conformations in which it is unstacked. By contrast, if A and B are unstacked, the excluded volume of A can prevent C adopting both stacked and unstacked configurations. As a consequence, C has a slightly higher tendency to stack if A and B are stacked, resulting in positive cooperativity. Similarly, end bases experience more freedom due to the reduction in excluded volume, and are therefore less likely to stack.

²Simulations are performed in the canonical ensemble, and hence should be described in terms of energy and entropy changes. I assume that, as dilute DNA strands contribute a very small partial pressure, discrepancies between constant volume and constant pressure results are small: we therefore use the term 'enthalpy' to describe what are in fact energies in the model, for consistency with experimental literature.

The statistical model is very successful. Figure 6.1 compares its predictions to the results for a strand length (14 bases) and temperature (300 K) that are well outside the ranges with which it was fitted.

6.2 Duplex formation

Hydrogen bonding between bases can lead to the formation of bound pairs of DNA strands, which adopt the canonical B-helix structure over a wide range of conditions due to stacking interactions. In contrast to the stacking transition, there is a reasonable consensus in the experimental literature on the melting temperature (T_m) of duplexes.

The model was fitted to the two-state model and parameters of Reference [91], which is known to give a very good prediction of experimental T_m . Note that the model is not limited to two-state thermodynamics, as will be shown in Section 6.2.2. Rather, I am using Reference [91] as a useful quantification of experimental results for melting. As my model contains no differentiation between A-T and G-C base pairs, I compare the results to strands consisting of ‘average bases’, the parameters for which, $\Delta h_{SL}^{step} = -8.2375 \text{ kcal mol}^{-1}$ and $\Delta s_{SL}^{step} = -20.019 \text{ cal mol}^{-1} \text{ K}^{-1}$, are obtained from averaging over all possible complementary base-pair steps in Reference [91]. I also use the average terminal corrections $\Delta h_{SL}^{term} = 1.1 \text{ kcal mol}^{-1}$ and $\Delta s_{SL}^{term} = 3.45 \text{ cal mol}^{-1} \text{ K}^{-1}$, the initiation parameters $\Delta h_{SL}^{init} = 0.2 \text{ kcal mol}^{-1}$ and $\Delta s_{SL}^{init} = -5.7 \text{ cal mol}^{-1} \text{ K}^{-1}$ and an additional salt correction of $\Delta s_{SL}^{salt} = -0.12754 \text{ cal mol}^{-1} \text{ K}^{-1}$ per phosphate for $[\text{Na}^+] = 500 \text{ mM}$, again taken from Reference [91].³

To explore the model’s representation of duplex hybridization, I simulated pairs of complementary oligonucleotides for a range of strand lengths between 5 and 20 bases. For each system, four simulations of 4×10^{10} VMMC steps were performed in periodic cells with a length of 20 simulation units,⁴ corresponding to a concentration of $3.36 \times 10^{-4} \text{ M}$. Umbrella

³Averaging over the parameters of Reference [91] gives an extremely convenient metric for comparison. An alternative approach (at least for the purposes of comparing T_m) would be to average over the T_m of all possible sequences. This second method gives results which are approximately 0.5 K lower for a 5 bp duplex and quickly converges on the first as duplex size increases.

⁴Simulations of duplexes with more than 12 bp necessitated using a larger periodic cell, and hence a lower concentration. The fraction of bound duplexes was scaled to the higher concentration assuming the

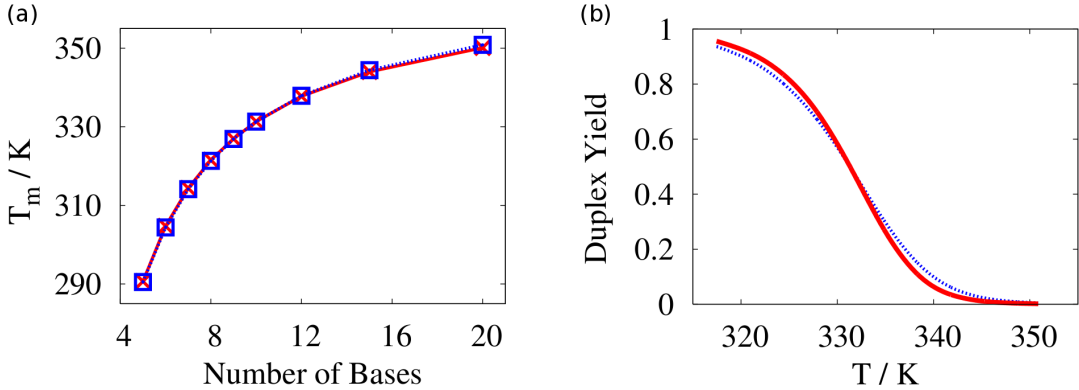


Figure 6.2: (a) T_m of duplexes as a function of strand length, with both strands having a total concentration of 3.3×10^{-4} M, as given by my model (crosses connected by a solid line) and averaged parameters from Reference [91] (squares connected by a dashed line). (b) Fraction of 10-base strands bound in duplexes at a concentration of 3.36×10^{-4} M as a function of temperature, from my model (solid line) and using the parameters of Reference [91] (dashed line). The typical range of the estimates of T_m from the four simulations at each duplex length is < 1 K, so errors are minimal on this scale.

sampling, using the number of base pairs with a hydrogen-bonding energy below -0.1 units (-0.60 kcal mol $^{-1}$) as an order parameter, was used to ensure good sampling. This cutoff is a factor of 7 lower than typical energies, and doubling it had no significant effect on the results. T_m was taken as the temperature at which half of the strands would be bound in a bulk solution.

The variation in melting temperature with duplex length is shown in Figure 6.2 (a), where it is compared to the predictions of the model of Reference [91]. The agreement in the dependence of T_m on length is extremely good: this dependence is essentially a measure of the cooperativity of the duplex forming transition, which is most strongly influenced by the relative contributions of hydrogen-bonding and stacking/cross-stacking to duplex stability.

The polynucleotide melting temperature (the melting temperature for infinitely long strands) at 500 mM [Na $^+$] for a strand of 50% C-G content, is predicted by the empirical relations given by Blake and Delcourt[200] and Frank-Kamenetskii[201] as 372.5 K and 369.0 K respectively. A crude estimate for my model can be made by simulating a pair of

separate species are approximately ideal, as justified in Chapter 4. Extrapolation to a range of temperatures was performed using single histogram re-weighting: the accuracy of such a method was checked for 8 bp duplexes, and no significant systematic errors were found.

long, complementary strands in a partially bound state, and finding the temperature at which the free-energy change of adding an additional base pair to a partially formed duplex is zero. Simulations of partially formed 100 bp strands (with the duplex/single-stranded DNA interface at a variety of points) gave values of T in the range 363.7 to 366.8 K, around 6 K below the empirical relations. The model's estimate, however, is very rough (as it neglects the stabilizing effect of bubbles and the destabilizing effect of intrastrand hairpins).

Figure 6.2 (b) compares the 10-bp duplex yield as a function of temperature for our model with the predictions of Reference [91]. The widths of the transitions are consistent to within a few degrees Kelvin, with my model consistently producing a marginally sharper transition for all duplex lengths. The width of the transition determines the response of the system to changes in concentration. Consider, for example, a simple two-state model of DNA hybridization, as used in Reference [91] and expressed in Equation 1.1. Assuming equal total concentrations of each strand ($[A_0]$), the width of the transition scales approximately as:

$$\Delta T \sim \frac{k_B T_m^2}{\Delta H}, \quad (6.9)$$

and the change in T_m with concentration is given by:

$$\frac{dT_m}{d[A_0]} = -\frac{k_B T_m^2}{[A_0] \Delta H} \sim \frac{\Delta T}{[A_0]}, \quad (6.10)$$

and hence agreement in both T_m and the transition width at a given concentration imply agreement in T_m over a range of concentrations.

6.2.1 Free energy profile of duplex formation and fraying

The free energy of duplex formation of a 15-bp duplex is plotted in Figure 7.3 as a function of the number of base pairs (the order parameter of umbrella sampling). To avoid complicating features in the free-energy profile due to hairpins and misbonds, which can conceal the underlying trends at low numbers of bonds, only base pairs that are present in the desired duplex were given a non-zero strength of hydrogen bonding in this simulation. The general form of the free-energy profile is qualitatively similar to that found for another coarse-grained model of DNA [121], with an initial entropy penalty for the formation of the first

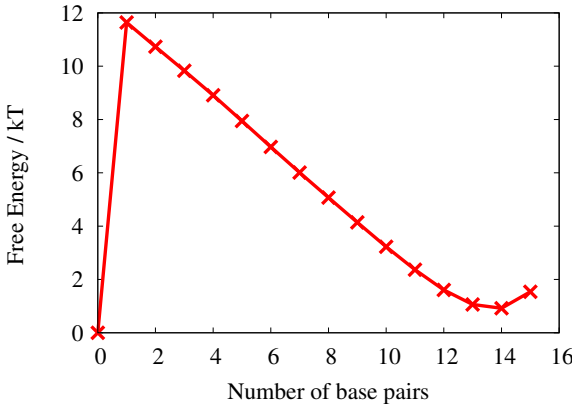


Figure 6.3: Free-energy profile of formation for a single 15-bp duplex from two strands in a cell of side length 30 simulation units, as a function of the number of base pairs, at 343 K.

base pair, followed by a downhill slope as the duplex ‘zips up’ in a cooperative fashion. As can be seen, the formation of the final base pair is actually free-energetically unfavourable, and the typical state consists of a duplex with ‘frayed’ ends. This fraying arises because bases at the end of the duplex lack the stabilizing influence of neighbouring base pairs on either side and entropy favours the open state.

Although fraying is a widely accepted phenomenon [202], experimental data is rather sparse, though it is established that weaker AT ends fray more easily than CG capped helices [203]. Nonin *et al.* inferred fraying probabilities of terminal AT bps of around 0.375 and 0.7 at 273 K and 298 K respectively, and found 0.015 and 0.12 for GC pairs at the same temperatures (at moderate salt concentrations) [203]. Patel *et al.* found much higher melting temperatures for terminal base AT pairs, concluding that they were around 50% frayed at 313 K at high salt concentration [204]. My model shows approximately 10% fraying at 273 K, increasing to around 20% at 300 K and reaching 50% at approximately 337 K, reasonable values for ‘average’ base pairs. It is worth noting that fraying probability for actual DNA will not only depend on the identity of the final base pair, but also its neighbour due to the sequence dependence of stacking interactions, an additional sequence-dependent effect that will not be reproduced by my model. Note that in many cases, particularly at low temperature, end bps in our model break but remain stacked, adopting conformations to maximize stacking at the expense of hydrogen bonding.

6.2.2 A statistical model of duplex formation

I attempted to fit the duplex yield as a function of temperature, for each strand length l , using a two-state model of the form in Equation 1.1.

$$\frac{[A_l B_l]}{[A_l][B_l]} = v \frac{Z_{ll}}{Z_l^2} = \exp \left(-\beta(\Delta H_l - T\Delta S_l) \right), \quad (6.11)$$

where $[A_l]$ is the concentration of strand A of length l and $[B_l]$ and $[A_l B_l]$ are the concentrations of its complementary strand and the bound pair. v is the volume simulated, Z_{ll} and Z_l are the statistical weights (contributions to the partition function) of the duplexes and single-stranded states to a simulation and ΔH_l and ΔS_l the (assumed T -independent) enthalpy and entropy of the transition. It was found, however, to be an unsatisfying fit to the melting curves, and further attempts to fit ΔH_l and ΔS_l as a linear function in l (by analogy with the nearest-neighbour model), were unsuccessful. The failure of a simple two-state model should not come as a surprise, however, as several authors have indicated that the entropy and enthalpy of duplex formation show temperature dependence due to the single-stranded stacking transition [140, 184, 191, 193].

Two-state descriptions typically fail when the macrostates that constitute the reactants and products show significant temperature dependence. In the rest of this section, I attempt to factor out the temperature dependence of reactant and product macrostates, thereby producing a multi-state statistical model that highlights the cause of the full model's description of duplex thermodynamics.

It is possible to devise a statistical model of duplex formation that takes the stacking transition into account. In particular, one can use the results of Section 6.1.1 to factor out the temperature dependence of the unbound state.

$$\frac{[A_l B_l]}{[A_l][B_l]} = v \frac{Z_{ll}}{Z_l^2} = v \frac{\exp \left(-\beta(\Delta H'_l - T\Delta S'_l) \right) (Z_l^u)^2}{Z_l^2}, \quad (6.12)$$

where in this case $\Delta H'_l$ and $\Delta S'_l$ are the enthalpy and entropy difference between the duplex and unstacked single-stranded macrostates. It might be expected that, having factored out the temperature dependence of the single-stranded state, $\Delta H'_l$ and $\Delta S'_l$ should be temperature independent. Unfortunately, although fitting to Equation 6.12 with constant $\Delta H'_l$ and

$\Delta S'_l$ was more successful than assuming constant ΔH_l and ΔS_l , it overcorrected for the variations in ΔS_l and ΔH_l with temperature. This failure arises primarily from neglecting the dependence of the bound state on temperature, which has two main contributions:

- As temperature increases, increased fraying leads to smaller entropy and enthalpy differences between typical bound states and completely unstacked single strands, as bound states become more disordered.
- Frayed ends themselves undergo a stacking transition, once more resulting in the entropy and enthalpy of bound states relative to unstacked strands becoming less negative with temperature.

To incorporate these effects within a statistical model, I split the duplex macrostate into states with y out of l possible base pairs formed. Stacking of the frayed ends is treated by viewing the $2(l - y)$ unpaired bases as undergoing stacking with the same Δh^{st} and Δs^{st} given in Section 6.1.1. Cooperativity and end effects are ignored for these bases as it would be difficult to include them consistently when stacking is initiated adjacent to a duplex region.

Therefore, defining $Z_{ll}(y)$ as the statistical weight of a duplex state with y out of l base pairs:

$$Z_{ll} = \sum_y Z_{ll}(y) = \sum_y Z_{ll}^u(y) \left(1 + \exp(-\beta(\Delta h^{st} - T\Delta s^{st}))\right)^{2(l-y)}. \quad (6.13)$$

Here $Z_{ll}^u(y)$ is the statistical weight of a duplex state with y out of l base pairs formed, and the other bases unstacked.

The hypothesis of the multi-state model is that the temperature dependence of the macrostates measured by $Z_{ll}^u(y)$ and Z_l^u should be minimal, and hence that:

$$v \frac{Z_{ll}^u(y)}{(Z_l^u)^2} = \exp\left(-\beta(\Delta H_l^0(y) - T\Delta S_l^0(y))\right), \quad (6.14)$$

with constant $\Delta H_l^0(y)$ and $\Delta S_l^0(y)$, which represent the enthalpy and entropy differences between unstacked single strands and the states contributing to $Z_{ll}^u(y)$.

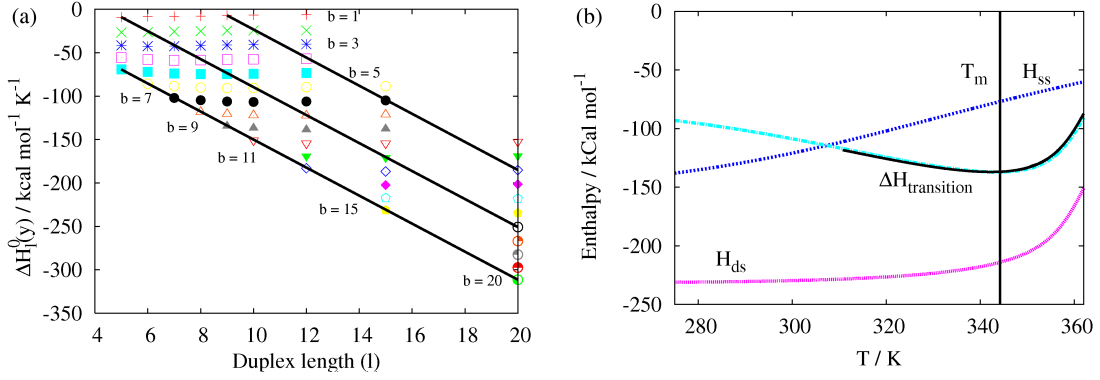


Figure 6.4: (a) $\Delta H_l^0(y)$ against duplex length l . Points are styled according to the total number of bonds formed, $b = l - y$. The solid lines are linear fits for the dependence of $\Delta H_l^0(y)$ on l for fixed y . (b) Variation with T of enthalpies associated with the formation of a 15-bp duplex. Solid lines represent simulation results, dashed lines the predictions of the statistical model outlined in the text. The lines labeled $\Delta H_{\text{transition}}$ give the enthalpy change upon duplex formation for the simulations and the statistical model. The lines labeled H_{ss} and H_{ds} are the enthalpies of the duplex and single strands relative to a completely unstacked state. The transition enthalpy in the statistical model is the difference between the latter two curves.

It is possible to extract the probability of forming l base pairs in a simulation of duplex formation, and the stacking parameters can be taken from Section 6.1.1. Hence $\Delta H_l^0(y)$ and $\Delta S_l^0(y)$ can be fitted via:

$$\begin{aligned} (\Delta H_l^0(y) - T\Delta S_l^0(y))/k_B T &= -\ln(v \frac{Z_l^u(y)}{(Z_l^u)^2}) \\ &= -\ln\left(\frac{v\Phi_l(y)(Z_l/Z_l^u)^2}{(1+\exp(-\beta(\Delta h^{st}-T\Delta s^{st})))^{2(l-y)}}\right), \end{aligned} \quad (6.15)$$

where $\Phi_l(y)$ is the relative probability (extracted from simulation) of being in a state with y base pairs compared to the probability of having zero base pairs. Temperature independent $\Delta H_l^0(y)$ and $\Delta S_l^0(y)$ fit the data extremely well, indicating that the statistical model successfully captures the temperature dependence of the bound and unbound states.

Furthermore, as shown in Figure 6.4, $\Delta H_l^0(y)$ (and $\Delta S_l^0(y)$, which is not shown) are linear in l for fixed y to an excellent approximation. Thus, having factored out sources of variation with temperature in the initial and final states, I obtain a result similar to the initial hypothesis of the nearest-neighbour model: adding an extra bp to a helix (i.e., increasing the length of the strands by one base, and forming one extra base pair, so that the number of unpaired bases is constant) contributes a constant enthalpy and entropy change relative to unstructured single strands.

The argument above suggests an extension of the nearest-neighbour model to non-two-state behaviour to incorporate fraying and stacking, and thus predict the values of $\Delta S(T)$ and $\Delta H(T)$ for oligonucleotides. Such a model, however, would require extensive parameterization (particularly when sequence dependence is considered), and may be impractical at the current time.

Interpreting duplex thermodynamics

Using the model described in the previous section, it is possible to describe the hybridization transition using completely temperature independent parameters. Combining Equations 6.6, 6.11, 6.13, and 6.14:

$$K_{eq} = \exp \left(-\beta (\Delta H_l - T \Delta S_l) \right) \\ = v \frac{Z_{ll}}{Z_l^2} = \frac{\sum_y \exp \left(-\beta (\Delta H_l^0(y) - T \Delta S_l^0(y)) \right) \left(1 + \exp \left(-\beta (\Delta h^{st} - T \Delta s^{st}) \right) \right)^{2(l-y)}}{\sum_r \exp \left(-\beta (\Delta h^{st} - T \Delta s^{st}) \right)^r \sum_x w^x \sum_p \sigma^p \Omega_{\{x,r,p;l\}}} \quad (6.16)$$

where K_{eq} is the equilibrium constant of the reaction.

$$\Delta H_l = -\frac{d}{d\beta} \ln K_{eq} \\ = \frac{\sum_y \left(\Delta H_l^0(y) + 2(l-y) \Delta h^{st} \frac{\exp(-\beta(\Delta h^{st} - T \Delta s^{st}))}{1 + \exp(-\beta(\Delta h^{st} - T \Delta s^{st}))} \right) Z_{ll}(y)}{Z_{ll}} - 2 \frac{\sum_r (r \Delta h_{st} Z_l(r))}{Z_l}. \quad (6.17)$$

The enthalpy changes at T_m for the model are slightly larger than predicted by Reference [91], which is to be expected as the transitions are slightly narrower. The discrepancy rises from about 6% for 5 bp duplexes to around 22% for 20 bp double strands. The behaviour of ΔS is similar.

To investigate the details of the temperature dependence of enthalpy changes in duplex formation, I simulated the formation of a 15 bp duplex over a wide range of temperatures (to avoid complications due to misbonds and hairpins, only ‘correct’ pairs were given an attractive hydrogen bonding interaction), with the data shown in Figure 6.4 (b). I find that at low temperatures, below 342 K, ΔH becomes more negative with increasing temperature, with a gradient that reaches a maximum size of around $-0.055 \text{ kcal mol}^{-1} \text{ K}^{-1}$ per base pair at approximately 312 K. At 342 K, however, ΔH reaches its most negative value, before

heading rapidly towards zero. The statistical model highlights the cause of this behaviour, as illustrated in Figure 6.4(b).

Well below 340 K, the enthalpy of the bound state is seen to be approximately constant whereas the enthalpy of the single strands becomes less negative with increased temperature as they unstack, causing the observed tendency for ΔH of the transition to become more negative. At higher temperatures, however, the enthalpy of the bound state becomes less negative due to fraying, as the typical bound state changes from being a fully formed duplex at low temperatures to a higher enthalpy partially-melted state at higher temperatures. As the polynucleotide melting temperature is approached, fraying becomes more significant, resulting in the observed decrease in the magnitude of ΔH for the transition.

A change in enthalpy due to the stacking transition has been observed experimentally by several groups [140, 184, 191, 193, 205], who deduced values for the typical enthalpy gradient of -0.050 , -0.095 , -0.05 to -0.1 , -0.068 to -0.87 and $-0.062 \text{ kcal mol}^{-1} \text{ K}^{-1}$ per base pair, respectively, in reasonable agreement with the model. These investigations were generally performed with either oligonucleotides with several CG pairs at the end [140, 184, 191, 193] or polynucleotides [205], both of which would massively reduce the impact of fraying: if I set the fraying contribution to zero in the statistical fit, I obtain a typical value of -0.06 to $-0.07 \text{ kcal mol}^{-1} \text{ K}^{-1}$, in even better agreement with experiment.

In addition, Jelesarov *et. al.* [193] considered another duplex with AT bps at the end of the helix, which showed ΔH becoming more negative with increasing T at low temperature, before flattening-off by around 320 K, in agreement with the predictions of the model for the consequences of fraying. Measurements were not performed at high enough T to check for an eventual decrease in the magnitude of ΔH , but my model predicts the effect should be observable. In particular, duplexes with large AT end regions and a stabilizing GC cores should demonstrate such an effect.

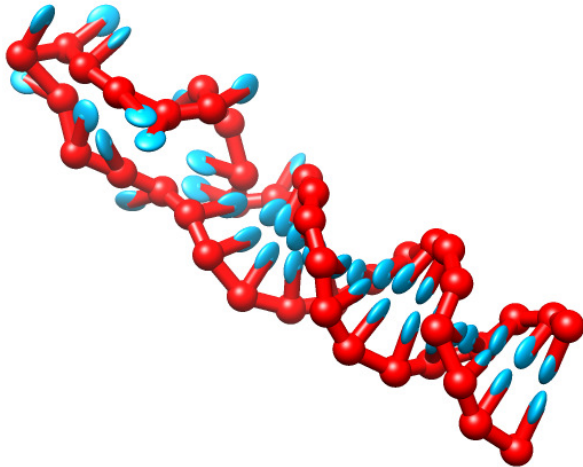


Figure 6.5: A hairpin with a 12 bp stem and an 18-base loop at 343 K.

6.2.3 Structural motifs

Hairpins

DNA hairpins, which occur when a self-complementary strand binds to itself, forming a duplex stem and an unhybridized loop (Figure 6.5), are a common structural motif. They have biological importance as a mechanism for release of superhelicity through cruciform formation [76], and relevance to nanotechnology as metastable states (either occurring by accident [206] or through design [66, 130]). In addition, they are an extremely common motif in biological RNA structures [207]. To date, there have been no coarse-grained models which have been applied to both hairpins and bimolecular duplexes. An approach in which the single strands have the potential to be extremely flexible allows for hairpins and duplexes to have appropriate relative stabilities.

To demonstrate the ability of the model to represent hairpins, I simulated them with stem sizes ranging from 5–12 bps, and loops of 5–18 bases. Four simulations for each hairpin were performed in the vicinity of T_m for 4×10^{10} VMMC steps, with umbrella sampling as a function of hydrogen-bonded base pairs used to ensure good statistics. In this case, I considered only states with at least one of the ‘native’ bps in the stem present as being a hairpin, as long loops had the potential to form transient base pairs with little relevance to the stability of the target structure. SantaLucia has presented parameters for estimating the

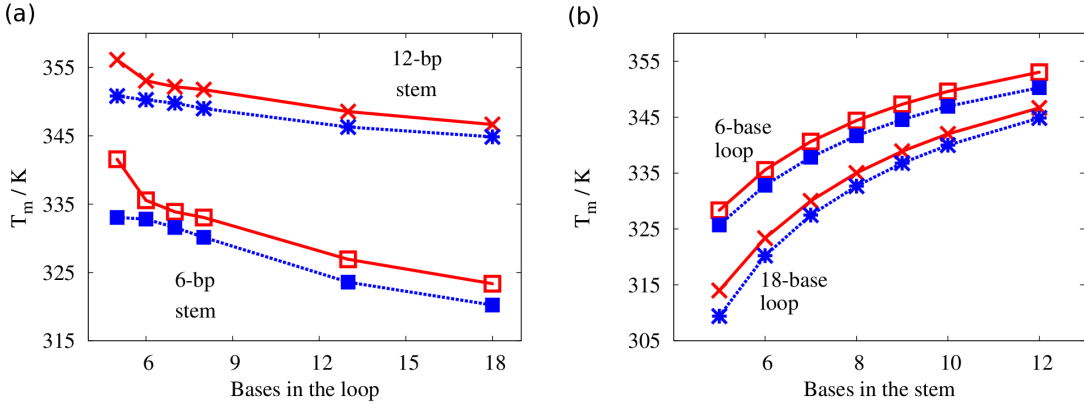


Figure 6.6: (a) Variation of hairpin melting temperature with loop length from our model (symbols connected by dashed lines) and from Reference [91] (solid lines). (b) Variation of hairpin melting temperature with stem length from our model (symbols connected by dashed lines) and from Reference [91] (solid lines).

melting temperature of hairpins [91], which I again take as a good representation of experimental results. These parameters include sequence independent entropy penalties for loop formation and enthalpy/entropy terms for the stabilizing effect of the first mismatched bp in the loop (called a ‘terminal mismatch’: I compare to an average $\Delta h_{SL}^{term} = -2.91 \text{ kcal mol}^{-1}$ and $\Delta s_{SL}^{term} = -7.33 \text{ cal mol}^{-1} \text{ K}^{-1}$). The results for T_m are compared to the predictions of Reference [91] in Figures 6.6 (a) and (b). T_m is defined as the temperature at which a strand is in a hairpin state half of the time.

The results indicate that the model slightly underestimates T_m for hairpins relative to the predictions of Reference [91] (and by extension, experiment): typically by approximately 3 K, which is slightly less than 1% of the absolute melting temperature (at the T_m predicted by Reference [91], model hairpins usually constitute approximately 25% of the ensemble rather than 50%). My model fails to reproduce the jump in stability associated with 5-base loops (and indeed smaller ones, which are not shown), and it is difficult to see how such an effect could be captured without specific stabilizing interactions (as it deviates strongly from the trend for longer loops). Such interactions have been postulated for tightly packed loops [208], and for 3- and 4-base loops Reference [91] includes loop sequence specific corrections, presumably for this reason.

Encouragingly, the trends with loop length (for loops longer than five bases) and stem

size are well reflected by the model (this is particularly pleasing, as the dependence on loop length was not used in parameterization), an indication that the majority of the physics of hairpin formation is well represented by the model. As found with duplex formation, model transition widths are slightly smaller than predicted by Reference [91] (the discrepancy is very similar to that indicated in Figure 6.2 (b)). Note that, if the duplex transitions were as broad as predicted by the SantaLucia model, hairpin stem formation would be somewhat more favourable at these temperatures well above the duplex T_m . The slightly narrower transitions of my model might therefore contribute to the lowering of hairpin T_m relative to Reference [91], as well as the representation of the loop itself.⁵

Mismatches, bulges and internal bubbles

A variety of other DNA motifs exist, such as duplexes involving mismatches between non-complementary base pairs or with one strand carrying extra, unpaired bases. SantaLucia [91] has provided parameters for the influence of these motifs on T_m . In many cases, they are highly sequence dependent and it is less clear than in the simple double-helix case (where the variations in parameters are relatively smaller) that averaging over ΔS and ΔH contributions for all sequences is a reasonable approach to find an average effect. It should, however, give a rough estimate of the typical change in melting temperature due to a motif.

I compared the effect of several motifs on model duplex T_m to the predictions of Reference [91], again averaged over all possible sequences (Table 6.1). The simplest possible case is that of a single unpaired base at the end of a strand, generally referred to as a ‘dangling end’. Typically, dangling ends are observed to provide a stabilizing influence, assumed to result from cross-stacking with the final base pair of the duplex, although the degree of stabilization is highly sequence dependent [91, 144]. The cross-stacking interaction included in the model provides such a stabilizing effect, and the degree of stabilization is in good agreement with the predictions of Reference [91]. In contrast to dangling ends, extra,

⁵For example, if the melting transition of an 8-bp duplex was as wide as predicted by Reference [91], one would expect the statistical weight of an 8-bp duplex in my model to be increased by around a factor of two at 341 K. This is the approximate melting temperature of a hairpin with an 8-bp stem and a 6-base loop: increasing the statistical weight of the hairpin by a factor of two would translate into an increase in T_m of around 2 K.

Motif	Complementary bp	Motif size	ΔT_m / K	
			My Model	Ref. [91]
Dangling end	5	1 base	+3.42	+4.24
	8	1 base	+1.33	+1.44
	15	1 base	+0.66	+0.70
Bulge	8	1 base	-17.98	-23.19
		2 bases	-23.92	-26.73
	15	1 base	-8.18	-12.36
		2 bases	-11.03	-11.58
		5 bases	-15.97	-13.11
Terminal mismatch	5	1 base / strand	+6.71	+8.60
	8	1 base / strand	+3.02	+2.71
	15	1 base / strand	+0.94	+0.31
Internal mismatch / bubble	8	1 base / strand	-8.81	-13.99
		2 bases / strand	-15.77	-21.86
		5 bases / strand	-25.70	-28.81
	15	1 base / strand	-4.92	-4.88
		2 bases / strand	-9.18	-11.51
		5 bases / strand	-15.37	-15.65

Table 6.1: Effect on the melting temperature of a complementary duplex due to the addition of a motif. In this table, ΔT_m is the difference between the T_m of a structure with the motif and a fully complementary duplex consisting of the same number of complementary bps as the motif structure. For internal mismatches, bulges and bubbles, the motif was placed at the centre of the duplex.

unpaired bases on one strand within the helix are highly destabilizing, as they disrupt the helix structure. In the terminology of SantaLucia, these are known as bulges. In general, our model slightly underestimates the destabilization of helices due to bulges compared to the predictions of Reference [91], although the melting observed temperatures remain within 2% of the predictions.

If a non-complementary pair of bases is added to an otherwise complementary duplex to form a mismatch, the effect is generally stabilizing at the end of a duplex (this is a ‘terminal mismatch’) and destabilizing in the interior. The model reproduces this tendency as shown in Table 6.1, and also captures the increase in destabilization if the mismatch region is extended (to form an internal ‘bubble’). Once again, the destabilizing effect of motifs

internal to the duplex tend to be slightly underestimated relative to the predictions of Reference [91], but the observed melting temperatures remain within 2% of the predictions.

Such motifs provide a good test of the model, as many were not considered in parameterization (although the dangling ends and terminal mismatches were used to constrain the strength of cross-stacking). In addition, misbonded structures involving these motifs may have a role in the kinetics of nanostructure assembly, and hence it is important that the model provides a reasonable representation of them. Although in some cases the quantitative agreement with Reference [91] is not perfect, the model represents these motifs in a physically sensible way and the trends in stability at least qualitatively reflect the average properties of DNA. Furthermore, the typical magnitudes of ΔT_m are reasonable, with the T_m remaining within 2% of the average predictions of Reference [91]. It is possible that an underestimate of the disruptive effect of extra bases on the helical structure [76], perhaps because the excluded volume of bases is smaller than in reality, causes the underestimate of ΔT_m due to internal motifs. Another consideration may be that as the model hybridization transition is slightly narrower than predicted by Reference [91], a given destabilizing ΔG will have a smaller effect on model T_m than for the statistical description of SantaLucia.

Given the good agreement between the model and Reference [91] for a single mismatch added to a 15-bp duplex, I investigated how the position of the mismatch affected stability. T_m is plotted against the position of the mismatch in Figure 6.7 (a). As can be seen, there are two distinct regimes, with the melting temperature initially decreasing as the mismatch is moved from the end of the strand (where it is stabilizing) towards the centre. Eventually, however, it reaches a plateau at around five bases from the end of the strand.

The cause of this plateau can be identified from examining the free energy profiles for duplexes with mismatches located one and six bp from the end (Figure 6.7 (b)). The first point to note is that the stability of duplexes with the maximum number of base pairs (15) is nearly identical, despite the difference in mismatch position. This suggests that provided a mismatch is surrounded by base pairs on either side, changing its location has little effect on the total free energy. The difference in T_m arises instead from a difference in the nature of the lowest free-energy state.

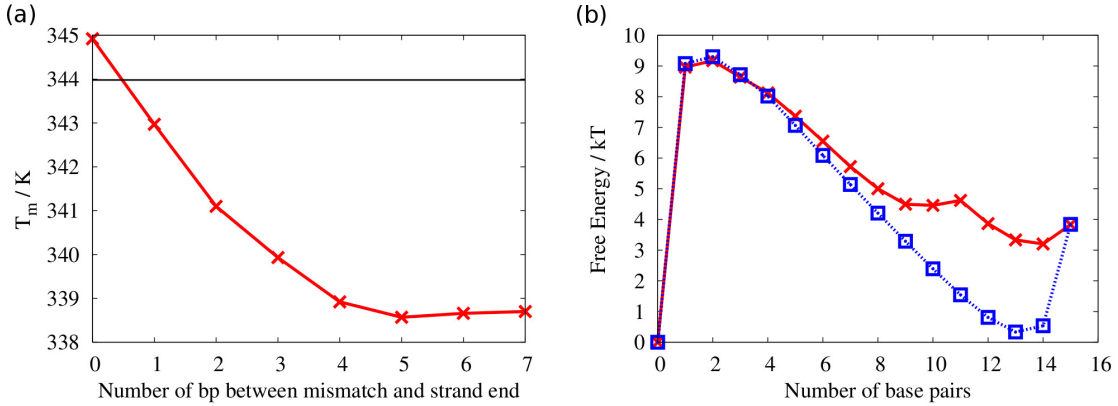


Figure 6.7: (a) Melting temperature of 15-bp complementary helix with an additional mismatch added against the distance of that mismatch from the end of the strand. The melting temperature in the absence of a mismatch is indicated via the horizontal line. (b) Free energy profile at 339 K for a 15-base-pair duplex with one additional mismatch placed 1 base from the end (squares) and 6 bases from the end (crosses)

When the mismatch is near to the strand end (in the regime where T_m depends on mismatch position), the most stable state consists of the larger section of duplex formed with the bases beyond the mismatch unpaired. In this regime, the total free energy gain from pairing the bases beyond the mismatch does not compensate for the cost of enclosing a mismatch in a helix. As the mismatch is moved towards the centre, the larger section loses bases and so becomes less stable, with the consequence that T_m drops. At some point, however, the possible number of base pairs in the region beyond the mismatch makes it favourable for the base pairs in this region to form. From this point onwards, the most stable state consists of the two duplex regions surrounding the mismatch. When this occurs, the net effect of moving the mismatch towards the centre can be viewed (in the average base-pair case) as the transfer of a base pair from the centre of the larger duplex to the centre of the smaller duplex, where the contribution to stability will be unchanged. As a result a plateau in T_m is observed.

As the temperature is lowered, the free-energy gain from base pair formation increases. As a consequence, the number of bases required before the region beyond the mismatch is stable as a duplex decreases. For example, I find that for a mismatch two bases from the end of a duplex, the enclosed mismatch state becomes the most stable just below 320 K.

It is claimed in Reference [91] that the stability of a mismatch is independent of its

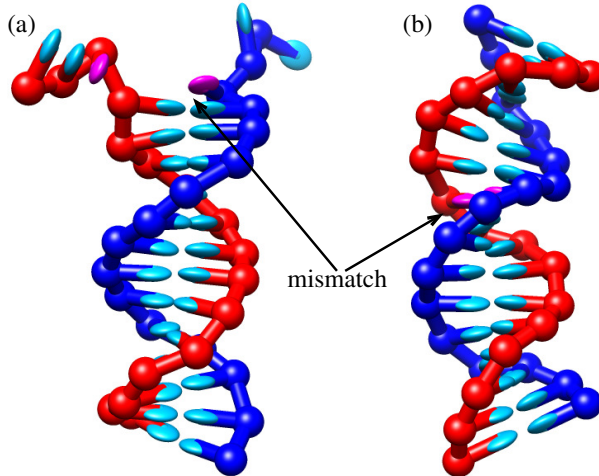


Figure 6.8: Typical configurations of a duplex with 15 complementary bp and 1 internal mismatch at 335 K. a) Mismatch two bp from the end of the strands, with unpaired bases after the mismatch. b) Mismatch six bp from the end of the strand, enclosed by two intact helices.

position, except for terminal mismatches and mismatches occurring one base from the end, which may cause the final base pair to be unstable. My simulations suggest, however, that the distance of the mismatch from the duplex end at which T_m plateaus should increase with strand length (as longer strands melt at higher temperature). Further, a similar temperature dependent influence of motif location should hold for all destabilizing internal bubbles and bulges, as the plateau simply indicates the point at which it is free-energetically favourable to form the second helix beyond the disruption.

A dependence of T_m on mismatch location has been reported in the DNA microarray literature [209, 210]. In particular, You *et al.* [209] observed that 21 bp duplexes had their melting temperatures reduced by 1.8-2.2 K due to a mismatch 3 bp from the end, and by 3.5 K due to a central mismatch, values comparable to those found here (although slightly smaller, as would be expected from a longer duplex with a narrower transition). Naiser *et al.* [210] have even attempted to explain the effect of mismatch location by using the model of Reference [91] to predict the relative population of partially bound states, in an approach similar to that of the Zuker group [211, 212]. This methodology, however, is somewhat suspect – the SantaLucia parameters are fitted to reproduce the free-energy difference between the bound and unbound ensembles, and hence partially-formed states are already included implicitly in the fitting. Using the parameters to separately evaluate

the free energy of partially bound states therefore represents a form of double-counting.

My model therefore reproduces the observed tendency of mismatch location to influence melting temperature, and even supports a hypothesis for the cause of this non-trivial effect. This is particularly pleasing, as the model was in no way parameterized for this purpose.

6.2.4 Coaxial stacking

Coaxial stacking occurs when non-neighbouring bases (often from different strands) stack with each other. Such an interaction has been used to explain the stabilization of duplexes by adjacent hairpin stems [91, 135, 136, 137, 138, 139] and the association of DNA origami tiles [42].

The strength of the coaxial stacking interaction has been measured experimentally in two ways. The Frank-Kamenetskii group have attempted to infer the probability that a nicked duplex is in a state with stacking across the nick from gel mobility studies [142, 143]. Other authors have considered the thermodynamic stabilization of duplexes due to the presence of an adjacent double-stranded region, with which coaxial stacking can occur [91, 135, 136, 137, 138, 139].

The direct measurements of the coaxial stacking/unstacking equilibrium from the Frank-Kamenetskii lab were performed by measuring the mobility of duplexes undergoing gel electrophoresis. The mobilities of duplexes with a nick in one of the backbones were compared to the same duplexes without a nick and gapped duplexes containing two single G bases. Assuming the coaxially stacked state of nicked duplexes had the same mobility as the unnickled duplexes, and that the unstacked state had the same mobility as the gapped duplexes, the authors were able to infer a probability of DNA being in each configuration.⁶

Initial experiments [142] were performed at 37°C and low monovalent salt concentrations (around 15 Mm Na⁺). These results give $\Delta G_{\text{coax}} \approx -0.3$ to $-3.5 kT$ for a range of sequences, where $e^{-\Delta G_{\text{coax}}/kT}$ is the ratio of occupancy of stacked and unstacked configurations. A noticeable trend is that stacks of CG base pairs seem to be much stronger. Later

⁶Note that the experiments were performed in a range of concentrations of denaturing urea, then extrapolated to zero urea concentration.

work explored higher salt concentrations, with ΔG_{coax} typically becoming more negative by around $0.65 kT$ between 15 mM and 100 mM $[\text{Na}^+]$, giving an average of around $-2.62 kT$ at 100 mM $[\text{Na}^+]$ [143].

A number of other groups have tried to estimate the stabilizing contribution to the free energy of hybridization, which I shall call $\Delta\Delta G_{\text{hybr.}}$, from the presence of an adjacent duplex. The Santalucia lab has inferred $\Delta\Delta G_{\text{hybr.}}$ from analyzing the formation hybridization of six-base strands to the dangling ends of hairpins at high salt, finding an average stabilization of around $-2.74 kT$ at 37°C (although there is strong sequence dependence). Pyshnyi and Ivanova considered 7 bp duplexes stabilized by hairpins at high salt, and inferred an average $\Delta\Delta G_{\text{hybr.}}$ of around $-2.62 kT$ at approximately 37°C , noting not only dependence on the bases at the coaxial stack site, but also on the next-nearest neighbours [136, 137]. Lane *et al.* measured the hybridization of a 13-base strand to the dangling end of a hairpin at 115 mM monovalent salt, comparing cases in which the duplex was adjacent to the hairpin stem and in which there was a two nucleotide gap. The first duplex was found to be more stable by about $-1.87 kT$ at around 325 K – as the authors point out, however, the gapped system is stabilized by dangling ends and hence this value is expected to be somewhat smaller than the true $\Delta\Delta G_{\text{hybr.}}$. Finally, Vasiliskov *et al.* considered the hybridization of a 5-base strand to the overhanging end of a duplex that was tethered to surface in a microarray [139]. The results are difficult to draw direct comparisons with, as the system may be quite distinct from solution-based assays, but the authors report that stabilization is much stronger if adenine is involved in the coaxial stacking.

It is tempting to draw direct comparisons between $\Delta\Delta G_{\text{hybr.}}$ and ΔG_{coax} , but they would be fallacious, as the quantities represent free energy differences between different pairs of macrostates. For my model, they are quite different, and ΔG_{coax} is significantly larger than $\Delta\Delta G_{\text{hybr.}}$. The main reason for the difference is that single-stranded dangling ends can be stacked onto the end of a duplex, and indeed one would expect them typically to be stacked as dangling ends tend to stabilize duplexes. This stacking of dangling ends tends to reduce the value of $\Delta\Delta G_{\text{hybr.}}$, as it makes the unbound state more favourable.

Modelling coaxial stacking

The coaxial stacking interaction in my model, described in Chapter 2, is based on the nearest-neighbour stacking interaction, with several differences. In particular, I have allowed the strength of the interaction to be different from the nearest-neighbour term – given that the stacking interaction implicitly incorporates a host of backbone effects, it is not surprising that this is necessary. As the interaction is only important in certain situations, the model was initially parameterized without it [213], and it has since been included with the interaction strength as a fitting parameter.

To compare the resulting interaction to experiment, I have simulated a system analogous to that studied by the Frank-Kamenetskii group. Specifically, I have measured the probability that a 20 base duplex with a nick in the centre unstacks about the nick site, at 37°C, finding $\Delta G_{\text{coax}} = -4.3 kT$. This is somewhat more negative than the Frank-Kamenetskii group’s average value, even if the salt dependence is extrapolated to $[\text{Na}^+] = 500 \text{ Mm}$.

I have also attempted to compare directly to the thermodynamic measurements of $\Delta\Delta G_{\text{hybr.}}$. In particular, I simulated 6-, 7- and 8-bp duplex formation adjacent to a hairpin stem⁷. The duplex melting temperatures were raised by 9.88 K, 7.66 K and 6.25 K respectively by the presence of the hairpin stem. By comparison, the averaged Santalucia parameters for coaxial stacking predict slightly larger stabilizations of 11.7 K, 8.5 K and 6.4 K. The results of Pyshnyi and Ivanova for $\Delta\Delta G_{\text{hybr.}}$ are similar on average to those of the SantaLucia lab, and hence give similar stabilizations if they are used in conjunction with the remaining Santalucia nearest-neighbour parameters. The actual value of $\Delta\Delta G_{\text{hybr.}}$ at 37°C is given by my model as $-2.43 kT$, slightly less negative than the estimates of References [136], [137] and [141]. Note also the difference between $\Delta\Delta G_{\text{hybr.}}$ and ΔG_{coax} in my model, as discussed above.

There are issues with all of these comparisons. The thermodynamic results are limited, as References [136], [137] and [141] considered systems with fairly low T_m , and the predictive power of their parameterizations have not been tested (particularly for systems with a higher

⁷Hairpin stems contained 12 bp and loops consisted of six bases. For each system, four VMMC simulations were performed for 4×10^{10} steps, using umbrella sampling to improve equilibration.

intrinsic T_m). The measurements of the Frank-Kamenetskii group also require some care: in particular, they rely on the assumption that the unstacked state of the nicked duplex has the same flexibility as a duplex containing two G nucleotides [142]. This will only be a reasonable assumption if the stacking along the single-stranded gap is always broken. This seems to be a very strong assumption, and if it is violated the effect would be to overestimate the probability of the nicked duplex being in the unstacked state.

The chosen parameterization gives a stabilization of duplexes by coaxial stacking which is similar to (if slightly smaller than) the average predicted by the Santalucia lab [141] and Pyshnyi and Ivanova [136, 137]. For the stacked/unstacked equilibrium of a nicked duplex, this parameterization gives ΔG_{coax} as ~ 1 to $1.5 kT$ more negative than that found by the Frank-Kamenetskii group [142, 143]. Neither discrepancy is overly large, and if stacking can occur across the two G nucleotides in a gapped duplex, this might make the agreement even better.

6.3 Summary

Overall, the model gives a good description of the average properties of the thermodynamics associated with the ssDNA to B-duplex transition. It is the only model to date to consistently describe stacking, duplex hybridization and hairpin formation, and it also provides a reasonable representation of various DNA motifs. Sequence-dependent effects are currently absent, but in some cases this can be an advantage. For example, the temperature dependence of mismatch location (see Section 6.2.3) may have been somewhat obscured by sequence-dependent variation. Further discussion of model properties and its accuracy is given in Chapter 9.

Chapter 7

Modelling DNA Tweezers

As mentioned in Chapter 1, DNA tweezers were initially introduced by Yurke *et al.* in 2000 as a prototypical DNA nanodevice [54]. The tweezers, consisting of strands which form the arms α and β , and the hinge (h), can be switched between open and closed states by the addition of fuel (f) and antifuel (\bar{f}) strands, as shown in Figure 7.1. The tweezers demonstrate the possibility of using DNA hybridization and toehold-mediated strand displacement to perform mechanical operations. As such, they have inspired the growing field of DNA nanodevices, as outlined in Chapter 1.

I have studied DNA tweezers to prove the versatility and efficiency of my model. When the simulations were originally performed [206] (with an older version of the model), no DNA nanodevice had ever been simulated with a coarse-grained approach. This is partly because of the complexity of such systems and partly due to the need to simultaneously represent rigid duplexes, flexible single strands and the hybridization transition. To date, I am unaware of any other simulations of nanodevices, excepting subsequent studies performed with my model (such as the two-footed walker simulation presented in Chapter 8).

7.1 Tweezer simulation methods

7.1.1 The model system

For computational simplicity, I chose to model a system approximately half the size of that used by Yurke *et al.*: the sequences are given in Appendix E. The duplex regions which form the body of the tweezers are 10 bp in length, and the overhanging sections for fuel

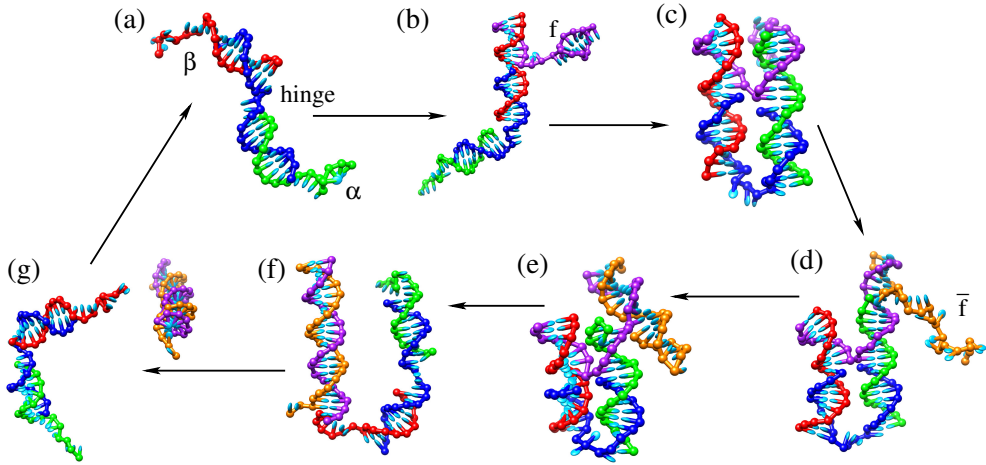


Figure 7.1: Simulation snapshots showing stages of operation of DNA tweezers. a) Tweezers initially open. b) Fuel (f) is added and binds to one arm (β). c) Fuel binds to the second arm (α) and closes the tweezers. d) Antifuel (\bar{f}) is added and binds to the toehold of the fuel. e) Antifuel begins to displace first arm of the tweezers. f) Tweezers open as first arm is displaced, and antifuel displaces the second arm. g) Antifuel fully hybridizes to fuel and the waste duplex is formed.

binding are 8 bases long. The toehold which facilitates the displacement of the tweezers is also taken to be 8 bases in length. I have sampled the free energy landscape of the system consisting of one set of tweezers and a single f and \bar{f} at 300 K, in a periodic cell of length 40 reduced units (3.96×10^{-20} litres).

7.1.2 Sampling the transitions

The sampling of the free energy landscape was performed using the VMMC algorithm, as introduced in Chapter 3. Every stage of the cycle is observable using unbiased simulations at 300 K. To obtain the free energy landscape, however, umbrella sampling was used to bias the ensemble and favour multiple transitions between (meta)stable states, as discussed in Chapter 3. As in the simulations of duplex formation performed in Chapter 6, the number of hydrogen-bonded base pairs (with energy below -0.1 simulation units) is a convenient order parameter to divide configuration space into regions which can then be biased. Due to the complexity of the tweezer system I found it necessary to introduce a four-dimensional order parameter $\mathbf{Q} = (Q_1, Q_2, Q_3, Q_4)$, with:

- Q_1 as the number of correct base pairs between α and f .

- Q_2 as the number of correct base pairs between β and f .
- Q_3 as the number of correct base pairs between f and \bar{f} , restricted to the bases of f which bind to α or are in the toehold.
- Q_4 as the number of correct base pairs between f and \bar{f} , restricted to the bases of f which bind to β .

In this context, *correct* base pairs are those which are intended to form during tweezer operation. Note that other base pairs are not forbidden, they are simply not included in the order parameter. Even an order parameter as complex as this was not sufficient to sample the tweezer cycle, particularly the transitions accompanying the displacement of the final bp of α and β from f . To estimate the free energy change associated with these processes, it is necessary to encourage the displaced strand to reattach. This requires that the displaced strand comes into close proximity with the duplex from which it was displaced, and that the duplex undergoes some fraying to allow it to reattach – a process not particularly well quantified by \mathbf{Q} as it stands.

\mathbf{Q} was therefore augmented with a fifth dimension, Q_5 , which depends on the actual separation of strands in addition to hydrogen-bonding matrices. Any configuration of the system is within one of 15 discrete values of Q_5 – the definitions are given in Appendix E.

Even with this complex order parameter, capturing the entire tweezer cycle in one simulation would be impractical. Instead, individual simulations were restricted to sampling small, overlapping ranges of \mathbf{Q} (for instance, the binding of the fuel to the first arm of the tweezers), and the results combined to give the free energy of the entire cycle. Ten simulations were performed in each sampling window for 4×10^{10} VMMC steps each. The windows used are detailed in Appendix E. The possibility of such an approach was suggested in the original work of Torrie and Valleau on umbrella sampling [156]. Kumar *et al.* [157] have proposed the weighted-histogram analysis method (WHAM) for systematically combining the results from overlapping simulation windows. This is particularly important for my system, as it has 185895 possible values of \mathbf{Q} , and 11 separate simulation windows which must be patched together.

In order to facilitate the windowed umbrella sampling scheme, several restrictions were imposed on the system during the simulations. These restrictions are outlined in Appendix E.

Simulation validity

When performing windowed sampling, the system has restricted movement through state space, which can lead to sampling errors. For example, it is conceivable that the state space corresponding to a certain order parameter range could be split into two non-connected regions. If this were the case, only one of these regions would be sampled in any simulation restricted to this range of the order parameter (ergodicity would be broken).

For the windowing scheme outlined in Appendix E, it seems unlikely that such a problem would arise. To be sure, however, I calculated the average energy as a function of the order parameter, $\langle E(\mathbf{Q}) \rangle$, in each simulation. For each pair of overlapping windows, the value of $\langle E(\mathbf{Q}) \rangle$ was compared for two frequently sampled values of \mathbf{Q} . In all cases, $\langle E(\mathbf{Q}) \rangle$ was found to be consistent between windows to within the error of the simulations (details are given in Appendix E). As a result, I am confident that simulations of each overlapping pair of windows had access to the same microstates, and therefore that the state space for each window was ergodically sampled.

The accuracy of the simulations can also be checked by considering that in going from the initial to the final state of the cycle, f and \bar{f} form a duplex, and the tweezers are in the open state in both cases. Thus the free energy change between the states with $Q_1 = Q_2 = Q_3 = Q_4 = 0$ and those with $Q_1 = Q_2 = 0, Q_3 > 0, Q_4 > 0$ should be the same as that for the formation of an isolated duplex of f and \bar{f} (assuming interactions between unbound molecules are minimal, as justified in Chapter 4). For the windowed tweezer simulations, ΔG was found to be $48.88 kT$, and from the isolated duplex simulations I obtained $\Delta G = 48.83 kT$. These values appear to agree well – in order to perform a statistical test of the agreement, 10 separate estimates of ΔG for the tweezer pathway were obtained by patching individual simulation results together. I compared this set to the six individual estimates of ΔG obtained for the simulation of duplexes in isolation using

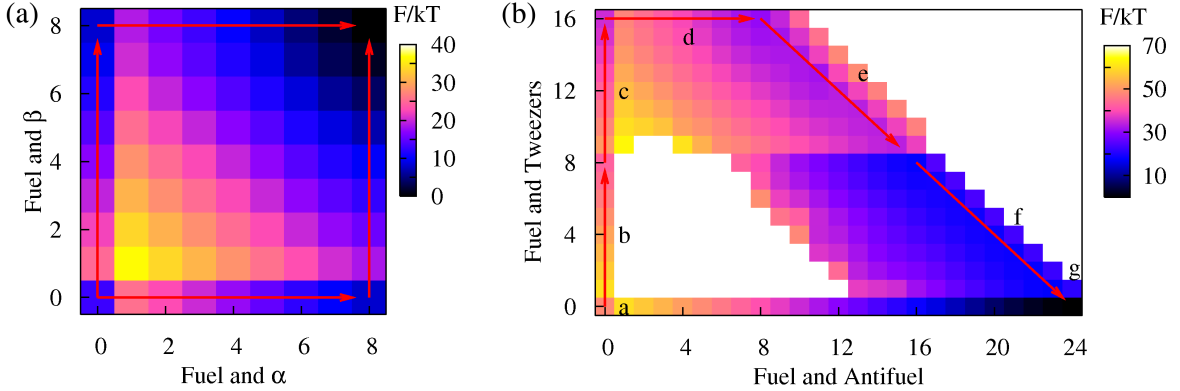


Figure 7.2: (a) Free energy F as a function of the number of f/α and f/β base pairs during initial binding of the fuel. The arrows indicate the cooperative lowest free-energy pathways. (b) Free energy F plotted as a function the number of f/\bar{f} and $f/\text{tweezer}$ base pairs for DNA tweezers at 300 K. White areas indicate regions of high free-energy relative to their environment that were unsampled.

Welch's unpaired t-test [214]. The result was a test statistic of 0.9413, suggesting that any systematic errors are smaller than the random error of the simulations.

7.2 Results

The free-energy landscape of the tweezer cycle is shown in Figure 7.2. To study the cycle in detail, it is convenient to consider a one-dimensional pathway through the landscape; I use that shown by the arrows in Figure 7.2 (b).

The gross features of the free energy landscape are as expected. Duplex formation is highly cooperative; the pairing of two strands involves a high entropic cost for forming the first base pair, then a downhill slope in free energy as additional bonds are formed. This is reflected in Figure 7.3 (a) by stages ‘b’, ‘c’ and ‘d’ which essentially involve duplex formation. The large cooperativity suggests that f will fully bind to one arm of the tweezers before binding to the second. The displacement processes (indicated by ‘e’ and ‘f’ in Figure 7.3) are comparatively flat as the total number of interstrand base pairs is constant. Returning the tweezers to the open state (between ‘e’ and ‘f’) and the decoupling of the $f\bar{f}$ duplex from the tweezers (‘g’) release the free energy stored in bringing strands together, resulting in large decreases in free energy.

Simulations allow for a detailed inspection of processes like displacement. Thus, Figure

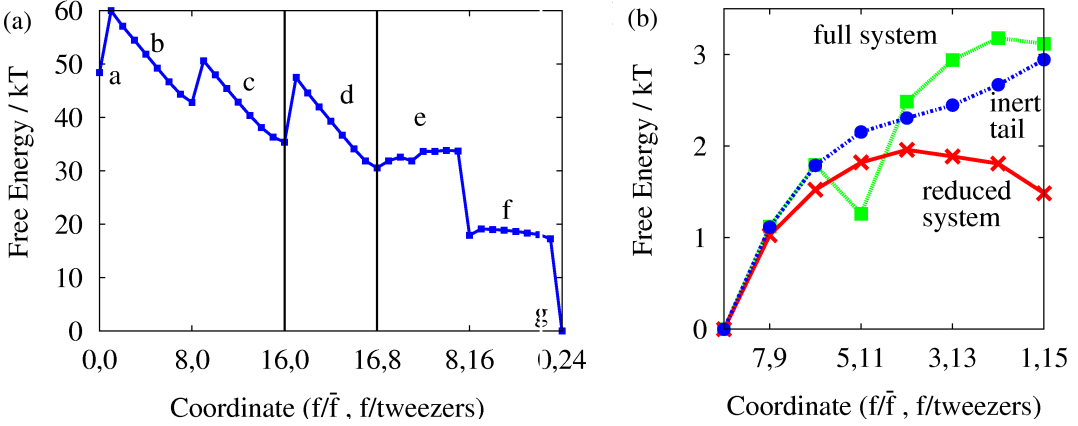


Figure 7.3: (a) Free energy profile along the one-dimensional pathway indicated in Figure 7.2 (b). Coordinates indicate the number of f/\bar{f} and f /tweezer base pairs. (b) The displacement process ‘e’ in more detail. Squares represent the original system, circles a system with the tail of \bar{f} unable to form a hairpin and crosses a system with the last eight bases of \bar{f} and most of the f/β arm removed.

7.3 shows that there is actually an increase in free energy of $\sim 3.2 \text{ kT}$ during the displacement of the first arm of the tweezers, α , even though the total number of interstrand base pairs in the system stays constant. Conversely, the displacement of the second arm (β) shows a decrease in free energy as displacement proceeds (after an unfavourable first step). These free-energy changes suggest a significant difference in speed for the two processes. To test this hypothesis, I initiated 20 unbiased Langevin simulations from both of the macrostates directly preceding displacement of the two arms, $\mathbf{Q} = (8, 8, 8, 0, 4)$ and $\mathbf{Q} = (0, 8, 16, 0, 8)$, and recorded when displacement occurred. In all 30 simulations initiated from $\mathbf{Q} = (0, 8, 16, 0)$, complete displacement was observed, requiring an average of 2.26×10^8 time steps. By contrast, only 83% of simulations initiated from $\mathbf{Q} = (8, 8, 8, 0)$ resulted in displacement before 275×10^9 time steps. This is consistent with the first stage of displacement being ~ 7 times slower.

The displacement of the α arm of the tweezers is shown in greater detail in Figure 7.3 (b). The first thing to note is that the free energy changes non-monotonically with the number of displaced bases of the α arm. The cause of the dip after three bases have been displaced is shown in Figure 7.1 (e): a metastable hairpin (with a three bp stem) is capable of coaxially stacking with the partial $f\bar{f}$ duplex, stabilizing the macrostate. Displacing more of the α arm means this hairpin cannot form, which costs free energy – this explains the large increase

in free energy associated with the displacement of the next base.¹

This is not enough, however, to explain the majority of the increase in free energy with displacement. To demonstrate this, I have simulated the displacement of the α arm, but with the final eight bases of f forbidden from forming bp (Figure 7.3(b)). As a consequence, the metastable hairpin cannot form, and the non-monotonic behaviour is no longer present. Furthermore, the free energy barrier of displacement is indeed reduced relative to the unaltered system, but remains fairly large ($\sim 3kT$).²

Excluding the rise in free energy associated with disrupting the hairpin, the largest increase occurs on displacing the first base of the α arm. To understand this, consider Figure 7.4. Before displacement, a single base of \bar{f} is forced to unstack and adopt a restricted conformation to avoid clashing with the two helices involved in displacement. When the first base of the α arm is displaced, it too must unstack and is restricted in its conformational freedom, which costs free energy. Alternatively, the coaxial stacking between the helices can be disrupted – but this is also costly. Essentially, opening up a second single-stranded region is an unfavourable process, and this explains the initial rise in free energy as displacement begins. It does not, however, explain why the free energy continues to increase as further bases are displaced.

To understand this, note that the tweezers are a fairly bulky object, and as displacement proceeds the single-stranded tails are increasingly drawn into the body of the tweezers. This is entropically unfavourable, as the conformations of the flexible single strands are restricted by the tweezer unit, and this effect contributes to the increase in free energy with displacement. To demonstrate this, I considered a system in which the final eight bases of \bar{f} and all but the first bp of the $f\beta$ duplex were removed. The results, shown in Figure 7.3(b), clearly demonstrate that a significant contribution to the rise in free energy from displacement is due to steric clashes involving these sections.

¹The existence of a dip in free energy at (5,11) is the only significant difference between this work and Reference [206], which used an earlier version of the model that lacked coaxial stacking.

²If Figure 7.3(b) is analyzed closely, one finds that the final step is actually less favourable for the inert hairpin than the normal system. This is because there is some tendency for the two single-stranded tails of \bar{f} and α to transiently bind at this point (lowering the free energy), which is obviously impossible for the inert tail.

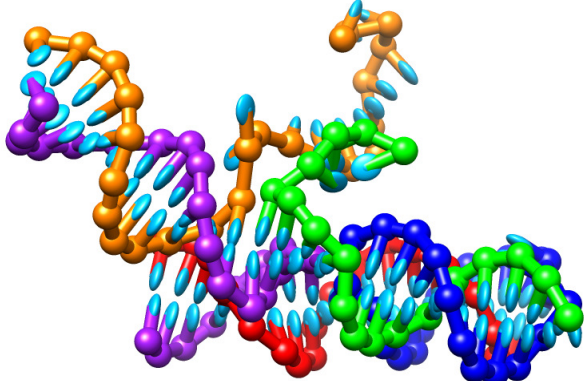


Figure 7.4: Tweezers (viewed from the side) during the displacement of the α arm ($Q_1 = 4$, $Q_2 = 8$, $Q_3 = 12$, $Q_4 = 0$). Note the restricted confirmations of both single-stranded tails.

The displacement of the β arm of the tweezers, shown in Figure 7.1 (f), is noticeably different. There is an initial free-energy penalty for opening up a second single-stranded section, as before, but once displacement has started it becomes favourable to displace more bases. As displacement proceeds, the $\bar{f}f$ duplex moves further away from the body of the tweezers, reducing the excluded volume interactions between them and favouring states with greater degrees of displacement.

7.3 Discussion

This chapter has shown the potential of the model to study nanotechnology. Due to its efficiency and physically reasonable description of DNA, it was possible to simulate the entire cycle of DNA tweezers, the first time a DNA nanomachine has been studied in this way.

Although the simulation was primarily undertaken as a proof of principle, non-trivial results were obtained. In particular, the model demonstrated that displacement processes can involve gradients in free energy, a result that was initially surprising. The causes of these gradients are, however, physically reasonable. Firstly, the necessary disruption of a metastable hairpin contributed to the free-energy barrier which opposed displacement of the α arm. The SantaLucia parameters predict that such a hairpin would indeed be metastable at 300K (in fact, due to the presence of two CG base pairs in the stem, it

would be significantly more stable than average, with a melting temperature predicted as 313.7 K). Furthermore, although this small system could be designed so as to eliminate such a hairpin, careful design would be needed to prevent similar structures forming when longer scale displacements are required. As hairpins will tend to form either at the start or end of displacement, when long single-stranded regions are available, they will constitute a free energy barrier in the middle stages of displacement, thereby slowing down the process.

The second contribution to the free energy rise during displacement is the initial cost of creating two single-stranded sections, which necessitates disrupting stacking interactions. Although there is little experimental data with which to compare this prediction, its motivation is physically reasonable. Furthermore, it is possible that in reality the sharp bending of single strands to avoid the branch point of displacement may incur some energy penalty that is not captured by the model, which would exaggerate the effect.

Finally, there is a significant entropy cost associated with bringing the single-stranded displacement tails into a region of high DNA density (the centre of the tweezers). Again, the physical basis of this effect is applicable to real DNA. Indeed, it is possible that electrostatic effects would exaggerate it further, particularly at moderate or low salt concentrations. In addition, as the tweezers of Yurke *et al.* are larger than those simulated here and thus have longer single-stranded tails, the consequences of steric interactions may be more significant for their system.

For the tweezers, in which the displacement process is fast compared to the initial association of duplexes [54], the consequences of any free-energy barrier to displacement are probably fairly minimal. The observation of a barrier to displacement, however, may help to explain an initially puzzling result. Yurke and Mills have shown that the rate of toehold-mediated strand displacement depends exponentially on the toehold length (or equivalently the free energy of toehold binding), up to a length of at least six bases when displacing 26 bp [53]. Zhang and Winfree have reported similar results, and have also demonstrated that the rate plateaus for longer toeholds [215]. Such a result could only be fitted with kinetics that assumed displacement was essentially a two-state process with a single rate

constant, rather than a random walk in which each intermediate state was equally probable. My simulations suggest that metastable hairpins and the consequences of having two single-stranded regions rather than one make the intermediate states less favourable. Such arguments make displacement look more like a process in which the system must hop from one low free-energy basin to another than an unbiased random walk, and hence may go some way to explaining the results of References [53] and [215].

As ever, all results should be viewed with the approximations of the model in mind. In this case, the tweezer arms are brought into close proximity when they are closed by the fuel – electrostatic repulsion will probably be relevant here, even at high salt. In addition, the lack of restriction on the conformation of ssDNA in the model may mean that the stability of the $f\alpha - f\beta$ junction is overestimated.³ Either of these considerations may mean that the stability of the closed state of the tweezers is exaggerated by the model, but they should not affect the primary conclusions related to the displacement process drawn from the study.

³It is plausible that the destabilizing effect of electrostatics could account for the population of partially open tweezers inferred from the single-molecule FRET analysis of tweezers performed by the Simmel group [216].

Chapter 8

Modelling a DNA walker

In this chapter I study the operation of a two-footed DNA walker proposed by Bath *et al.* [65]. The walker consists of two single-stranded ‘foot’ domains which are linked together by a duplex (Figure 8.1). The feet are intended to bind to adjacent sites on a single-stranded track. The binding sites overlap, meaning that the feet compete for binding to the track. As a consequence, a single-stranded region of one foot or the other is always exposed. A fuel strand that is also present in solution can bind to either exposed toehold. If the fuel has bound to the back foot,¹ then it is able to compete with the track for binding to the rest of the foot. The fuel can thus displace the track and cause the foot to be raised. If, alternatively, the fuel has bound to an exposed toehold of the front foot, conventional toehold-mediated strand displacement is impossible and the fuel should eventually detach.

The fuel contains a recognition site for the nicking enzyme N.BbvC IB, which can cleave the fuel when it is in a duplex state (the duplex partner is not cleaved). With the fuel in two separate pieces, the fuel/foot duplex is unstable and the fuel will eventually detach, allowing the foot to rebind to the track. If it binds in front of the other foot, a forwards step is taken – if it binds behind the other foot, the system has returned to its original state and an idle step has occurred.

Due to the intended asymmetry of foot-lifting, the walker is expected to undergo unidirectional, autonomous motion. For the walker to function in this way (with the potential to perform work), it must catalyze the equilibration of a non-equilibrium system – in this

¹Throughout this chapter, the terms *back* or *backwards* are defined by the 5' direction of the track, and *front* or *forwards* by the 3' direction.

case, it facilitates the enzymatic cleavage of ssDNA fuel strands.

To date, the principle of operation of the walker has been demonstrated on a short track of two sites [65]. Preferential lifting of the front foot by a fuel strand has been achieved (as has lifting of the back foot by a ‘reverse fuel’). The release of the fuel has also been shown, and walkers have been found to successfully catalyze the hydrolysis of ≥ 64 fuel strands.

8.1 Walker simulations

I have simulated every stage of walker operation, with the exception of the hydrolysis of the fuel strand (for obvious reasons). As the sizes of individual components are potentially pivotal to walker operation, I considered the system exactly as introduced by Reference [65], at a temperature of 310 K (the sequences are given in Figure 8.1).

8.1.1 Binding of a foot to the track

Consider the stage in the walker’s cycle corresponding to moving from state (v) to (vi) or (i) in Figure 8.1 (a).² To study this process, I performed 50 Langevin simulations of a system consisting of a track of three binding sites, with one foot attached to the middle site of the track, and the other initially in a raised position (Figure 8.2 (b)). Figure 8.3 shows the state of these simulations after 5×10^9 steps, at the end of the simulations.

Typically, two types of initial binding were observed:

- binding to the front or back site in the intended (correct) manner (Figures 8.2 (a) and (c)).
- binding to the front or back site in an unintended manner (forming misbonds - Figures 8.2 (d), (e), (f), (g) and (h)).

Misbonds occurred for one of two reasons – either, as in Figure 8.2 (d) and (g), because the foot possesses two competition domains and these can attach to the track in the wrong place, or as in Figure 8.2 (e), (f) and (h), because the sequence of the toehold domain is

²In fact, it is not clear that the fuel must completely detach before the foot rebinds to the track – the possible consequences of this will be discussed in Section 8.1.5.

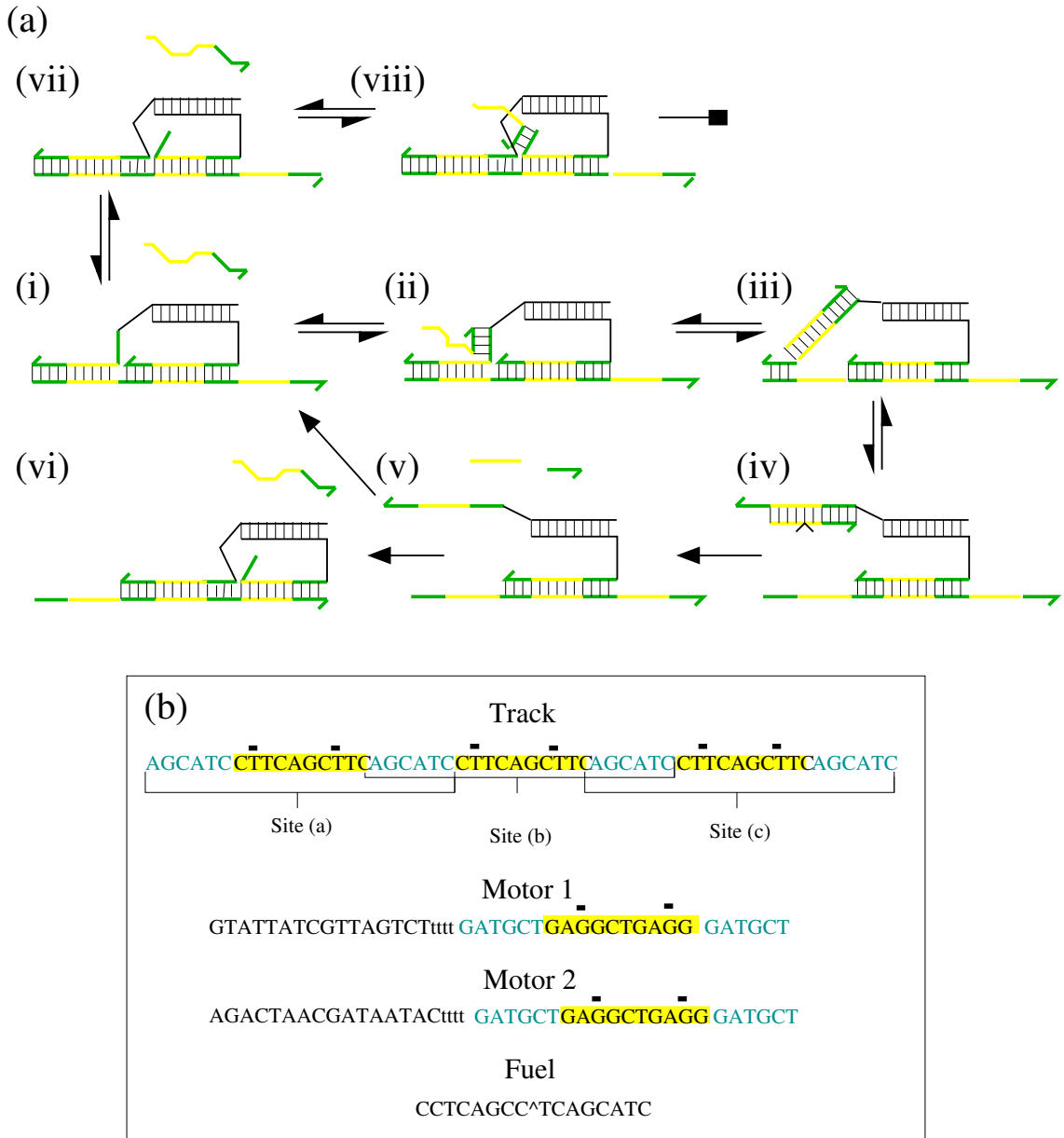


Figure 8.1: (a) Schematic diagram of walker operation, adapted from Reference [65]. Reactions which are expected to show a large decrease in free energy are shown as irreversible. (i) and (vii) Walker bound to track, with competition between the two feet for binding. (ii) Fuel binds to raised toehold of the back foot. (iii) Fuel competes with the track for binding to the back foot. (iv) Back foot detaches from track and nicking enzyme binds to the recognition site. (v) Fuel is nicked by the enzyme and detaches. The raised foot can then rebind either in front (vi) or behind (vii) the attached foot, corresponding to taking an active step or idling respectively. As an alternative to (ii), the fuel can bind to the raised toehold of the front foot: due to the geometry of the strands, however, conventional displacement cannot proceed from this point. (b) Sequences used in the tweezers (written in 5' to 3' notation). Green highlighting indicates competition (or toehold) domains, and yellow the non-competition (non-toehold) domains associated with binding to the track. Three adjacent binding sites of the track are shown here. The lower case 't's represent the bases which link the walker body duplex to the feet. To prevent cleavage of the track by the nicking enzyme, mismatches are included between foot and track: these mismatches are indicated by bars. The nicking site of the enzyme within the fuel is indicated by the [^] symbol.

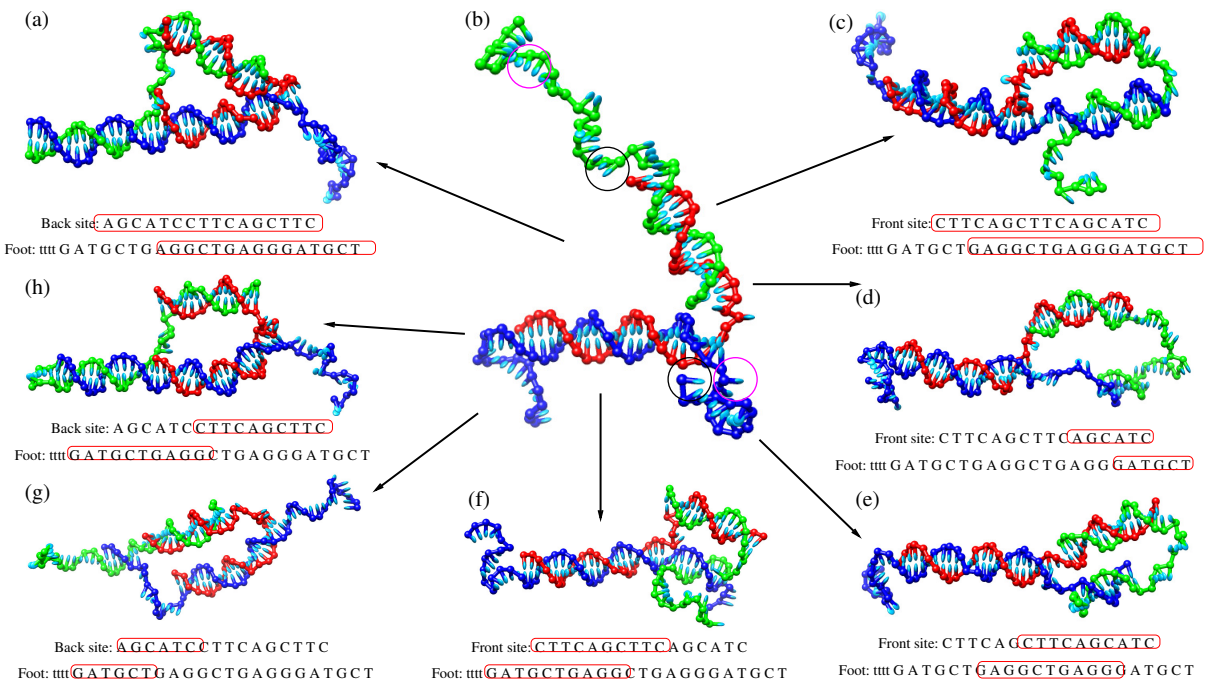


Figure 8.2: The binding of a walker to the track as observed in simulations. (a) The walker bound to the back site. (b) The walker with foot raised; magenta and black circles indicate the typical separation of bases that could form correct base pairs. (c) The walker bound to the front site. (d) – (f): common misbonds with the front site. (g), (h): common misbonds with the back site. Highlighted regions show the motifs involved in pairing, all sequences are listed in 5' to 3' notation.

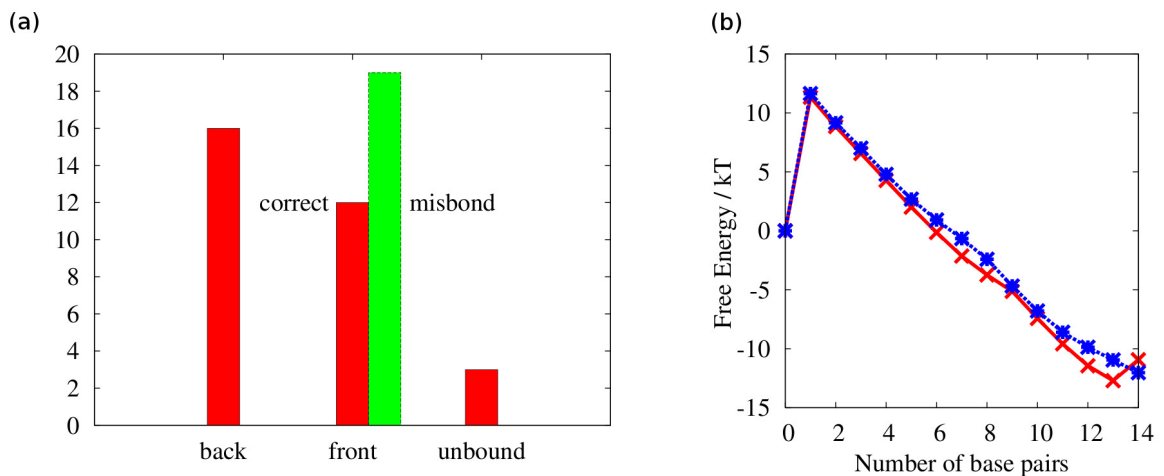


Figure 8.3: Results of foot-attachment simulations. (a) The final state of 50 Langevin simulations initiated with one foot in the raised state. (b) The free energy profile of bonding to the front (red) and back (blue) sites as a function of the number of correct base pairs, measured relative to the state with no bonds. The free energy profiles were obtained from 10 VMMC simulations of 4×10^{10} steps, with umbrella sampling used to improve equilibration.

very similar to the non-toehold domain (which itself contains a repeated motif). The most common (and most stable) misbonds were those shown in Figure 8.2 (d) and (e).

In some simulations, misbonds were displaced by correct bonding to the track. The most obvious way that this can occur is in an ‘inchworm’ fashion, wherein some correct base pairs are formed, creating a bulge that is eventually pushed through the misbond, leaving the correct bonding in place. An intermediate stage of displacing a misbond such as that in Figure 8.2(h) is shown in Figure 8.4(a). For this particular misbond, the displacement process is facile as correct base pairs can form without disrupting the misbond, acting as a toehold for displacement. Other misbonds do not provide obvious toeholds, and hence the rate of displacement is suppressed.

An alternative mechanism exhibited by my model is far less obvious. In this case, one strand reaches back to form a ‘double-X’ structure, shown in Figure 8.4(b). There is no obvious route to *fully* displace the misbonding from this position, but the misbond is somewhat destabilized (partially due to the loss of dangling end stabilization) and hence has a greater tendency to melt than in a normal configuration. If the misbonding melts before the other half of the structure, displacement has, in effect, taken place. This process has been observed for the misbonds of type (d) and (g), for which the inchworm form of displacement is particularly slow. Misbonds of type (f) are reasonably rare, and the only example of displacement involving this structure caused it to be converted into a type (d) misbond by a double-X mediated process.

I have also observed displacement wherein the back site displaces a misbond with the front. This process is similar to normal strand displacement, except it is generally slowed by pauses at the mismatch locations.

In some other cases, misbonds were observed to detach from the track, allowing the foot to rebind. Whether through displacement or melting, all misbonded feet would be expected to find the lower free-energy states of correct bonding eventually.³

³Unbiased VMMC simulations performed at the same temperature demonstrate similar behaviour, suggesting that these results are not overly dependent on the details of the simulation technique. Although statistics are fairly poor, the VMMC simulations appear to give a somewhat higher probability of any misbonds melting before they are displaced – nevertheless, binding through displacement is still observed.

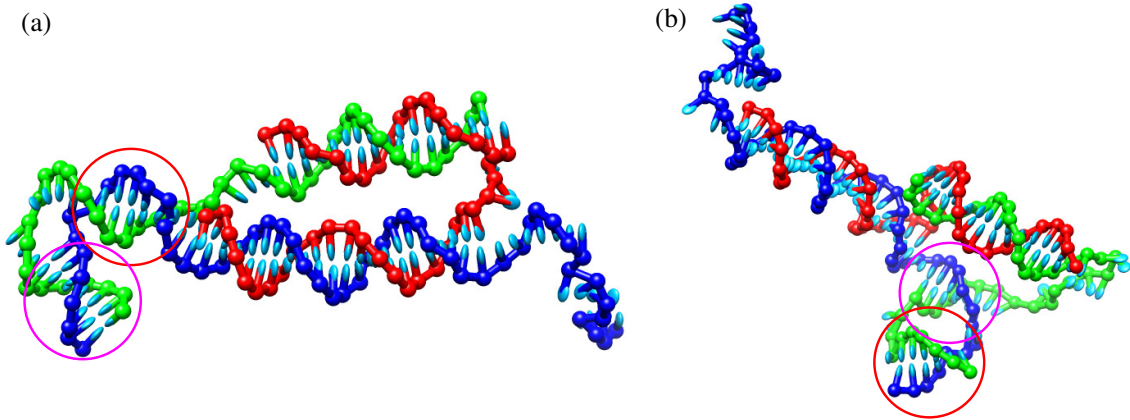


Figure 8.4: Displacement as observed during simulations. (a) ‘Inchworm’-style displacement of a back foot misbond. (b) ‘double-X’ structure containing a misbond with the front site. Misbonded sections are highlighted with red circles, and correctly-bonded regions with magenta circles.

The stability of the misbonds as isolated strands can be estimated using the nearest-neighbour model of Reference [91]: they are predicted to have melting temperatures slightly above that for an average 6-bp duplex with two dangling ends.⁴ This level of stability suggests that the misbonds could be kinetically relevant in the physical system, as they have a role in my simulations where the type of misbond shown in Figure 8.2 (d) is a close approximation to an average 6-bp duplex with two dangling ends.

The absence of a foot replacement bias

The walker was not originally designed to have a bias associated with foot replacement. From looking at Figure 8.2 (a), however, there is clearly an asymmetry between the forward and backward directions, as the raised foot is attached to the front end of the track-bound foot. As mentioned in the previous section, misbond formation is far more common with the front site. Generating a bias for foot replacement would be extremely beneficial for the walker – without such a bias, its efficiency is limited to 50 %.

From the evidence of Figure 8.3 (a), the geometrical asymmetry does not appear to translate to a bias towards stepping forwards correctly. Instead, a roughly equal number

⁴Using the parameters of Reference [91], I calculate melting temperatures of 316 K, 311.7 K and 312.8 K at 0.000336 M for strands carrying the motifs causing misbonds in Figures 8.2 (d) and (g), (e), and (f) and (h) respectively. This should be compared to that of an ‘average’ 6-bp duplex with two dangling ends, which has a T_m of 309.4 K.

of simulations were observed to be bound correctly in front of or behind the attached foot at the end of the simulations. Even if all of the simulations that were ended while in a misbonded state were destined to eventually attach correctly to the front site, the ratio of forwards to backwards steps would still only be $\sim 2 : 1$. It is clear that my model predicts that any bias associated with foot replacement is minimal.

The lack of an obvious bias was initially surprising, as Figure 8.2 (b) suggests that the raised foot is inherently closer to the front site. Part of the reason may be the greater ease with which back misbonds can be converted into correct binding to the back site, as discussed in the previous section. Many simulations, however, directly formed correct bonding, and simulations in which misbonding was forbidden provided little evidence of a substantial bias for stepping forwards.

The flexibility of single strands ensures that this difference in proximity of the front and back sites is somewhat illusory. In order to form a correct base pair between the front site of the track and the raised foot, it is necessary for a single-stranded region to stretch across the length of the 16-bp duplex which links the two feet together – this is decidedly unfavourable (see Figure 8.2(b)). By contrast, forming a correct base pair with the back site does not require such stretching (only the approximate anti-alignment of the walker body and the track/foot duplex). This visual argument is borne out by the free energy of bonding for the front and back sites, which indicate only a marginally lower free energy cost for forming the first few (correct) base pairs with the front site than with the back (Figure 8.3 (b)).

Binding of the feet to a track under tension

In the previous section, it was argued that the asymmetry of the walker did not result in a large bias for binding correctly to the front site over the back site due to the contractility of ssDNA. A possible method of overcoming this difficulty, and thereby generating biased foot replacement, would be to apply a tension to the track. The question of how the walker behaves on a stretched track is also an interesting problem in its own right, as tracks will have to be stretched to generate motion in a chosen direction.

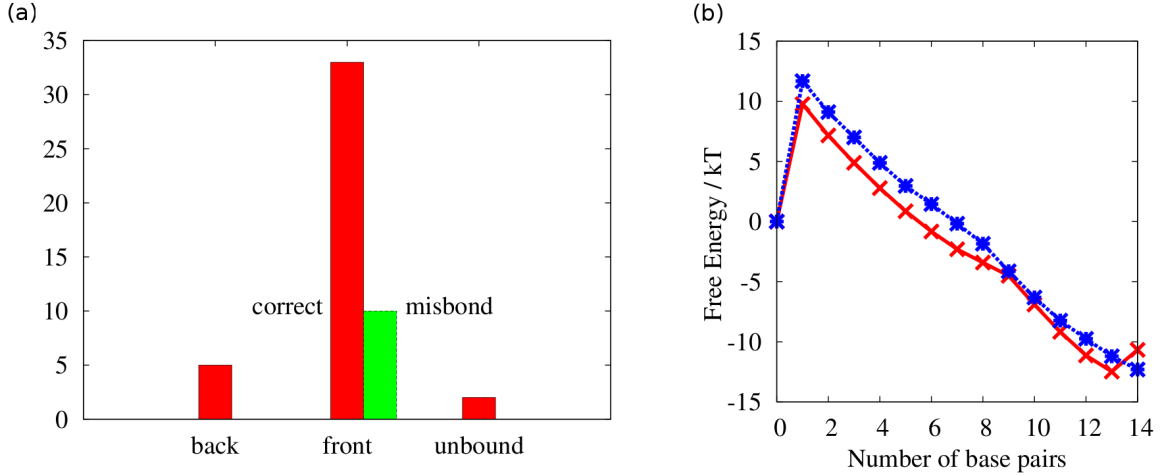


Figure 8.5: Results of foot-attachment simulations with a tension of 14.6 pN applied to the track. (a) The final state of 50 Langevin simulations initiated with one foot in the raised state. (b) The free energy profile of bonding to the front (red) and back (blue) sites as a function of the number of correct base pairs, measured relative to the state with no bonds. The free energy profiles were obtained from 10 VMMC simulations of 4×10^{10} steps, with umbrella sampling used to improve equilibration.

I performed 50 Langevin simulations of 5×10^9 steps in which the track was subjected to a constant tension of 14.6 pN, a physically reasonable force to apply to a DNA (enough to cause significant extension of single strands, as shown in Chapter 5, but not sufficient to cause overstretching of long duplexes). The results, shown in Figure 8.5 (a), now indicate a significant (although not overwhelming) preference for stepping forwards (similar results were obtained using unbiased VMMC and with tracks under half the tension). The majority of this bias is a result of the increased rate of directly binding to the front site in the correct manner, as the bases at the end of the track are now automatically stretched into a more convenient location for binding. Such an explanation is supported by the free energy profiles of binding under tension shown in Figure 8.5 (b), which show that the result of the force is to make the formation of the first few correct base pairs with the front site far easier. The mechanism is further highlighted by considering the probability of having a single correct base pair with different bases along the track as measured in VMMC umbrella sampling simulations. The application of tension dramatically increases the weight of being paired to bases at the far end of the front site (by approximately a factor of 7), whilst barely affecting the probability of pairing with bases closer to the attached foot, and slightly suppressing pairing with bases at the far end of the back site.

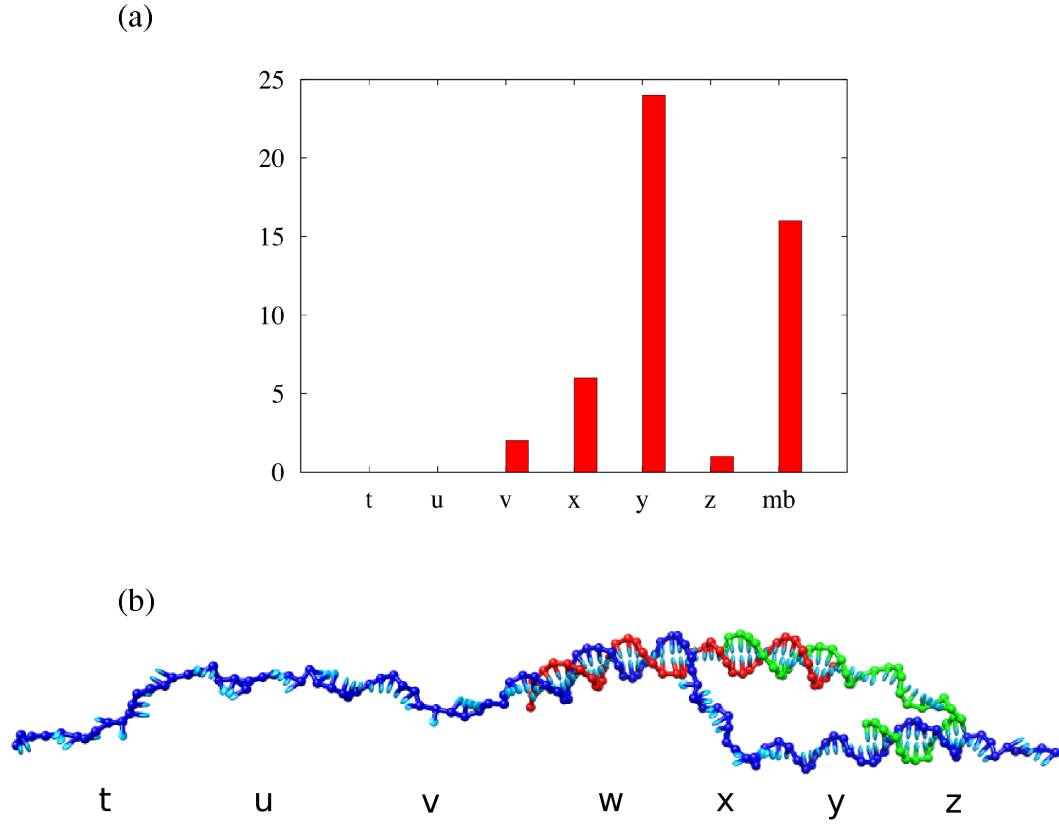


Figure 8.6: Results of foot-attachment simulations with a tension of 14.6 pN applied to a track with six available binding sites. (a) The final state of 50 Langevin simulations initiated with one foot in the raised state. (b) A double over-stepped configuration, exhibiting considerable fraying.

Binding to an extended track

If the walker is to be useful, it will have to take many consecutive steps along an extended track. To test the feasibility of such a process, simulations were performed for 3.5×10^9 steps on a track with seven binding sites (hereafter labelled *t*–*z* from back to front), with one foot initially bound to site *w* and the other raised. The track was subjected to a tension of 14.6 pN. The results, shown in Figure 8.6 (a), indicate that the most common result is binding to site *y*; a single overstep. This would be a disaster for the walker – when overstepped, there is no competition between the two feet and hence the lifting of either foot is strongly suppressed. In this configuration, the walker is essentially stuck.

It is even possible to double overstep, by binding to site *z*. Such binding, however, requires considerable stretching and hence walkers bound in this configuration have a tendency for the front foot to fray away from the track (Figure 8.6 (b)). Consequently, the fuel

binding site can be revealed and the foot lifted.

Misbonds are also relatively common – generally they are based on the motifs of Figure 8.2 (e) and (f). The extra length of track in these simulations allows the formation of a few extra base pairs. To form these extra pairs, the duplex must enclose several mismatches or internal loops. In my model, these structures are stable at 310 K – as discussed in Chapter 6, however, my model generally overestimates the stability of structures involving internal loops and mismatches relative to Reference [91]. It is therefore probable that these kinetic traps would be less of an impediment for the actual walker than in my system, although they may still be kinetically relevant. In particular, the most common misbond in simulations involves binding to site y in the manner of Figure 8.2 (f), with some additional pairing with site x . It is likely that the additional pairing would likely be less stable in reality than in my model – this would open up a toehold for an inchworm-type displacement of the misbond by correct binding to the y site. No backwards steps were observed in this set of simulations. However, this is likely to be due to the low statistics, given that backwards steps were observed for a short track under tension.

Note that the mismatches such as those shown in Figures 8.2 (d) and (g) do not have an analogue in this larger system – with an extended track, these misbonds constitute partial correct binding at the adjacent site.

Simulations performed with a tension of 7.3 pN gave similar results. Differences are as would be expected – binding to site z is more frequent and stable at lower tension, for example.

8.1.2 Competition between feet

Figure 8.7 (a) shows the free-energy landscape associated with the competition between the two feet of the walker (attached to the front two sites of a short track). The maximum number of base pairs for each foot is 20 (that foot has then ‘won’ the competition, and the toehold on the other foot is raised), and the maximum number of base pairs with the non-toehold domain is 14. There appears to be only a small free energy difference between having the front and back foot toeholds raised. Note that when the front foot has lost the

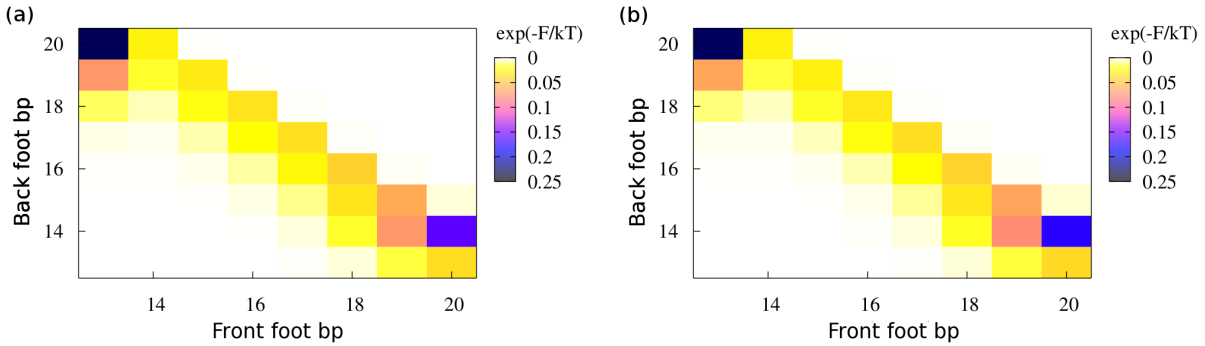


Figure 8.7: Free-energy landscapes associated with the competition between toehold domains, obtained from 10 VMMC simulations of 4×10^{10} steps each. (a) With no applied tension. (b) With the track under a tension of 14.6 pN.

competition, it tends to fray as it is not favourable to enclose a mismatch only one base pair from the end of the duplex.

The important point to note is that there is remarkably little difference between Figure 8.7 (a) and Figure 8.7 (b) (which shows the free-energy landscape for displacement under a tension of 14.6 pN). The similarity suggests that such a tension should have a limited effect on the bias of foot-lifting, as the relative exposure of toeholds is virtually unchanged. The effect of the tension appears to be to differentially influence the transition states of foot-binding, and hence the probability of binding to the front and back sites, but not the final states of foot-binding. This is perhaps not surprising, as the bases in the track are essentially transferred from one duplex to another by competition, and so the track's extensibility is similar in both cases.

8.1.3 Fuel binding and displacement

Figure 8.8 (a) shows the free-energy landscape of fuel binding to the feet of a walker (attached to the front two sites of a short track) without tension. The probability of binding to each foot is roughly equal, with a slight bias towards the front foot toehold (possibly because when the front toehold is exposed, it is not constrained by being part of a loop). When bound to the back foot, displacement can proceed: the free-energy profile for the displacement of the fuel and release of the foot is shown in Figure 8.8 (b). The steps at 7 and 12 bp between the track and the foot indicate points at which mismatches are repaired. These mismatches

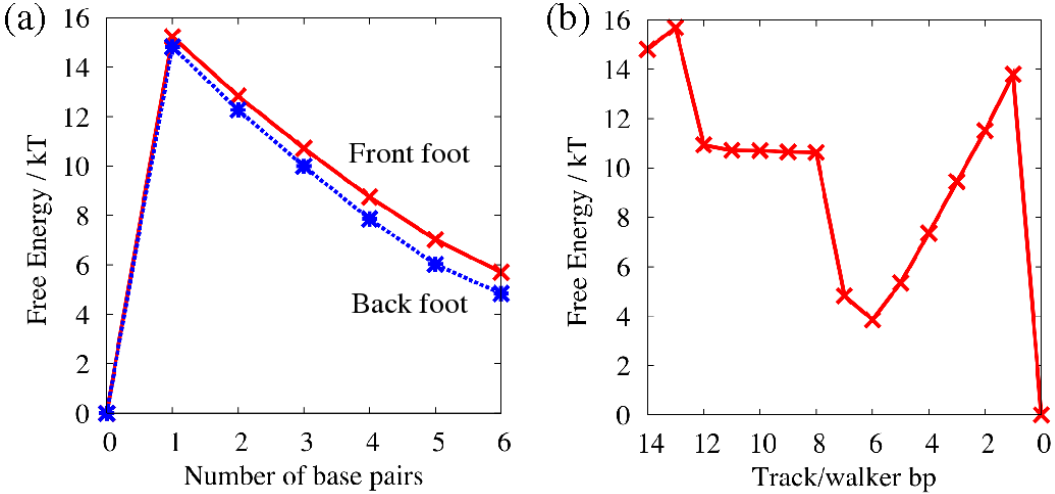


Figure 8.8: Free-energy profiles associated with (a) the binding of fuel to the raised toeholds and (b) the displacement of the track from the back foot by the fuel. Each profile was obtained from 10 VMMC simulations of steps, performed in a periodic cell of side length 80 reduced units.

give a strong bias favouring displacement, which in this case is likely to be beneficial as the fuel is unable to displace all the bp between track and foot, and must wait for the final 6 bp to detach spontaneously.

8.1.4 Lifting of the wrong foot

Bath *et al.* [65] performed an experiment in which one of the feet had its non-competition domain mutated so as to be non-complementary with the fuel. The rate of fuel binding was estimated by addition of a fluorescently-labeled fuel molecule whose signal would increase upon binding. The signal was observed to increase around 30 times faster when the mutant foot was the front foot, suggesting that the bias for lifting the back foot was approximately 30:1. For back-foot lifting, the signal was well modeled by a second-order process; the lifting of the front foot appeared to be more complex.

Two possible mechanisms which could lead to this leak current are:

- The front foot is able to fray from its far end, partially revealing the binding site for the fuel and allowing displacement.
- The fuel can bind to the toehold of the front foot, and then somehow displace the front foot from the track using this as a toehold.

Explanations which involve some fraction of the DNA having errors or being in a misformed state when the fuel is introduced seem to be unlikely, as by the end of the front-foot experiments around 50 % of the walkers had raised feet (see supplementary material of Reference [65]).

Yurke *et al.* [53] and Zhang *et al.* [215] have demonstrated that the blunt-ended displacement involved in the first mechanism is highly suppressed relative to toehold-mediated strand displacement. Both groups found that toeholds of 6 bp accelerate displacement of ~ 20 bp duplexes by around a factor 10^6 relative to blunt-ended displacement at room temperature. Given these values, a factor of only 30 for the walker seems surprisingly low. One should note that for the walker, only 8 bp must be displaced before a mismatch can be repaired, which will make blunt-ended strand displacement more likely to succeed once it has started. Furthermore, it is possible that being attached to the body of the walker makes the end of the foot/track duplex less stable, again favouring displacement.⁵

Exploratory simulations showed the possibility that, whilst bound to the front toehold, the fuel could loop round and displace base pairs between the front foot and the non-competition domain. This suggests a possible mechanism of incorrect foot-lifting, in which the fuel detaches from the front toehold whilst bound to then non-competition domain, allowing displacement of the track from the foot to proceed. Such a process, involving a double-X-like structure, is illustrated in Figure 8.9.

I performed forward flux sampling (FFS), as outlined in Chapter 3, to estimate the flux from a state in which the fuel is bound to the toehold to one in which it has formed 8 bp with the non-toehold region.⁶ The sampling details are given in Appendix F. 50 more unbiased simulations were initiated from this point, of which 25 returned to the initial state with no binding to the non-toehold region, 25 completed the displacement and one failed to

⁵There is no evidence from my simulations that being attached to the body of the walker causes a significant unpeeling tension on the end of the front foot/track duplex. The presence of explicit electrostatics, however, would tend to generate some repulsion between the walker body and the foot/track duplex. This would favour conformations in which they are separated by a greater distance, perhaps providing some tendency for the foot/track duplex to peel from its front end.

⁶8 bp between fuel and foot was found to be a metastable minimum of free energy. At this point, the fuel has repaired the two mismatches between track and foot, which is favourable, but forming any more base pairs is geometrically difficult without melting the fuel/toehold duplex (see Figure 8.9).

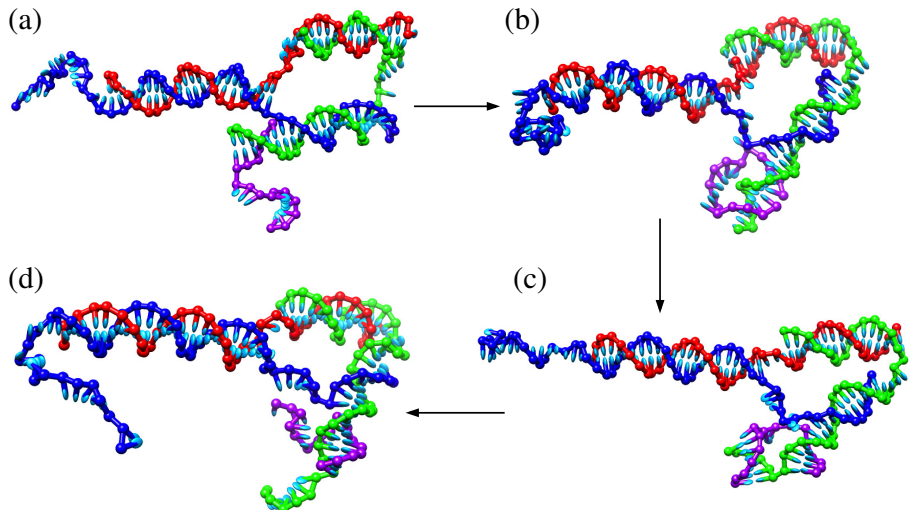


Figure 8.9: Postulated mechanism for raising the front foot via displacement. (a) Fuel is bound to raised front toehold. (b) Fuel is bound to first three bases of the non-competition domain, having displaced the track. Fuel displaces more bases, forming a double-X-like structure. Fuel detaches from toehold, allowing displacement to proceed.

do either within the simulation time. I compared the overall flux for displacement to that for simple dissociation of the fuel from the walker.

The two estimates give a relative flux of 286:1 (error $\sim 17.2\%$) in favour of simply melting the fuel/toehold duplex, rather than melting the toehold whilst displacing the track. Thus $\sim 0.35\%$ of systems with toehold binding to the front foot should displace the track from the foot rather than simply dettaching.

Thus the model suggests that this mechanism should have a rate of order 300 times slower than the lifting of the back foot by the fuel. This number should be treated with great caution – a number of limitations in the model may cause it to be significantly different for real DNA (for instance, electrostatic effects may suppress the configurations required, or the extra destabilization of the foot/track duplex due to mismatches may favour the mechanism). Nonetheless, the model suggests that this mechanism for a leak current is not completely implausible.

It should be relatively simple to distinguish between the two displacement mechanisms in experiment. By mutating one or other of the two toehold domains of the front foot, either process can be effectively prevented from happening. If the leak current is unaffected by

the change, the mechanism in question can be eliminated as a cause.

Establishing the dominant cause of lifting the wrong foot is important for optimizing the walker. In particular, the mechanism analyzed here is strongly favoured by the location of a mismatch near to the end of the non-toehold domain, which leads to increased fraying and favours displacement. If this mechanism is discovered to dominate the leak current, the sequence design could be reconsidered. Furthermore, the pathway explored above is unlikely to be particularly sensitive to whether the walker is attempting to do work against a force. Such a force would favour peeling of the front end of the foot/track duplex, and hence exacerbate the alternative blunt-ended displacement.

8.1.5 Fuel dissociation

In Section 8.1.1, it was assumed that reattachment of the foot occurs after the severed fuel and nicking enzyme have completely dissociated from the foot. This may not, however, be the case. One plausible scenario is that one or other of the halves of the fuel, along with the enzyme, will dissociate first, leaving the other half still attached. What consequences could the presence of half of the fuel have for reattachment (here, the terms *proximal* and *distal* refer to the proximity of the fuel to the body of the walker – see Figure 8.10(a)):

- Proximal half of the fuel still attached. In this case, the section of the foot which would bind to the back site is mostly exposed, whereas the front binding site is largely inhibited.
- Distal half of the fuel still attached. In this case both binding sites are partially inhibited by the presence of the fuel.

I have performed exploratory simulations for each of systems with both long and short tracks under tension in which the walker is initiated with either half of the fuel still attached to the raised foot. Currently data is limited, but the preliminary findings are given below.

- Proximal half of fuel remaining: misbonds of type (d) and (e) are very common, but displacement by the correct site is prevented by the presence of the fuel remnant. The

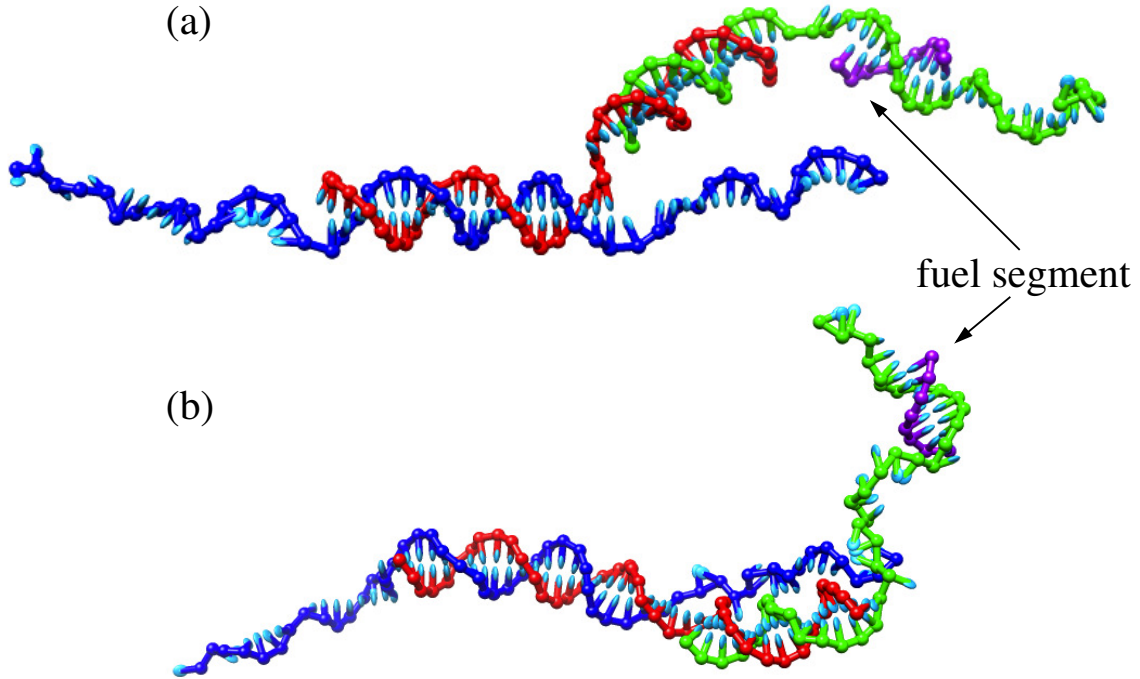


Figure 8.10: Raised foot with (a) proximal and (b) distal half of fuel attached.

12 available bp with sites v and y mean that the foot often binds in these configurations (overwhelmingly with y for an extended track). Displacement of the remainder of the fuel is reasonably facile for site y , but not for v as the other foot blocks displacement of more than 2 bp. Presumably, however, detachment of the fuel will eventually occur. Successful binding to site x (involving fuel displacement) has been observed, but appears to be rare.

- Distal half of fuel remaining: binding to the 8 available bases at the front of the x site is most probable. Binding to site v appears to be suppressed, although it has been observed, and binding to site y can also occur. Although no full displacement of the fuel by site x has been observed in this system to date (a mismatch hampers progress), the eventual outcome is likely to be that the fuel is displaced by the track, as this will have a lower free-energy barrier than simply melting the track/foot duplex.

Fully binding to any site on the track requires that the fuel fragment dissociate. This could occur whilst the foot is raised, which would leave the system in a state identical to

that considered in Section 8.1.1. As the fuel remnant/foot duplex should be more stable than any misbond, it seems likely that misbonded feet will detach from the track before fuel dissociation, and so misbonds are likely to constitute only a kinetic trap rather than a pathway to binding.

Partially binding in the correct fashion, however, may lead directly to full binding. In particular, the track has been seen to displace the fuel from the foot, although such a process is suppressed by the existence of mismatched bp between the track and foot that are absent in the fuel/foot duplex. To date, I have only observed displacement of the proximal half of the fuel by correct binding to site y , but it should be possible in other cases. The only exception would be when the foot binds to site v with the proximal half of the fuel still attached. In this case, 12 bp form with the track without displacing the fuel and so full displacement of more than two base pairs of the fuel is impossible. Nonetheless, one would expect the partial binding to the track to be more stable than the foot/fuel duplex. It therefore seems likely that once in this state, the fuel fragment would eventually detach, allowing complete binding.

Overall, the preliminary simulations seem to suggest that a foot with the proximal half of the fuel still attached will be more likely to overstep or step backwards than in a system with no fuel remnant. By contrast, a foot with the distal half of the fuel present seems to be likely to bind correctly to the site immediately in front of the attached foot, although overstepping is still possible. It would be beneficial to have greater experimental characterization of the behaviour of the nicking enzyme, but it seems likely that to preserve any forward-stepping bias it would be useful to ensure that the proximal half of the fuel detaches first. This might be achieved by using more AT base pairs in this half of the fuel strand, or alternatively moving the nicking site so that it is closer to the proximal end of the fuel.

8.2 Discussion

In this chapter I have presented the results of simulations into the operation of a two-footed DNA walker, the first time a simulation of a system such as this has been attempted. The model and simulation techniques were able to successfully describe a system this complex, and several novel predictions were made.

The model suggests that:

- In the original system introduced by Bath *et al.*, if the walker is released from a state with one foot raised, it can either bind directly to the front or back site, or initially misbond with the track. Misbonds will eventually melt or be displaced.
- The foot/track duplex can displace incorrect bonding via an ‘inchworm’ mechanism, or via a novel ‘double-X’ state.
- There is little or no bias for a raised foot to step forwards correctly.
- Applying a tension of around 15 pN to the track favours directly binding with the front site over the back site, and will tend to bias the walker towards stepping forwards.
- Applying tension has a very small effect on the competition between feet once both are bound.
- If the track is extended to include multiple binding sites, the most probable result is a single overstep (even under tension). The singly-overstepped state does not appear to be particularly susceptible to fraying from the front end (which would encourage fuel binding), meaning that it would be difficult for the walker to recover from such a configuration.
- A possible method by which the fuel could lift the front foot has been suggested, involving initial binding to the raised toehold before reaching back to displace the track from the foot.

- It is plausible that the one or other of the halves of the fuel will remain attached to the track after the other has detached. Early results suggest that if the half that is closest to the body of the walker remains, it will strongly inhibit binding to the desired site (whilst permitting backwards stepping and overstepping). By contrast, if the half of fuel that is furthest from walker's body is still present, the bias for stepping forward seems to be preserved, and overstepping is not strongly favoured compared to the case with no fuel present.

The majority of these predictions rely primarily on the basic geometrical, thermodynamic and mechanical properties of DNA, which the model reproduces reasonably well. In particular, it is difficult to see how the conclusion that overstepping is probable could be avoided, and similarly the consequences of having half of the fuel still attached seem physically robust.

The relative probability of reaching correct binding directly or through a displacement-based pathway is quite sensitive to the rates of different types of binding. The model does, however, highlight the possibility of previously unexpected binding pathways from a system with one raised foot (and no fuel attached). Although misbonding is still possible with fuel fragments present, the pathway to full binding is strongly suppressed.

My model predicts that tension will favour stepping forward primarily by increasing the rate at which correct pairing will occur with the front site relative to the back site. Note, however, that tension will also prevent the back site from displacing misbonds that occur with the front site. As a consequence, tension could increase any bias for stepping forwards regardless of which pathway (direct binding or via displacement) is more common in reality. The model's representation of a mechanism for lifting the front foot indicates that it is plausible, but the number of factors which contribute to the process mean that it should be treated with caution.

The 'double-X' configuration does not appear to put the DNA under unusual strain, but, due to the simplicity of the description of the backbone in the model, this predicted displacement mechanism can only be described as plausible. The inchworm method, however,

only requires the formation of bulged duplexes, which are known to exist.

The main caveat is that the walker, like the tweezers, often brings duplexes into reasonably close proximity. As a consequence, it is possible that electrostatic repulsion may have a significant role. Duplexes tend to be closest when in the fully bound state. In this case, electrostatic repulsion may tend to push the body of the walker away from the track, which would encourage more fraying than is observed in my simulations. Some misbonds also tend to involve looser, more open structures, and this means that they too might be penalized less by explicit electrostatics than correct bonding. If this is the case, the displacement route to correct bonding may be more common than observed in my simulations. Displacement itself, however, usually involves a higher density of strands (particularly when the double-X mechanism is involved), and so the effect on the kinetics is not obvious.

It is not easy to see whether electrostatics would penalize the initial states of binding to either the front or back site more than the other. If so, however, this effect would influence the bias for binding correctly to the front foot under tension. If electrostatic interactions jeopardize the performance of the walker, higher salt concentrations could be considered.

8.2.1 Considerations for design modifications

Having studied the walker using my model, several modifications to the design appear to be sensible. Most importantly, the tendency to overstep must be reduced. One approach would be to have two distinct toehold domains, two distinct non-toehold domains, and two fuel strands (see Figure 8.11). This would prevent overstepping by a single site, and would also have the benefit of suppressing the possible leak current suggested in this work.

The most stable misbonds (ignoring those which correspond to overstepping) arise because the toehold domain and non-toehold domain have regions of very similar sequence, and due to a repeated motif in the non-toehold domain. The behaviour of the system could be simplified by removing this similarity, as direct binding to the correct sites would dominate. It is not clear, however, that such a system would be any more efficient. Furthermore, if the fuel typically detaches in stages, the role of misbonds as intermediates to correct binding appears to be reduced. In the original work, the sequence was somewhat limited

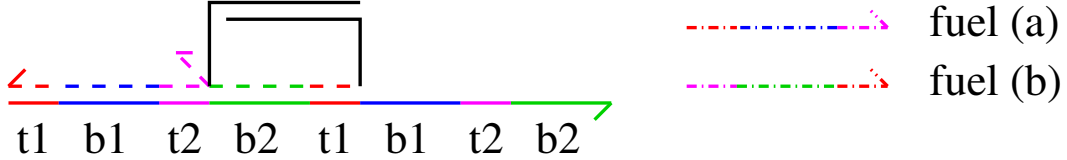


Figure 8.11: Schematic diagram of a proposed alternative walker design, with like colours indicating complimentarity. The track has two distinct competition domains (t_1 and t_2), and two distinct non-competition domains (b_1 and b_2). Two types of fuel are required - fuel (a) to lift the foot currently at the rear of the track, and fuel (b) to lift the foot currently towards the front.

by the desire to make the walker able to move in either direction. Lifting this requirement would give more sequence flexibility.

Applying tension to an extended track appears to be important, not only because it allows the direction of the walker to be defined. Firstly, it will assist with the recovery of walkers in a double-overstepped configuration (and reduce the frequency with which this occurs) by encouraging fraying from the front end. My model also suggests that tension will, to a limited degree, favour stepping forward. An additional benefit would be that the invasion of the foot/track duplex by other sites on the track [65] (which would effectively lead to an overstepped walker) would be strongly suppressed.

Finally, designing the fuel so that the half closest to the walker body is likely to detach first will probably improve the efficiency of the walker – this could be achieved by making it shorter than the other half, or including more AT base pairs. A remaining concern is whether the enzyme will detach before or after the fuel segments, and whether it will interfere with the reattachment to the track if it remains after one half of the fuel has dissociated.

Chapter 9

Conclusions

This thesis introduces a novel coarse-grained model of DNA which is specifically designed to capture the structural, mechanical and thermodynamic properties associated with single-stranded and duplex DNA, and the transition between the two. The model has been shown to give a physically reasonable description of a range of DNA properties, and has been applied to two nanotechnological systems (DNA tweezers and a two-footed DNA walker). These studies represent the first investigations into dynamic DNA devices using a coarse-grained model.

Unlike many alternatives in the literature, my model aims to reproduce DNA behaviour in a manner which is as physically motivated as possible. In particular, the structure of duplexes is enforced through the directionality of stacking and hydrogen-bonding interactions, rather than through backbone constraints. This approach allows for a stacking transition for single strands (which is absent in the majority of alternative models). Such a transition allows single strands to be extremely flexible when unstacked, and makes the simulation of hairpins and more complex motifs possible. To implement directional interactions, it was found to be important to represent the planarity of bases in the model, another feature that was novel when the model was introduced.

9.1 Utility of the model

In Chapters 5 and 6, the model's properties were discussed in great detail. I have shown that a model which is fairly simple in principle can give a good description of a wide range of

DNA properties (a far wider range than has been considered by other authors). Specifically, my model has been shown to:

- Give an approximate representation of the structure of B-DNA.
- Have a broad, almost uncooperative stacking transition.
- Reproduce the average melting temperatures and transition widths of short duplexes extremely well (when compared to the SantaLucia parameterization of the two-state model [91]).
- Give a reasonable representation of the stability of structures involving hairpins, bulges, mismatches and dangling ends. In particular, the dependence of hairpin stability on loop and stem size agrees well with Reference [91].
- Quantitatively reproduce the tendency of stacked single strands to be stiffer than unstacked ssDNA, with duplex DNA seen to be even less flexible.
- Give a good description of the response of duplexes and single strands to tension, and the response of dsDNA to applied torsion.

Many of the properties mentioned above were used to parameterize the model. Although it is noteworthy that such a range of features can be described by a model of this kind, to be really useful, it is important that the model makes predictions that extend beyond its parameterization (a feature somewhat lacking in the field of coarse-grained DNA modelling to date). The model reproduces some well-established physical properties of DNA that were in no way considered in parameterization. For example, the propellor twist of base pairs arises naturally in the model for the same reason as in real DNA. Similarly, the dependence of T_m on motif location for duplexes with mismatches was initially surprising, but actually reflects experimentally reported behaviour.

In addition, the model also makes non-trivial predictions, which are based on sufficiently generic properties that they should prove robust to the approximations present in my description of DNA. Below I highlight some of these predictions that may be amenable to experimental investigation:

- The model suggests that at low temperatures, the magnitude of the enthalpy associated with duplex formation should increase with temperature, as has been observed experimentally. At high enough temperatures, however, the model predicts that duplex fraying will cause this gradient to change sign. Such a feature should be particularly easy to detect for a duplex with a CG-rich core and AT bases towards the ends, as fraying will be significant close to the melting temperature of the duplex.
- It was found that the strand displacement process during the operation of DNA tweezers was far from flat in free energy, as is generally assumed. In part this was due to specific features of the system, but a component was due to the unfavourability of opening up a second single-stranded region when displacement begins, a feature which should apply to all displacement processes. An initial increase in free energy during displacement may help to explain the observed dependence of displacement rate on toehold length, the strength of which is currently difficult to justify [215].
- Simulation of a two-footed walker on a track demonstrated a range of non-trivial behaviour. Most significantly, it was found that the current design would typically lead to overstepping, an extremely undesirable event. The possibility that rebinding may occur before both halves of the hydrolyzed fuel have detached may also tend to favour overstepping and stepping backwards. Finally, applying a tension to the track was shown to improve the probability of stepping forwards, assuming the difficulties mentioned above can be overcome. Potential design changes to eliminate these problems are discussed in Chapter 8.

9.2 Limitations of the model

Any coarse-grained model is a compromise, and my description of DNA certainly neglects or heavily simplifies many features of potential importance. In particular, the simplistic representation of electrostatic interactions is probably the major factor limiting the predictive power of the model at this stage. This is particularly true for much of nanotechnology, which

often involves duplexes being brought into close proximity (DNA origami, for instance, is a dense array of double helices).

The next most obvious simplification is the limited sequence dependence of model. Features such as the tendency of bubbles to form in regions of weaker base-pairing cannot therefore, be captured – an average base description has the advantage, however, of not disguising general trends through sequence-specific effect.

The model’s structural and mechanical description of DNA on the level of one base is unlikely to be very accurate, as the representation of a nucleotide is necessarily simplistic and the model is fitted to long length scale properties. In particular, the simplicity of the backbone interaction between nucleotides may allow structures that in reality are penalized. Note also that minor and major grooving are absent in the model, asymmetrical groove sizes being important in determining the stresses involved in DNA origami [59].

9.3 Future work

The most obvious avenues for further work involve improving the current limitations of the model, by introducing a more physical representation of electrostatics, increased sequence dependence and exploring the description of DNA mechanics on the level of one base. None of these goals will be simple to achieve. The electrostatic behavior of DNA is potentially very complex, particularly for systems which involve divalent cations. In fact, an understanding of the consequences of electrostatics for the stability of systems like DNA origami would be interesting in its own right. For instance, it is a puzzle as to why ‘CANDO’ (a package with no explicit electrostatics) is able to reproduce the structures of complex 3-dimensional DNA origami [38]. Introducing sequence dependence of all interactions to the model would result in an enormous number of parameters, which would be difficult to constrain given the experimental data available (this is particularly true of stacking). The description of DNA on the level of a single base could potentially be compared to atomistic simulations, such as those of the Maddocks group [97], which would show which modes of deformation are too easy or too stiff in my model. It will not necessarily be trivial, however, to deduce

which interactions are responsible for the differences between force fields.

Despite the caveats mentioned in the previous section, I believe the model is capable of providing insight into a range of phenomena as it stands. For example, I am beginning to investigate the fundamental principles of strand displacement (in a range of scenarios) and duplex formation under tension, topics which were inspired by the simulations of nanotechnology presented in this thesis. Other authors have published work on the pathways involved in duplex formation [120, 121]: it would be instructive to see if I find similar results, or whether the results are strongly model-dependent.

Further development of the model, as outlined above, will enable more systems to be considered, such as DNA origami. From a broader perspective, it would be interesting to investigate more biological behaviour. In particular, the model could be used to explore how the forces and torques due to protein binding might induce biologically useful behaviour, such as gene regulation and the origin of replication in bacteria. Such work would require collaboration with researchers who study the detailed mechanics of DNA/protein binding [96]. Another possible extension would be to consider modelling RNA with an approach similar to that presented in this thesis. Such a model may be of use in predicting the secondary structure of biologically relevant RNAs, and could also provide information on the pathways by which these molecules fold to their native state.

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Appendix A

Representing forces and torques using quaternions

Langevin simulations in this work were performed using an algorithm which requires a quaternion representation of rigid body orientation [153]. The details of quaternion dynamics are discussed in Appendix B – here I present the forms of the derivatives which are required by the algorithm.

A.1 Nucleotide Description

Any set of coordinates that are used to construct the potential must specify the position and orientation of each nucleotide. Firstly, position can be specified by a vector \mathbf{r} which gives the location of the centre of mass of a nucleotide. Associated with each nucleotide there is also a unit vector which indicates the direction of the base sites from the backbone site (\mathbf{b}) and a unit vector giving the normal to the plane of the base (\mathbf{n}). When represented in the space-fixed frame, call them \mathbf{b}^s and \mathbf{n}^s , and in a body-fixed frame \mathbf{b}^b and \mathbf{n}^b . Define the body-fixed frame such that $b_i^b = \delta_{i,1}$ and $n_i^b = \delta_{i,3}$. The vectors can be mapped between frames using the rotation matrix:

$$\mathbf{x}^s(t) = A^T(t)\mathbf{x}^b. \quad (\text{A.1})$$

The rotation matrix A gives the orientation of the rigid body in the space-fixed frame. A possesses only three independent degrees of freedom, and so can be uniquely specified by a four-dimensional unit vector. Defining a quaternion $\mathbf{q} = (q_0, q_1, q_2, q_3)$, any rotation matrix

can be constructed via:

$$A = \begin{pmatrix} q_0^2 + q_1^2 - q_2^2 - q_3^2 & 2(q_1 q_2 + q_0 q_3) & 2(q_1 q_3 - q_0 q_2) \\ 2(q_1 q_2 - q_0 q_3) & q_0^2 - q_1^2 + q_2^2 - q_3^2 & 2(q_2 q_3 + q_0 q_1) \\ 2(q_1 q_3 + q_0 q_2) & 2(q_2 q_3 - q_0 q_1) & q_0^2 - q_1^2 - q_2^2 + q_3^2 \end{pmatrix}, \quad (\text{A.2})$$

with $\sum_i q_i^2 = 1$. An explicit discussion of the relationship between \mathbf{q} and the Euler angles of a rotation is given in Reference [217]. The quaternion \mathbf{q} is then a convenient representation of the orientation of the nucleotide. Using the quaternion representation of orientation:

- Backbone-base orientation:

$$\mathbf{b}^s = A^T(t) \mathbf{b}^b = \begin{pmatrix} q_0^2 + q_1^2 - q_2^2 - q_3^2 \\ 2(q_1 q_2 + q_0 q_3) \\ 2(q_1 q_3 - q_0 q_2) \end{pmatrix}. \quad (\text{A.3})$$

- Normal orientation:

$$\mathbf{n}^s = A^T(t) \mathbf{n}^b = \begin{pmatrix} 2(q_1 q_3 + q_0 q_2) \\ 2(q_2 q_3 - q_0 q_1) \\ (q_0^2 - q_1^2 - q_2^2 + q_3^2) \end{pmatrix}. \quad (\text{A.4})$$

- Third axis (defined by $\mathbf{n} \times \mathbf{b}$, useful for the chiral term of stacking):

$$\mathbf{y}^s = A^T(t) \mathbf{y}^b = \begin{pmatrix} 2(q_1 q_2 - q_0 q_3) \\ q_0^2 - q_1^2 + q_2^2 - q_3^2 \\ 2(q_2 q_3 + q_0 q_1) \end{pmatrix}. \quad (\text{A.5})$$

- Interaction site position:

$$\mathbf{r}_X = \mathbf{r} + d_X A^T(t) \mathbf{b}^b = \mathbf{r} + d_X \begin{pmatrix} q_0^2 + q_1^2 - q_2^2 - q_3^2 \\ 2(q_1 q_2 + q_0 q_3) \\ 2(q_1 q_3 - q_0 q_2) \end{pmatrix}, \quad (\text{A.6})$$

where d_x is the distance of the interaction site x from the centre of mass along the \mathbf{b} direction.

A.2 Derivatives

The potential of the model is described in Chapter 2. As outlined in Appendix B, derivatives with respect to the centre of mass position and quaternion of each nucleotide are required to perform Langevin simulations. The potential is a sum over pairwise interactions, and for each interaction one must differentiate the potential with respect to the position vector and orientation quaternion for both nucleotides involved.

For a nearest-neighbour interaction:

$$\begin{aligned}
 & \frac{dV_{\text{FENE}}(\delta r_{\text{backbone}})}{d\delta r_{\text{backbone}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{backbone}}) + \sum_{x,y^*} \frac{df_3(\delta r_{xy})}{d\delta r_{xy}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{xy}) \\
 & + \frac{V_{\text{stack}}}{f_1(\delta r_{\text{stack}})} \frac{df_1(\delta r_{\text{stack}})}{d\delta r_{\text{stack}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{stack}}) + \sum_{i=4,5',6'} \frac{V_{\text{stack}}}{f_4(\theta_i)} \frac{df_4(\theta_i)}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
 \nabla_{\mathbf{q},\mathbf{r}} V_{\text{neighbour}} = & \\
 & + \frac{V_{\text{stack}}}{f_5(\cos(\phi_1))} \frac{df_5(\cos(\phi_1))}{d\cos(\phi_1)} \nabla_{\mathbf{q},\mathbf{r}}(\cos(\phi_1)) \\
 & + \frac{V_{\text{stack}}}{f_5(\cos(\phi_2))} \frac{df_5(\cos(\phi_2))}{d\cos(\phi_2)} \nabla_{\mathbf{q},\mathbf{r}}(\cos(\phi_2)),
 \end{aligned} \tag{A.7}$$

where the sum over x, y^* means all combinations of repulsion centres except for the two backbone sites. For non-neighbouring nucleotides,

$$\begin{aligned}
 & \sum_{x,y} \frac{df_3(\delta r_{xy})}{d\delta r_{x,y}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{xy}) + \frac{V_{\text{HB}}}{f_1(\delta r_{\text{HB}})} \frac{df_1(\delta r_{\text{HB}})}{d\delta r_{\text{HB}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{HB}}) \\
 & + \sum_{i=1-4,7,8} \frac{V_{\text{HB}}}{f_4(\theta_i)} \frac{df_4(\theta_i)}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
 & + \frac{V_{\text{cross_stack}}}{f_2(\delta r_{\text{HB}})} \frac{df_2(\delta r_{\text{HB}})}{d\delta r_{\text{HB}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{HB}}) + \sum_{i=1}^3 \frac{V_{\text{cross_stack}}}{f_4(\theta_i)} \frac{df_4(\theta_i)}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
 & + \sum_{i=1,7,8} \frac{V_{\text{cross_stack}}}{f_4(\theta_i)+f_4(\pi-\theta_i)} \frac{d(f_4(\theta_i)+f_4(\pi-\theta_i))}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
 \nabla_{\mathbf{q},\mathbf{r}} V_{\text{non-neighbour}} = & \\
 & + \frac{V_{\text{coax_stack}}}{f_2(\delta r_{\text{stack}})} \frac{df_2(\delta r_{\text{stack}})}{d\delta r_{\text{stack}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{stack}}) + \frac{V_{\text{coax_stack}}}{f_4(\theta_4)} \frac{df_4(\theta_4)}{d\theta_4} \nabla_{\mathbf{q},\mathbf{r}}(\theta_4) \\
 & + \frac{V_{\text{coax_stack}}}{f_4(\theta_1)+f_4(2\pi-\theta_1)} \frac{d(f_4(\theta_1)+f_4(2\pi-\theta_1))}{d\theta_1} \nabla_{\mathbf{q},\mathbf{r}}(\theta_1) \\
 & + \sum_{i=5}^6 \frac{V_{\text{coax_stack}}}{f_4(\theta_i)+f_4(\pi-\theta_i)} \frac{d(f_4(\theta_i)+f_4(\pi-\theta_i))}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
 & + \sum_{i=3}^4 \frac{V_{\text{coax_stack}}}{f_4(\cos(\phi_i))} \frac{df_4(\cos(\phi_i))}{d\cos(\phi_i)} \nabla_{\mathbf{q},\mathbf{r}} \cos(\phi_i),
 \end{aligned} \tag{A.8}$$

where in this case the sum over x, y means all combinations of repulsion centres of the two nucleotides. For each instance of an interaction, the parameters such as strength and cutoffs are constant and so they have been suppressed in the equations above.

A.2.1 Derivatives of functional forms

Equations A.7 and A.8 contain derivatives of the functional forms V_{FENE} and f_i with respect to their arguments, which are generally angles or distances. These are given by:

$$\frac{dV_{\text{FENE}}(r)}{dr} = \frac{\epsilon(r - r^0)}{\Delta^2 - (r - r^0)^2}. \tag{A.9}$$

$$\frac{df_1(r)}{dr} = \begin{cases} 2\epsilon a \exp(-(r - r^0)a)(1 - \exp(-(r - r^0)a)) & \text{if } r^{low} < r < r^{high}, \\ 2\epsilon b_{low}(r - r^{c,low}), & \text{if } r^{c,low} < r < r^{low}, \\ 2\epsilon b_{high}(r - r^{c,high}) & \text{if } r^{high} < r < r^{c,high}, \\ 0 & \text{otherwise.} \end{cases} \quad (\text{A.10})$$

$$\frac{df_2(r)}{dr} = \begin{cases} k(r - r^0) & \text{if } r^{low} < r < r^{high}, \\ 2kb_{low}(r - r^{c,low}), & \text{if } r^{c,low} < r < r^{low}, \\ 2kb_{high}(r - r^{c,high}) & \text{if } r^{high} < r < r^{c,high}, \\ 0 & \text{otherwise.} \end{cases} \quad (\text{A.11})$$

$$\frac{df_3(r)}{dr} = \begin{cases} \frac{4\epsilon}{\sigma} \left(6 \left(\frac{\sigma}{r} \right)^7 - 12 \left(\frac{\sigma}{r} \right)^{13} \right) & \text{if } r < r^*, \\ 2\epsilon b(r - r^c) & \text{if } r^* < r < r^c, \\ 0 & \text{otherwise.} \end{cases} \quad (\text{A.12})$$

$$\frac{df_4(\theta)}{d\theta} = \begin{cases} -2a(\theta - \theta^0) & \text{if } \theta^0 - \Delta\theta^* < \theta < \theta^0 + \Delta\theta^*, \\ 2b(\theta - (\theta^0 - \Delta\theta^c)) & \text{if } \theta^0 - \Delta\theta^c < \theta < \theta^0 - \Delta\theta^*, \\ 2b(\theta - (\theta^0 + \Delta\theta^c)) & \text{if } \theta^0 + \Delta\theta^* < \theta < \theta^0 + \Delta\theta^c, \\ 0 & \text{otherwise.} \end{cases} \quad (\text{A.13})$$

$$\frac{df_5(\cos(\phi))}{d\cos(\phi)} = \begin{cases} -2a \cos(\phi) & \text{if } \cos(\phi)^* < \cos(\phi) < 0, \\ 2b(\cos(\phi) - \cos(\phi)^c) & \text{if } \cos(\phi)^c < \cos(\phi) < \cos(\phi)^*, \\ 0 & \text{otherwise.} \end{cases} \quad (\text{A.14})$$

A.2.2 Derivatives with respect to the coordinates

The somewhat clumsy definitions of angles in Chapter 2 are now extremely useful, as it is natural to consider one nucleotide as being influenced by another when calculating the force or torque. Label the nucleotide for which the force or torque is being calculated as α , and its partner as β . Define $\delta\mathbf{r}_{xy}$ to be the vector **to** the interaction site x on α **from** site y on β . Let $\delta\hat{\mathbf{r}}$ be a normalized vector, and let $\tilde{\mathbf{b}}$, $\tilde{\mathbf{n}}$ and $\tilde{\mathbf{y}}$ represent the orientation of β . θ_2 , θ_5 , θ_7 , ϕ_1 and ϕ_3 involve angles with respect to the orientation of α , and θ_3 , θ_6 , θ_8 , ϕ_2 and ϕ_4 to involve angles with the orientation of β . The stacking interaction is intentionally asymmetric with respect to the 3' to 5' direction for neighbouring nucleotides. The result is that $\nabla_{\mathbf{r},\mathbf{q}}(\theta_{5'})$, $\nabla_{\mathbf{r},\mathbf{q}}(\theta_{6'})$, $\nabla_{\mathbf{r},\mathbf{q}}(\cos(\phi_1))$ and $\nabla_{\mathbf{r},\mathbf{q}}(\cos(\phi_2))$ all have a sign that depends on whether α is in the 5' direction of β : if it is, then the first option is taken, if not, then the alternative.

Derivatives with respect to position vectors

$$\nabla_{\mathbf{r}}(r_{xy}) = \delta \hat{\mathbf{r}}_{xy}. \quad (\text{A.15})$$

$$\nabla_{\mathbf{r}}(\theta_1) = 0. \quad (\text{A.16})$$

$$\nabla_{\mathbf{r}}(\theta_2) = \frac{1}{\sqrt{1 - (\mathbf{b} \cdot \delta \hat{\mathbf{r}}_{\text{HB}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{HB}}|} (\mathbf{b} - (\mathbf{b} \cdot \delta \hat{\mathbf{r}}_{\text{HB}}) \delta \hat{\mathbf{r}}_{\text{HB}}) \quad (\text{A.17})$$

$$\nabla_{\mathbf{r}}(\theta_3) = \frac{-1}{\sqrt{1 - (\tilde{\mathbf{b}} \cdot \delta \hat{\mathbf{r}}_{\text{HB}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{HB}}|} (\tilde{\mathbf{b}} - (\tilde{\mathbf{b}} \cdot \delta \hat{\mathbf{r}}_{\text{HB}}) \delta \hat{\mathbf{r}}_{\text{HB}}) \quad (\text{A.18})$$

$$\nabla_{\mathbf{r}}(\theta_4) = 0 \quad (\text{A.19})$$

$$\nabla_{\mathbf{r}}(\theta_5) = \frac{1}{\sqrt{1 - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{stack}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} (\mathbf{n} - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{stack}}) \delta \hat{\mathbf{r}}_{\text{stack}}) \quad (\text{A.20})$$

$$\nabla_{\mathbf{r}}(\theta_6) = \frac{-1}{\sqrt{1 - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{stack}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} (\tilde{\mathbf{n}} - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{stack}}) \delta \hat{\mathbf{r}}_{\text{stack}}) \quad (\text{A.21})$$

$$\nabla_{\mathbf{r}}(\theta_{5'}) = \frac{\mp 1}{\sqrt{1 - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{stack}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} (\mathbf{n} - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{stack}}) \delta \hat{\mathbf{r}}_{\text{stack}}) \quad (\text{A.22})$$

$$\nabla_{\mathbf{r}}(\theta_{6'}) = \frac{\mp 1}{\sqrt{1 - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{stack}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} (\tilde{\mathbf{n}} - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{stack}}) \delta \hat{\mathbf{r}}_{\text{stack}}) \quad (\text{A.23})$$

$$\nabla_{\mathbf{r}}(\theta_7) = \frac{1}{\sqrt{1 - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{HB}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{HB}}|} (\mathbf{n} - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{HB}}) \delta \hat{\mathbf{r}}_{\text{HB}}) \quad (\text{A.24})$$

$$\nabla_{\mathbf{r}}(\theta_8) = \frac{-1}{\sqrt{1 - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{HB}})^2}} \frac{1}{|\delta \mathbf{r}_{\text{HB}}|} (\tilde{\mathbf{n}} - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{HB}}) \delta \hat{\mathbf{r}}_{\text{HB}}) \quad (\text{A.25})$$

$$\nabla_{\mathbf{r}}(\cos(\phi_1)) = \frac{\pm 1}{|\delta \mathbf{r}_{\text{backbone}}|} (\mathbf{y} - (\mathbf{y} \cdot \delta \hat{\mathbf{r}}_{\text{backbone}}) \delta \hat{\mathbf{r}}_{\text{backbone}}) \quad (\text{A.26})$$

$$\nabla_{\mathbf{r}}(\cos(\phi_2)) = \frac{\pm 1}{|\delta\mathbf{r}_{\text{backbone}}|} (\tilde{\mathbf{y}} - (\tilde{\mathbf{y}} \cdot \delta\hat{\mathbf{r}}_{\text{backbone}}) \delta\hat{\mathbf{r}}_{\text{backbone}}) \quad (\text{A.27})$$

$$\begin{aligned} \nabla_{\mathbf{r}}(\cos(\phi_3)) &= -\left(\frac{\delta\mathbf{r}_{\text{stack}}}{|\delta\mathbf{r}_{\text{stack}}|^2} + \frac{\delta\mathbf{r}_{\text{backbone}}}{|\delta\mathbf{r}_{\text{backbone}}|^2}\right) ((\delta\hat{\mathbf{r}}_{\text{stack}} \times \delta\hat{\mathbf{r}}_{\text{backbone}}) \cdot \mathbf{b}) \\ &\quad + \frac{\mathbf{b} \times (\delta\mathbf{r}_{\text{stack}} - \delta\mathbf{r}_{\text{backbone}})}{|\delta\mathbf{r}_{\text{stack}}||\delta\mathbf{r}_{\text{backbone}}|} \end{aligned} \quad (\text{A.28})$$

$$\begin{aligned} \nabla_{\mathbf{r}}(\cos(\phi_4)) &= -\left(\frac{\delta\mathbf{r}_{\text{stack}}}{|\delta\mathbf{r}_{\text{stack}}|^2} + \frac{\delta\mathbf{r}_{\text{backbone}}}{|\delta\mathbf{r}_{\text{backbone}}|^2}\right) \left((\delta\hat{\mathbf{r}}_{\text{stack}} \times \delta\hat{\mathbf{r}}_{\text{backbone}}) \cdot \tilde{\mathbf{b}}\right) \\ &\quad + \frac{\tilde{\mathbf{b}} \times (\delta\mathbf{r}_{\text{stack}} - \delta\mathbf{r}_{\text{backbone}})}{|\delta\mathbf{r}_{\text{stack}}||\delta\mathbf{r}_{\text{backbone}}|} \end{aligned} \quad (\text{A.29})$$

Derivatives with respect to orientation quaternions

$$\nabla_{\mathbf{q}}(\delta r_{xy}) = \delta\hat{\mathbf{r}}_{xy} \cdot \frac{\partial \delta\mathbf{r}_{xy}}{\partial \mathbf{q}} \quad (\text{A.30})$$

$$\nabla_{\mathbf{q}}(\theta_1) = \frac{-1}{\sqrt{1 - (\mathbf{b}^s \cdot \tilde{\mathbf{b}}^s)^2}} \tilde{\mathbf{b}} \cdot \frac{\partial \mathbf{b}}{\partial \mathbf{q}} \quad (\text{A.31})$$

$$\nabla_{\mathbf{q}}(\theta_2) = \frac{1}{\sqrt{1 - (\mathbf{b} \cdot \delta\hat{\mathbf{r}}_{\text{HB}})^2}} \left(\frac{\partial \mathbf{b}}{\partial \mathbf{q}} \cdot \delta\hat{\mathbf{r}}_{\text{HB}} + \frac{1}{|\delta\mathbf{r}_{\text{HB}}|} \mathbf{b} \cdot \frac{\partial \delta\mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} - \frac{1}{|\delta\mathbf{r}_{\text{HB}}|^3} \mathbf{b} \cdot \delta\mathbf{r}_{\text{HB}} \left(\delta\mathbf{r}_{\text{HB}} \cdot \frac{\partial \delta\mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} \right) \right) \quad (\text{A.32})$$

$$\nabla_{\mathbf{q}}(\theta_3) = \frac{-1}{\sqrt{1 - (\tilde{\mathbf{b}} \cdot \delta\hat{\mathbf{r}}_{\text{HB}})^2}} \left(\frac{1}{|\delta\mathbf{r}_{\text{HB}}|} \tilde{\mathbf{b}} \cdot \frac{\partial \delta\mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} - \frac{1}{|\delta\mathbf{r}_{\text{HB}}|^3} \tilde{\mathbf{b}} \cdot \delta\mathbf{r}_{\text{HB}} \left(\delta\mathbf{r}_{\text{HB}} \cdot \frac{\partial \delta\mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} \right) \right) \quad (\text{A.33})$$

$$\nabla_{\mathbf{q}}(\theta_4) = \frac{-1}{\sqrt{1 - (\mathbf{n} \cdot \tilde{\mathbf{n}})^2}} \tilde{\mathbf{n}} \cdot \frac{\partial \mathbf{n}}{\partial \mathbf{q}} \quad (\text{A.34})$$

$$\nabla_{\mathbf{q}}(\theta_5) = \frac{1}{\sqrt{1 - (\mathbf{n} \cdot \delta\hat{\mathbf{r}}_{\text{stack}})^2}} \left(\begin{array}{l} \frac{\partial \mathbf{n}}{\partial \mathbf{q}} \cdot \delta\hat{\mathbf{r}}_{\text{stack}} + \frac{1}{|\delta\mathbf{r}_{\text{stack}}|} \mathbf{n} \cdot \frac{\partial \delta\mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \\ - \frac{1}{|\delta\mathbf{r}_{\text{stack}}|^3} \mathbf{n} \cdot \delta\mathbf{r}_{\text{stack}} \left(\delta\mathbf{r}_{\text{stack}} \cdot \frac{\partial \delta\mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \right) \end{array} \right) \quad (\text{A.35})$$

$$\nabla_{\mathbf{q}}(\theta_6) = \frac{-1}{\sqrt{1 - (\tilde{\mathbf{n}} \cdot \delta\hat{\mathbf{r}}_{\text{stack}})^2}} \left(\frac{1}{|\delta\mathbf{r}_{\text{stack}}|} \tilde{\mathbf{n}} \cdot \frac{\partial \delta\mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} - \frac{1}{|\delta\mathbf{r}_{\text{stack}}|^3} \tilde{\mathbf{n}} \cdot \delta\mathbf{r}_{\text{stack}} \left(\delta\mathbf{r}_{\text{stack}} \cdot \frac{\partial \delta\mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \right) \right) \quad (\text{A.36})$$

$$\nabla_{\mathbf{q}}(\theta_{5'}) = \frac{\mp}{\sqrt{1 - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{stack}})^2}} \begin{pmatrix} \frac{\partial \mathbf{n}}{\partial \mathbf{q}} \cdot \delta \hat{\mathbf{r}}_{\text{stack}} + \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} \mathbf{n} \cdot \frac{\partial \delta \mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \\ - \frac{1}{|\delta \mathbf{r}_{\text{stack}}|^3} \mathbf{n} \cdot \delta \mathbf{r}_{\text{stack}} \left(\delta \mathbf{r}_{\text{stack}} \cdot \frac{\partial \delta \mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \right) \end{pmatrix} \quad (\text{A.37})$$

$$\nabla_{\mathbf{q}}(\theta_{6'}) = \frac{\mp}{\sqrt{1 - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{stack}})^2}} \left(\frac{1}{|\delta \mathbf{r}_{\text{stack}}|} \tilde{\mathbf{n}} \cdot \frac{\partial \delta \mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} - \frac{1}{|\delta \mathbf{r}_{\text{stack}}|^3} \tilde{\mathbf{n}} \cdot \delta \mathbf{r}_{\text{stack}} \left(\delta \mathbf{r}_{\text{stack}} \cdot \frac{\partial \delta \mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \right) \right) \quad (\text{A.38})$$

$$\nabla_{\mathbf{q}}(\theta_7) = \frac{1}{\sqrt{1 - (\mathbf{n} \cdot \delta \hat{\mathbf{r}}_{\text{HB}})^2}} \left(\frac{\partial \mathbf{n}}{\partial \mathbf{q}} \cdot \delta \hat{\mathbf{r}}_{\text{HB}} + \frac{1}{|\delta \mathbf{r}_{\text{HB}}|} \mathbf{n} \cdot \frac{\partial \delta \mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} - \frac{1}{|\delta \mathbf{r}_{\text{HB}}|^3} \mathbf{n} \cdot \delta \mathbf{r}_{\text{HB}} \left(\delta \mathbf{r}_{\text{HB}} \cdot \frac{\partial \delta \mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} \right) \right) \quad (\text{A.39})$$

$$\nabla_{\mathbf{q}}(\theta_8) = \frac{-1}{\sqrt{1 - (\tilde{\mathbf{n}} \cdot \delta \hat{\mathbf{r}}_{\text{HB}})^2}} \left(\frac{1}{|\delta \mathbf{r}_{\text{HB}}|} \tilde{\mathbf{n}} \cdot \frac{\partial \delta \mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} - \frac{1}{|\delta \mathbf{r}_{\text{HB}}|^3} \tilde{\mathbf{n}} \cdot \delta \mathbf{r}_{\text{HB}} \left(\delta \mathbf{r}_{\text{HB}} \cdot \frac{\partial \delta \mathbf{r}_{\text{HB}}}{\partial \mathbf{q}} \right) \right) \quad (\text{A.40})$$

$$\begin{aligned} \nabla_{\mathbf{q}}(\cos(\phi_1)) &= \pm \left(\frac{\partial \mathbf{y}}{\partial \mathbf{q}} \cdot \delta \hat{\mathbf{r}}_{\text{backbone}} + \frac{1}{|\delta \mathbf{r}_{\text{backbone}}|} \mathbf{y} \cdot \frac{\partial \delta \mathbf{r}_{\text{backbone}}}{\partial \mathbf{q}} \right) \\ &\mp \frac{1}{|\delta \mathbf{r}_{\text{backbone}}|^3} \mathbf{y} \cdot \delta \mathbf{r}_{\text{backbone}} \left(\delta \mathbf{r}_{\text{backbone}} \cdot \frac{\partial \delta \mathbf{r}_{\text{backbone}}}{\partial \mathbf{q}} \right) \end{aligned} \quad (\text{A.41})$$

$$\begin{aligned} \nabla_{\mathbf{q}}(\cos(\phi_2)) &= \pm \frac{1}{|\delta \mathbf{r}_{\text{backbone}}|} \tilde{\mathbf{y}} \cdot \frac{\partial \delta \mathbf{r}_{\text{backbone}}}{\partial \mathbf{q}} \\ &\mp \frac{1}{|\delta \mathbf{r}_{\text{backbone}}|^3} \tilde{\mathbf{y}} \cdot \delta \mathbf{r}_{\text{backbone}} \left(\delta \mathbf{r}_{\text{backbone}} \cdot \frac{\partial \delta \mathbf{r}_{\text{backbone}}}{\partial \mathbf{q}} \right) \end{aligned} \quad (\text{A.42})$$

$$\begin{aligned} \nabla_{\mathbf{q}}(\cos(\phi_3)) &= -((\delta \hat{\mathbf{r}}_{\text{stack}} \times \delta \hat{\mathbf{r}}_{\text{backbone}}) \cdot \mathbf{b}) \left(\frac{\delta \mathbf{r}_{\text{stack}} \cdot \partial \delta \mathbf{r}_{\text{stack}} / \partial \mathbf{q}}{|\delta \mathbf{r}_{\text{stack}}|^2} + \frac{\delta \mathbf{r}_{\text{backbone}} \cdot \partial \delta \mathbf{r}_{\text{backbone}} / \partial \mathbf{q}}{|\delta \mathbf{r}_{\text{backbone}}|^2} \right) \\ &+ \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} \frac{1}{|\delta \mathbf{r}_{\text{backbone}}|} \left(\left(\frac{\partial \delta \mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \times \delta \mathbf{r}_{\text{backbone}} - \frac{\partial \delta \mathbf{r}_{\text{backbone}}}{\partial \mathbf{q}} \times \delta \mathbf{r}_{\text{stack}} \right) \cdot \mathbf{b} \right) \\ &+ \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} \frac{1}{|\delta \mathbf{r}_{\text{backbone}}|} \left(\frac{\partial \mathbf{b}}{\partial \mathbf{q}} \cdot (\delta \mathbf{r}_{\text{stack}} \times \delta \mathbf{r}_{\text{backbone}}) \right) \end{aligned} \quad (\text{A.43})$$

$$\begin{aligned} \nabla_{\mathbf{q}}(\cos(\phi_4)) &= -((\delta \hat{\mathbf{r}}_{\text{stack}} \times \delta \hat{\mathbf{r}}_{\text{backbone}}) \cdot \tilde{\mathbf{b}}) \left(\frac{\delta \mathbf{r}_{\text{stack}} \cdot \partial \delta \mathbf{r}_{\text{stack}} / \partial \mathbf{q}}{|\delta \mathbf{r}_{\text{stack}}|^2} + \frac{\delta \mathbf{r}_{\text{backbone}} \cdot \partial \delta \mathbf{r}_{\text{backbone}} / \partial \mathbf{q}}{|\delta \mathbf{r}_{\text{backbone}}|^2} \right) \\ &+ \frac{1}{|\delta \mathbf{r}_{\text{stack}}|} \frac{1}{|\delta \mathbf{r}_{\text{backbone}}|} \left(\left(\frac{\partial \delta \mathbf{r}_{\text{stack}}}{\partial \mathbf{q}} \times \delta \mathbf{r}_{\text{backbone}} - \frac{\partial \delta \mathbf{r}_{\text{backbone}}}{\partial \mathbf{q}} \times \delta \mathbf{r}_{\text{stack}} \right) \cdot \tilde{\mathbf{b}} \right) \end{aligned} \quad (\text{A.44})$$

Useful explicit forms of quaternion derivatives

Many of the expressions in the previous section contain terms such as $\frac{\partial \delta \mathbf{r}_{xy}}{\partial \mathbf{q}}$. These are actually matrices, with the explicit forms given below.

$$M_{ij} = \frac{\partial(\delta r_{xy})_i}{\partial q_j} = 2d_x \begin{pmatrix} q_0 & q_1 & -q_2 & -q_3 \\ q_3 & q_2 & q_1 & q_0 \\ -q_2 & q_3 & -q_0 & q_1 \end{pmatrix}. \quad (\text{A.45})$$

$$M_{ij} = \frac{\partial n_i}{\partial q_j} = 2 \begin{pmatrix} q_2 & q_3 & q_0 & q_1 \\ -q_1 & -q_0 & q_3 & q_2 \\ q_0 & -q_1 & -q_2 & q_3 \end{pmatrix}. \quad (\text{A.46})$$

$$M_{ij} = \frac{\partial y_i}{\partial q_j} = 2 \begin{pmatrix} -q_3 & q_2 & q_1 & -q_0 \\ q_0 & -q_1 & q_2 & -q_3 \\ q_1 & q_0 & q_3 & q_2 \end{pmatrix}. \quad (\text{A.47})$$

$$M_{ij} = \frac{\partial b_i}{\partial q_j} = 2 \begin{pmatrix} q_0 & q_1 & -q_2 & -q_3 \\ q_3 & q_2 & q_1 & q_0 \\ -q_2 & q_3 & -q_0 & q_1 \end{pmatrix}. \quad (\text{A.48})$$

Appendix B

Quaternion dynamics

The algorithm of Davidchack *et al.* [153] was used for the Langevin simulations in this thesis. This algorithm extends the method of Miller *et al.* [218], which generates Newtonian motion of rigid bodies described by quaternions. As neither paper fully explains the origin of the equations of motion, I derive them here from first principles.

B.1 Angular velocities represented in quaternions

In Appendix A, quaternions were defined in terms of a rotation matrix relating the orientation of a vector in the body-fixed frame \mathbf{x}^b to one in the space-fixed frame \mathbf{x}^s . Rapaport [217] has shown that the angular velocity of an object in the body-fixed frame is given by:

$$\begin{bmatrix} 0 \\ w_1^b \\ w_2^b \\ w_3^b \end{bmatrix} = 2 \begin{bmatrix} q_0 & q_1 & q_2 & q_3 \\ -q_1 & q_0 & q_3 & -q_2 \\ -q_2 & -q_3 & q_0 & q_1 \\ -q_3 & q_2 & -q_1 & q_0 \end{bmatrix} \begin{bmatrix} \dot{q}_0 \\ \dot{q}_1 \\ \dot{q}_2 \\ \dot{q}_3 \end{bmatrix}. \quad (\text{B.1})$$

I label the matrix in (B.1) as $S_{\mathbf{q}}^T$.

B.2 Motion without noise or damping

The full Lagrangian for my model, in the absence of any noise or damping, is given by:

$$L = \sum_i \left(\left(\frac{1}{2} m_i (\dot{\mathbf{r}}_i)^T (\dot{\mathbf{r}}_i) + \frac{2}{|\mathbf{q}_i|^4} (S_{\mathbf{q}_i}^T \dot{\mathbf{q}}_i)^T I_i (S_{\mathbf{q}_i}^T \dot{\mathbf{q}}_i) \right) + \lambda_i \left(\sum_j (q_i)_j (q_i)_j - 1 \right) \right) - U(\{\mathbf{q}\}, \{\mathbf{r}\}). \quad (\text{B.2})$$

The second term in Equation B.2 is simply the kinetic energy of rotation. At this stage, I have not fixed the normalization for each quaternion to unity, but instead imposed a

constraint term $\lambda_i \left(\sum_j (q_i)_j (q_i)_j - 1 \right)$ that will later be used to impose this condition on the trajectory. In this extended picture, I have introduced a fictional zeroth component of the moment of inertia tensor, so that the matrix is invertible:

$$I_i = \begin{bmatrix} (I_i)_0 & 0 & 0 & 0 \\ 0 & (I_i)_x & 0 & 0 \\ 0 & 0 & (I_i)_y & 0 \\ 0 & 0 & 0 & (I_i)_z \end{bmatrix}. \quad (\text{B.3})$$

I have also included a $1/|\mathbf{q}_i|^4$ term – I do this to make the Hamiltonian (when I derive it) look simpler. Eventually I will restrict myself to trajectories for which $|\mathbf{q}^i|^2 = 1$, and these added terms will have no consequences for the motion. The generalized momenta for this Lagrangian are:

$$\mathbf{p}_i = \frac{\partial L}{\partial \dot{\mathbf{r}}_i} = m_i \dot{\mathbf{r}}_i \quad (\text{B.4})$$

and

$$\boldsymbol{\Pi}_i = \frac{\partial L}{\partial \dot{\mathbf{q}}_i} = \frac{4}{|\mathbf{q}_i|^4} S_{\mathbf{q}_i} I_i S_{\mathbf{q}_i}^T \dot{\mathbf{q}}_i. \quad (\text{B.5})$$

Consequentially, the Hamiltonain can be shown to be:

$$\begin{aligned} \mathcal{H} = \sum_i & \left(\left(\frac{1}{2m_i} (\mathbf{p}_i)^T (\mathbf{p}_i) + \frac{1}{8} (S_{\mathbf{q}_i}^T \boldsymbol{\Pi}_i)^T (I_i)^{-1} (S_{\mathbf{q}_i}^T \boldsymbol{\Pi}_i) \right) - \lambda_i \left(\sum_j (\mathbf{q}_i)_j (\mathbf{q}_i)_j - 1 \right) \right) \\ & + U(\{\mathbf{q}\}, \{\boldsymbol{\Pi}\}). \end{aligned} \quad (\text{B.6})$$

Equations of motion can be determined via:

$$\dot{\mathbf{r}}_i = \frac{\partial}{\partial \mathbf{p}_i} \mathcal{H}, \quad \dot{\mathbf{p}}_i = -\frac{\partial}{\partial \mathbf{r}_i} \mathcal{H}, \quad \dot{\mathbf{q}}_i = \frac{\partial}{\partial \boldsymbol{\Pi}_i} \mathcal{H}, \quad \dot{\boldsymbol{\Pi}}_i = -\frac{\partial}{\partial \mathbf{q}_i} \mathcal{H}, \quad \frac{\partial}{\partial \lambda_i} \mathcal{H} = 0. \quad (\text{B.7})$$

The first two terms are trivial:

$$\dot{\mathbf{r}}_i = \frac{\mathbf{p}_i}{m_i} \quad \text{and} \quad \dot{\mathbf{p}}_i = -\frac{\partial \mathcal{U}}{\partial \mathbf{r}_i}. \quad (\text{B.8})$$

The next two terms are not so simple:

$$\dot{\mathbf{q}}_i = \frac{1}{4} S_{\mathbf{q}_i} I_i^{-1} S_{\mathbf{q}_i}^T \boldsymbol{\Pi}_i \quad (\text{B.9})$$

and

$$\dot{\boldsymbol{\Pi}}_i = -\frac{1}{8} \frac{\partial}{\partial \mathbf{q}_i} \left((S_{\mathbf{q}_i}^T \boldsymbol{\Pi}_i)^T (I_i)^{-1} (S_{\mathbf{q}_i}^T \boldsymbol{\Pi}_i) \right) + 2\lambda_i \mathbf{q}_i - \frac{\partial \mathcal{U}}{\partial \mathbf{q}_i}. \quad (\text{B.10})$$

Hence forth the sunscript i indicating the identity of the particle will be assumed, to avoid complication with index notation. I will also drop the \mathbf{q} label on $S_{\mathbf{q}}$. I can now use the $\frac{\partial}{\partial \lambda} \mathcal{H}$ term to impose $|\mathbf{q}|^2 = 1$, which means that $S_{\mathbf{q}} S_{\mathbf{q}}^T = 1$. The easiest way to eliminate λ is by evaluating the angular velocity ω . ω is given by:

$$\omega = 2S^T \dot{\mathbf{q}} = \frac{1}{2}(I)^{-1} S^T \boldsymbol{\Pi}, \quad (\text{B.11})$$

in which I have used Equation B.9, and the fact that S is orthorgonal once $|\mathbf{q}|^2 = 1$ is imposed. Therefore

$$\dot{\omega} = \frac{1}{2}(I)^{-1} \frac{d}{dt} (S^T \boldsymbol{\Pi}). \quad (\text{B.12})$$

Using $\dot{\boldsymbol{\Pi}} = -\frac{\partial}{\partial \mathbf{q}^i} \mathcal{H}$,

$$\frac{d}{dt} (S^T \boldsymbol{\Pi}) = \dot{S}^T \boldsymbol{\Pi} + S^T \left(-\frac{1}{8} \frac{\partial}{\partial \mathbf{q}} \left((S^T \boldsymbol{\Pi})^T (I)^{-1} (S^T \boldsymbol{\Pi}) \right) + 2\lambda \mathbf{q} - \frac{\partial U}{\partial \mathbf{q}} \right). \quad (\text{B.13})$$

The best way to treat these equations is to use tensor notation. Defining the third rank tensor $D_{\sigma\nu\rho}$ by:

$$D_{\sigma\nu\rho} \dot{q}_\sigma = (\dot{S}^T)_{\nu\rho}, \quad (\text{B.14})$$

and $F_{\rho\gamma\beta}$ by:

$$F_{\rho\gamma\beta} = \frac{\partial S_{\gamma\beta}}{\partial q_\rho}, \quad (\text{B.15})$$

I can express Equations B.12 and B.13 as:

$$\begin{aligned} \dot{\omega}_\mu &= \frac{1}{2}(I^{-1})_{\mu\nu} [D_{\sigma\nu\rho} \dot{q}_\sigma \Pi_\rho] \\ &+ \frac{1}{2}(I^{-1})_{\mu\nu} \left[(S^T)_{\nu\rho} \left(\frac{-1}{8} \Pi_\alpha (F_{\rho\alpha\beta}(I^{-1})_{\beta\gamma} (S^T)_{\gamma\sigma} + S_{\alpha\beta}(I^{-1})_{\beta\gamma} F_{\rho\sigma\gamma}) \Pi_\sigma + 2\lambda q_\rho - \frac{\partial U}{\partial q_\rho} \right) \right]. \end{aligned} \quad (\text{B.16})$$

Using the fact that $(I^{-1})_{\beta\gamma}$ is symmetric, it can be shown that the second and third terms are equal. Hence,

$$\dot{\omega}_\mu = \frac{1}{2}(I^{-1})_{\mu\nu} \left[D_{\sigma\nu\rho} \dot{q}_\sigma \Pi_\rho - \frac{1}{4} (S^T)_{\nu\rho} \Pi_\alpha (F_{\rho\alpha\beta}(I^{-1})_{\beta\gamma} (S^T)_{\gamma\sigma}) \Pi_\sigma + 2\lambda (S^T)_{\nu\rho} q_\rho - (S^T)_{\nu\rho} \frac{\partial U}{\partial q_\rho} \right]. \quad (\text{B.17})$$

Consider the first term in the bracket. Inverting the definitions in Equation B.11:

$$D_{\sigma\nu\rho} \dot{q}_\sigma \Pi_\rho = D_{\sigma\nu\rho} \frac{1}{2} S_{\sigma\alpha} \omega_\alpha \cdot 2S_{\rho\delta} I_{\delta\beta} \omega_\beta = D_{\sigma\nu\rho} S_{\sigma\alpha} S_{\rho\delta} I_{\delta\beta} \omega_\alpha \omega_\beta \quad (\text{B.18})$$

Similarly the second term in the bracket becomes:

$$\begin{aligned}
\frac{1}{4}(S^T)_{\nu\rho}\Pi_\alpha F_{\rho\alpha\beta}(I^{-1})_{\beta\gamma}(S^T)_{\gamma\sigma}\Pi_\sigma &= \frac{1}{4}(S^T)_{\nu\rho}(2S_{\alpha\phi}I_{\phi\theta}\omega_\theta)F_{\rho\alpha\beta}(I^{-1})_{\beta\gamma}(S^T)_{\gamma\sigma}(2S_{\sigma\chi}I_{\chi\psi}\omega_\psi) \\
&= (S^T)_{\nu\rho}S_{\alpha\phi}I_{\phi\theta}F_{\rho\alpha\beta}(I^{-1})_{\beta\gamma}\delta_{\gamma\chi}I_{\chi\psi}\omega_\theta\omega_\psi \\
&= (S^T)_{\nu\rho}S_{\alpha\phi}I_{\phi\theta}F_{\rho\alpha\beta}(I^{-1})_{\beta\gamma}I_{\gamma\psi}\omega_\theta\omega_\psi \\
&= (S^T)_{\nu\rho}S_{\alpha\phi}I_{\phi\theta}F_{\rho\alpha\beta}\delta_{\beta\psi}\omega_\theta\omega_\psi \\
&= (S^T)_{\nu\rho}S_{\alpha\phi}I_{\phi\theta}F_{\rho\alpha\beta}\omega_\theta\omega_\beta \\
&= (S^T)_{\nu\rho}S_{\gamma\sigma}I_{\sigma\alpha}F_{\rho\gamma\beta}\omega_\alpha\omega_\beta
\end{aligned} \tag{B.19}$$

So,

$$\dot{\omega}_\mu = \frac{1}{2}(I^{-1})_{\mu\nu} \left([D_{\sigma\nu\rho}S_{\sigma\alpha}S_{\rho\delta}I_{\delta\beta} - S_{\rho\nu}S_{\gamma\sigma}I_{\sigma\alpha}F_{\rho\gamma\beta}] \omega_\alpha\omega_\beta + 2\lambda(S^T)_{\nu\rho}q_\rho - (S^T)_{\nu\rho}\frac{\partial U}{\partial q_\rho} \right). \tag{B.20}$$

With

$$\begin{aligned}
D_{0\nu\rho} &= \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}, \quad D_{1\nu\rho} = \begin{pmatrix} 0 & 1 & 0 & 0 \\ -1 & 0 & 0 & 0 \\ 0 & 0 & 0 & 1 \\ 0 & 0 & -1 & 0 \end{pmatrix}, \\
D_{2\nu\rho} &= \begin{pmatrix} 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & -1 \\ -1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \end{pmatrix}, \quad D_{3\nu\rho} = \begin{pmatrix} 0 & 0 & 0 & 1 \\ 0 & 0 & 1 & 0 \\ 0 & -1 & 0 & 0 \\ -1 & 0 & 0 & 0 \end{pmatrix},
\end{aligned} \tag{B.21}$$

and

$$\begin{aligned}
F_{0\nu\rho} &= \begin{pmatrix} 1 & 0 & 0 & 0 \\ 0 & 1 & 0 & 0 \\ 0 & 0 & 1 & 0 \\ 0 & 0 & 0 & 1 \end{pmatrix}, \quad F_{1\nu\rho} = \begin{pmatrix} 0 & -1 & 0 & 0 \\ 1 & 0 & 0 & 0 \\ 0 & 0 & 0 & -1 \\ 0 & 0 & 1 & 0 \end{pmatrix} \\
F_{2\nu\rho} &= \begin{pmatrix} 0 & 0 & -1 & 0 \\ 0 & 0 & 0 & 1 \\ 1 & 0 & 0 & 0 \\ 0 & -1 & 0 & 0 \end{pmatrix}, \quad F_{3\nu\rho} = \begin{pmatrix} 0 & 0 & 0 & -1 \\ 0 & 0 & -1 & 0 \\ 0 & 1 & 0 & 0 \\ 1 & 0 & 0 & 0 \end{pmatrix}.
\end{aligned} \tag{B.22}$$

The summations can be done explicitly, giving:

$$\begin{aligned}
\dot{\omega}_0 &= \lambda/I_0 + \tau_0/I_0, \\
\dot{\omega}_1 &= \frac{I_y - I_z}{I_x}\omega_2\omega_3 + \tau_1/I_x, \\
\dot{\omega}_2 &= \frac{I_z - I_x}{I_y}\omega_3\omega_1 + \tau_2/I_y, \\
\dot{\omega}_3 &= \frac{I_x - I_y}{I_z}\omega_1\omega_2 + \tau_3/I_y,
\end{aligned} \tag{B.23}$$

with the torque τ given by $\tau_\mu = -\frac{1}{2}(S^T)_{\mu\rho}\frac{\partial U}{\partial q_\rho}$. Imposing $q_\mu q_\mu = 1$ implies that $\omega_0 = 2q_\mu \dot{q}_\mu = 0$ and hence that $\dot{\omega}_0 = 0$. The Lagrange multiplier can therefore be identified as $\lambda = -\tau_0$. When this is substituted back into the expression for $\dot{\Pi}$, it cancels the effect of any component of $\frac{\partial U}{\partial \mathbf{q}}$ along \mathbf{q} , which is exactly what is needed to retain $q_\mu q_\mu = 1$. So, returning to the explicit use of particle labels and $S_{\mathbf{q}_i}^T$, the expression for $\dot{\Pi}_i$ becomes:

$$\dot{\Pi}_i = -\frac{1}{8}\frac{\partial}{\partial \mathbf{q}_i} \left((S_{\mathbf{q}_i}^T \Pi_i)^T (I_i)^{-1} (S_{\mathbf{q}_i}^T \Pi_i) \right) - \left(\frac{\partial U}{\partial \mathbf{q}_i} - \left(\mathbf{q}_i^T \frac{\partial U}{\partial \mathbf{q}_i} \right) \mathbf{q}_i \right). \quad (\text{B.24})$$

Note that once the condition $|\mathbf{q}|^2 = 1$ has been applied, $(I_i)_0$ cancels out from Equations B.9 and B.24. In fact, in the work of Miller *et al.* and Davidchack *et al.*, the kinetic term is written (equivalently) as:

$$\frac{1}{8} (S_{\mathbf{q}_i}^T \Pi_i)^T (I_i)^{-1} (S_{\mathbf{q}_i}^T \Pi_i) = \sum_{k=1}^3 h_k(\mathbf{q}_i, \Pi_i), \quad (\text{B.25})$$

with

$$h_k(\mathbf{q}_i) = \frac{1}{8I_{ik}} [\Pi_i^T \mathbf{P}_k(\mathbf{q}_i)]^2, \quad (\text{B.26})$$

in which

$$\mathbf{P}_1(\mathbf{q}_i) = (-(q_i)_1, (q_i)_0, (q_i)_3, -(q_i)_2)^T, \quad \mathbf{P}_2(\mathbf{q}_i) = (-(q_i)_2, -(q_i)_3, (q_i)_0, (q_i)_1)^T \quad (\text{B.27})$$

$$\text{and } \mathbf{P}_3(\mathbf{q}_i) = (-(q_i)_3, (q_i)_2, -(q_i)_1, (q_i)_0)^T$$

Quaternions and generalized quaternion momenta then evolve as:

$$\dot{\mathbf{q}}_i = \sum_{k=1}^3 \frac{\partial}{\partial \Pi_i} h_k(\mathbf{q}_i, \Pi_i) \quad \text{and} \quad \dot{\Pi}_i = -\sum_{k=1}^3 \frac{\partial}{\partial \mathbf{q}_i} h_k(\mathbf{q}_i, \Pi_i) - \left(\frac{\partial U}{\partial \mathbf{q}_i} - \left(\mathbf{q}_i^T \frac{\partial U}{\partial \mathbf{q}_i} \right) \mathbf{q}_i \right). \quad (\text{B.28})$$

Interestingly, if the $-\left(\mathbf{q}_i^T \frac{\partial U}{\partial \mathbf{q}_i} \right) \mathbf{q}_i$ term is left out of this expression, there are no consequences for the dynamics of physical observables. By dropping this term, the component of Π parallel to \mathbf{q} is not fixed at zero, as it technically should be. $\frac{\partial}{\partial \Pi, \mathbf{q}_i} h_k(\mathbf{q}_i, \Pi_i)$ is easily shown to be independent of any term of Π parallel to \mathbf{q} , however – therefore such a term has no effect on the motion or energy of the system. $\dot{\mathbf{q}}_i$ also remains orthogonal to \mathbf{q}_i , as it must.

B.3 Incorporating noise and damping

Equations B.8 and B.28 can be integrated to give Newtonian dynamics of rigid bodies, as outlined in Reference [218]. In this work, I am interested in motion driven by noise and damping, implicitly representing the presence of a solvent. Davidchack *et al.* [153] have developed an algorithm for this purpose [153] – here I discuss some aspects in more detail. For convenience, the $-\left(\mathbf{q}_i^T \frac{\partial U}{\partial \mathbf{q}_i}\right) \mathbf{q}_i$ term is not included in the equation of motion by the authors of Reference. From now on, the moment of inertia tensor I is taken to be three-dimensional again.

One can attempt to add noise and damping in the following simple fashion:

$$\begin{aligned} \dot{\mathbf{r}}_i &= \frac{\mathbf{p}_i}{m_i}, \\ \dot{\mathbf{p}}_i &= -\frac{\partial U}{\partial \mathbf{r}_i} - \gamma \mathbf{p}_i + b \mathbf{w}_i(t), \end{aligned} \quad (\text{B.29})$$

$$\begin{aligned} \dot{\mathbf{q}}_i &= \sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \quad \text{and} \\ \dot{\boldsymbol{\Pi}}_i &= -\sum_{k=1}^3 \frac{\partial}{\partial \mathbf{q}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) - \frac{\partial U}{\partial \mathbf{q}_i} - \Gamma G(\mathbf{q}_i, \boldsymbol{\Pi}_i) + B \mathbf{W}_i(t), \end{aligned} \quad (\text{B.30})$$

where $\mathbf{w}_i(t)$ and $\mathbf{W}_i(t)$ are 3- and 4-dimensional Wiener processes with unit variance and independent dimensions.¹

To specify the damping and noise terms that will result in sampling of the canonical ensemble, it is helpful to find the Fokker-Planck description equivalent to Equations B.29 and B.30. For a Langevin process with additive and independent noise, $\dot{\mathbf{v}} = \mathbf{A}(\mathbf{v}) + C \mathbf{W}(t)$ (where C is a constant diagonal matrix), the equivalent Fokker-Planck equation for a probability density $\rho(\mathbf{v}, t)$ is given by [152]:

$$\frac{\partial \rho(\mathbf{v}, t)}{\partial t} = -\frac{\partial}{\partial \mathbf{v}} \cdot (\mathbf{A}(\mathbf{v}) \rho(\mathbf{v}, t)) + \frac{1}{2} \sum_i C_{ii}^2 \frac{\partial^2 \rho(\mathbf{v}, t)}{\partial v_i^2}. \quad (\text{B.31})$$

For sampling of the canonical ensemble it is necessary that $\partial \rho_0 / \partial t = 0$, where

$$\rho_0 \propto \exp \left(-\beta \left(\sum_i \left(\sum_{k=1}^3 h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) + \mathbf{p}_i^2 / 2m \right) \right) + U(\{\mathbf{r}\}, \{\mathbf{q}\}) \right) \quad (\text{B.32})$$

is the Boltzmann distribution. Substituting Equation B.32 into Equation B.31, it is immediately obvious that cancellation of terms will occur separately for each particle, and also

¹Having different strengths of noise and damping for each particle would be a simple addition.

separately for linear and angular components. For the linear terms,

$$-\frac{\mathbf{p}_i}{m_i} \cdot \frac{\partial \rho_0}{\partial \mathbf{r}_i} + \left(\frac{\partial U}{\partial \mathbf{r}_i} + \gamma \mathbf{p}_i \right) \cdot \frac{\partial \rho_0}{\partial \mathbf{p}_i} + 3\gamma \rho_0 + \frac{b^2}{2} \frac{\partial^2 \rho_0}{\partial \mathbf{p}_i^2} = 0, \quad (\text{B.33})$$

$$\implies \frac{\beta \mathbf{p}_i}{m_i} \cdot \frac{\partial U}{\partial \mathbf{r}_i} \rho_0 - \left(\frac{\partial U}{\partial \mathbf{r}_i} + \gamma \mathbf{p}_i \right) \cdot \frac{\beta \mathbf{p}_i}{m_i} \rho_0 + 3\gamma \rho_0 + \frac{b^2}{2} \left(\frac{-3\beta}{m} + \frac{\beta^2 \mathbf{p}_i^2}{m^2} \right) \rho_0 = 0. \quad (\text{B.34})$$

It is fairly trivial to infer that the appropriate relationship between b and γ is:

$$\gamma = b^2 \frac{\beta}{2m}. \quad (\text{B.35})$$

For the angular terms,

$$\begin{aligned} & - \left(\sum_{k=1}^3 \frac{\partial}{\partial \mathbf{q}_i} \cdot \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \rho_0 - \left(\sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \cdot \frac{\partial \rho_0}{\partial \mathbf{q}_i} \\ & + \left(\sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} \cdot \frac{\partial}{\partial \mathbf{q}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \rho_0 + \Gamma \left(\frac{\partial}{\partial \boldsymbol{\Pi}_i} \cdot G(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \rho_0 \\ & + \left(\sum_{k=1}^3 \frac{\partial}{\partial \mathbf{q}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) + \frac{\partial U}{\partial \mathbf{q}_i} + \Gamma G(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \cdot \frac{\partial \rho_0}{\partial \boldsymbol{\Pi}_i} + \frac{B^2}{2} \frac{\partial^2 \rho_0}{\partial \boldsymbol{\Pi}_i^2} = 0. \end{aligned} \quad (\text{B.36})$$

It is fairly simple to show that:

$$\frac{\partial \rho_0}{\partial \mathbf{q}_i} = \left(-\beta \sum_{k=1}^3 \frac{\partial}{\partial \mathbf{q}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) - \beta \frac{\partial U}{\partial \mathbf{q}_i} \right) \rho_0, \quad (\text{B.37})$$

$$\frac{\partial \rho_0}{\partial \boldsymbol{\Pi}_i} = -\beta \left(\sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \rho_0 \quad (\text{B.38})$$

and

$$\frac{\partial^2 \rho_0}{\partial \boldsymbol{\Pi}_i^2} = \left(\beta^2 \left(\sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right)^2 - \beta \sum_{k=1}^3 \frac{\partial^2}{\partial \boldsymbol{\Pi}_i^2} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \rho_0 \quad (\text{B.39})$$

Thus,

$$\begin{aligned} & \Gamma \left(\frac{\partial}{\partial \boldsymbol{\Pi}_i} \cdot G(\mathbf{q}_i, \boldsymbol{\Pi}_i) - \beta G(\mathbf{q}_i, \boldsymbol{\Pi}_i) \cdot \sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \\ & = \frac{B^2}{2} \left(\beta^2 \left(\sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right)^2 - \beta \sum_{k=1}^3 \frac{\partial^2}{\partial \boldsymbol{\Pi}_i^2} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \right) \rho_0. \end{aligned} \quad (\text{B.40})$$

Equation B.40 is evidently satisfied by:

$$\Gamma G(\mathbf{q}_i, \boldsymbol{\Pi}_i) = \frac{B^2}{2} \beta \sum_{k=1}^3 \frac{\partial}{\partial \boldsymbol{\Pi}_i} h_k(\mathbf{q}_i, \boldsymbol{\Pi}_i) \quad (\text{B.41})$$

One way to express this result is:

$$\Gamma = \frac{\beta B^2}{2M}, \quad M = \frac{4}{\text{Tr}((I_i)^{-1})} \quad \text{and} \quad G(\mathbf{q}_i, \boldsymbol{\Pi}_i) = J(\mathbf{q}_i) \boldsymbol{\Pi}_i, \quad (\text{B.42})$$

and:

$$J(\mathbf{q}_i) = \frac{M}{4} \left(\sum_{k=1}^3 \frac{1}{(I_i)_k} \mathbf{P}_k(\mathbf{q}_i) \mathbf{P}_k^T(\mathbf{q}_i) \right) = \frac{S(\mathbf{q}) \begin{pmatrix} 0 & 0 \\ 0 & I^{-1} \end{pmatrix} S^T(\mathbf{q})}{\text{Tr}(I^{-1})}, \quad (\text{B.43})$$

exactly as stated in Chapter 3. Davidchack *et al.* [153] have introduced a numerical method for integrating Equations B.29 and B.30, and I have used this method in this work.

B.3.1 Subtleties related to the use of quaternions

In the previous section, it was shown that the dynamics of Equations B.29 and B.30, with the damping and noise terms related by Equations B.35, B.42 and B.43, will have a correct equilibrium distribution. It will also result in motion that conserves $|\mathbf{q}_i|^2$, as the expression for $\dot{\mathbf{q}}_i$ is orthogonal to \mathbf{q}_i . A subtlety, however, is that the Boltzmann distribution shown to be stationary is one *over all values of \mathbf{q}_i and $\boldsymbol{\Pi}_i$* , whereas only $|\mathbf{q}_i|^2 = 1$ and $\mathbf{q}_i \cdot \boldsymbol{\Pi}_i = 0$ are physically relevant. However, the dynamics of Equations B.29 and B.30 do not connect states with $|\mathbf{q}_i|^2 = 1$ to other values. Consequentially, the Boltzmann distribution at $|\mathbf{q}_i|^2 = 1$ must be stationary *in its own right*. Furthermore, as the value of $\boldsymbol{\Pi}_i$ parallel to \mathbf{q}_i has no influence on the energy or the dynamics of the simulated system, it cannot affect the distribution with respect to the other variables.

Appendix C

Validation of simulation techniques

It is important to ensure that the algorithms have been implemented correctly, and in the case of Langevin dynamics, that the time step of simulations is small enough to give accurate results.

C.1 Comparison of Langevin and VMMC energies

Langevin and VMMC methods can both be used to estimate the average energy of a system. As the methods are independent, a consensus is a good indicator that both algorithms are correctly sampling the system.

Simulations were performed on a system in which all the interactions present in the model occur frequently – a 30-base strand bound to two 10-base strands. The shorter strands bind to adjacent regions of the longer strand, and hence can undergo coaxial stacking across the joint.

Four VMMC simulations (of 5×10^9 steps per simulation), and four Langevin simulations each at a range of time steps h (for 5×10^8 steps per simulation) were performed at 300 K. The average potential energies measured in each simulation are show in Table C.1. As can be seen, the values agree well, with the average at each step size within 0.04% of the average obtained from VMMC simulations. No noticeable dependence of the maximum possible time step on the values of friction coefficients in the vicinity of $\gamma = 1$ and $\Gamma = 3$ was observed.

Simulation Method Timestep h	VMMC		Langevin		
	N/A	$h = 0.005$	$h = 0.003$	$h = 0.001$	$h = 0.0005$
$\langle U \rangle$ in each simulation	-59.0067	-59.0360	-58.9919	-58.9409	-58.9648
	-59.0118	-59.0146	-59.0042	-58.9646	-59.0070
	-59.0253	-59.0165	-59.0202	-59.1566	-59.0763
	-59.0290	-58.9882	-59.0313	-58.9267	-59.0295
Overall $\langle U \rangle$	-59.0182	-59.0138	-59.0119	-58.9972	-59.0194
Test statistic	N/A	0.7152	0.5681	0.7232	0.9630

Table C.1: Comparison of average potential energies obtained from VMMC and Langevin simulations. Timesteps and energies are quoted in the reduced units of the model. The test statistic is the result of applying Welch's unpaired t-test [214] to the distributions of $\langle U \rangle$ in the VMMC and Langevin code. For $h = 0.003$ (5.12 ps), the difference between VMMC and Langevin estimates of $\langle U \rangle$ is around 0.01% of the total, and is not statistically significant.

C.2 Comparison of hairpin folding speed as a function of step size in Langevin Dynamics

As discussed in Section C.1, Langevin dynamics with a step size of $h = 0.003$ in reduced units appears to provide thermodynamic averages consistent with the results of VMMC. Such a result, however, does not necessarily imply that the kinetics of simulations at these time steps are reliable.

As Langevin dynamics is used in this thesis to explore the kinetics of processes, it is important that the time step is small enough to reproduce them reliably. To test this, simulations were performed to measure the rate of spontaneous folding of a small hairpin as a function of step size. I considered a hairpin consisting of a stem of six base pairs and a loop of five bases, measuring the average time (from 100 simulations) required for a strand (initialized in the open state) to fold into a fully-formed hairpin at 300 K. The results are shown in Figure C.1.

As is evident from Figure C.1, there is no systematic dependence on hairpin folding rate on the step size h up to $h = 0.005$ simulation units. Therefore I use $h = 0.003$ with confidence throughout this work.

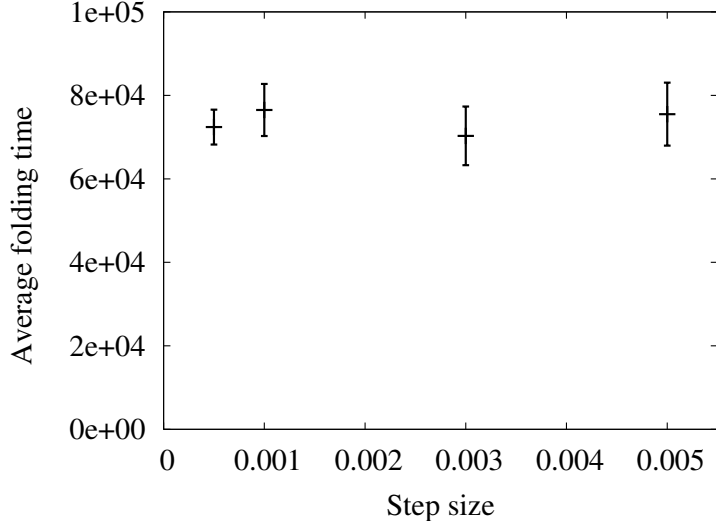


Figure C.1: Average simulation time for hairpin folding as a function of step size (in reduced units T , $T = 1$ corresponding to 1.706 ps). Error bars are calculated assuming that hairpin folding is a Poisson process.

C.3 Comparison of unbiased and biased VMMC simulations

Umbrella sampling is central to much of the work presented here. Due to its (often heavy) biasing of some configurations, it can be very sensitive to errors in algorithm implementation and typically responds differently from unbiased simulations. As a test of both the algorithm in general and the umbrella sampling code itself, therefore, simulations of the melting transition of a five-base pair duplex were performed both with and without biasing.

I performed four unbiased simulations at 297.03 K for 2×10^{11} VMMC steps each, and four biased simulations at 294.11 K for 4×10^{10} VMMC steps. The results (extrapolated to nearby temperatures using single-histogram re-weighting) are given in Table C.2. The results are clearly consistent, supporting the accuracy of my implementation of the VMMC algorithm and umbrella sampling.

Temperature	Unbiased		Biased	
	Average yield	σ	Average Yield	σ
309.28	5.31E-02	4.33E-04	5.27E-02	7.46E-04
306.12	8.87E-02	6.89E-04	8.81E-02	1.16E-03
303.03	1.41E-01	1.01E-03	1.40E-01	1.66E-03
300.00	2.11E-01	1.33E-03	2.10E-01	2.16E-03
297.03	2.96E-01	1.58E-03	2.95E-01	2.53E-03
294.12	3.89E-01	1.71E-03	3.87E-01	2.70E-03
291.26	4.82E-01	1.70E-03	4.81E-01	2.67E-03
288.46	5.70E-01	1.60E-03	5.69E-01	2.48E-03
285.71	6.48E-01	1.43E-03	6.47E-01	2.21E-03

Table C.2: Comparison of the bulk duplex yield inferred from four biased and four unbiased simulations of the formation of a five bp duplex (extrapolation to the bulk limit performed as outlined in Chapter 4). The average and standard deviation (σ) over the four simulations are shown.

Appendix D

Finite size effects for more complex systems

In Chapter 4, finite size effects relating to dimerization processes simulated in the canonical ensemble were discussed. In this appendix, the formalism is extended to multi-component assembly (of relevance to systems such as virus capsids and micelles, as well as more complex assemblies of DNA). Simulations using the grand canonical ensemble, and systems with one species immobilized, are also considered.

D.1 Monodisperse large homoclusters

Consider the formation of larger clusters of a specific size from identical monomers, a case relevant to the assembly of virus capsid-like objects [49, 50, 147, 219, 220, 221], and homomeric protein complexes [222]. If the formation of a single cluster is simulated in the canonical ensemble, once again, the statistics of the various cluster sizes do not directly correspond to bulk properties, but under the assumption that interactions which do not constitute bonding are negligible it is possible to extrapolate to large system sizes. Firstly, some definitions:

- n is the number of monomers needed to form the target, equal to the number of monomers simulated.
- z_i is the partition function for species i (a cluster of i identical monomers), in the simulation volume v , with the internal degrees of freedom treated *distinguishably*.

- $Z_{i,j,k\dots}$ is the partition function of a system of volume v when in a state which contains one cluster of species i , one of species j etc. This partition function is calculated using *distinguishable statistics*. $i, j, k\dots$ can therefore be taken to specify a macrostate corresponding to the set of clusters $i, j, k\dots$
- η_m the number of clusters of size m in the macrostate – consequentially, an alternative way to specify the macrostate is via the set $\{\eta_m\}$ ($Z_{i,j,k\dots} = Z_{\{\eta_m\}}$).
- $Z(n)$ is the total partition function of the n -particle system in a volume v , calculated using *distinguishable statistics*.
- as all monomers are identical, the A index will be omitted for clarity.

The thermodynamically relevant quantities are the $q_i/(q_1)^i$, because given these it is a simple task to calculate the bulk concentrations of each species using:

$$\frac{N_i}{(N_1)^i} = \frac{q_i}{(q_1)^i} \quad (\text{D.1})$$

and

$$\sum_i i N_i = n D = N. \quad (\text{D.2})$$

The quantities that are directly accessible from simulation are $Z_{i,j,k\dots}/Z$. Exactly how these can be accessed depends on how the system is sampled. A sensible choice, however, is to sample states by the largest cluster size – this neatly divides the partition function Z into n parts, and I label these subdivisions Ω_i . There are now n equations, one for each Ω_i :

$$\frac{\Omega_i}{Z} = \sum_{j,k\dots} \frac{Z_{i,j,k\dots}}{Z}, \quad (\text{D.3})$$

where the summation over $j, k\dots$ is the sum over all sets of indices such that $j, k\dots \leq i$ and the indices sum to n . $Z_{i,j,k\dots}$ are given by:

$$Z_{i,j,k\dots} = n! \prod_m^i \frac{(z_m/m!)^{\eta_m}}{\eta_m!}, \quad (\text{D.4})$$

I now have n simultaneous equations for $z_i/Z^{i/n}$ in terms of the measured quantities Ω_i/Z . In addition, these simultaneous equations have already been decoupled as each Ω_i/Z

expression contains only z_m with $m \leq i$, and thus finding $z_i/Z^{i/n}$ amounts to solving a polynomial of order i . All that remains is to find q_i in terms of z_i . This is reasonably simple:

$$q_i = D \frac{z_i}{i!}, \quad (\text{D.5})$$

where in dividing by $i!$ I account for the reduction in states imposed by indistinguishability.

I can then obtain the right hand side of (D.1) by:

$$\frac{q_i}{(q_1)^i} = \frac{D z_i}{i! (D z_1)^i} = \frac{D z_i / Z^{i/n}}{i! (D z_1 / Z^{1/n})^i}, \quad (\text{D.6})$$

in which the right hand side is expressed in terms of the known quantities $z_i/Z^{i/n}$. I can now eliminate the arbitrary large factor D by converting to concentrations (which equates to multiplying both sides by $(Dv)^{(i-1)}$), giving:

$$\frac{[N_i]}{[N_1]^i} = v^{i-1} \frac{z_i / Z^{i/n}}{i! (z_1 / Z^{1/n})^i}. \quad (\text{D.7})$$

Once again, the system of equations can be closed by conserving the total monomer number:

$$\sum_i i [N_i] = n/v. \quad (\text{D.8})$$

To illustrate the form of finite size corrections, consider the artificial example of completely cooperative hexamer formation (in which we approximate clusters of intermediate size as having zero probability). The complicating effects of additional states will be discussed in Section D.2. For comparison with Section 4.1.3, I will assume that the small system can be described by an equivalent two-state model, so that the yields of hexamers and homodimers are identical in the small simulation volume:

$$Z_6 / Z_{1,1,1,1,1,1} = \exp(-\Delta E/T + \Delta S), \quad (\text{D.9})$$

with $\Delta E = 2$ and $\Delta S = 15$ in reduced units. The result, plotted in Figure D.1 (c), indicates once again a much broader transition in the bulk case, this time with a slightly adjusted midpoint. Furthermore, this broadening is much more pronounced for hexamers than dimers. This trend is a general one, with larger clusters experiencing greater broadening due to finite-size corrections. This is because smaller relative concentration fluctuations are required to push the system towards a yield of approximately 50% for a clustering transition involving many monomers as opposed to dimers, as illustrated in Figure D.1 (b).

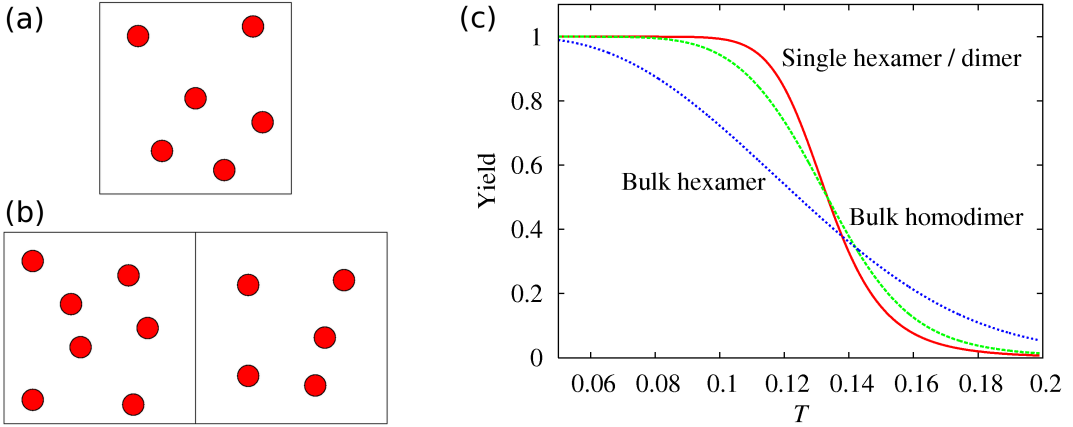


Figure D.1: (a) The top image shows six identical particles in a volume v . Doubling the volume means that only a relatively minor concentration fluctuation is required to make the formation of two hexamers impossible (and the formation of one hexamer more likely), compared to the equivalent situation for dimers. As a consequence, the broadening effect of bulk corrections increases with the size of the target structure. (b) Fractional yield of hexamers in the statistical model of equation (D.9). Plotted are the yields for a single cluster system (red), for hexamers in bulk (blue) and for an equivalent extrapolation to bulk for homodimer formation (green).

D.2 Homocluster convergence

Many canonical simulations of self-assembly are performed using systems large enough to form several or many clusters [49, 50, 219, 220, 223]. I apply the formalism of the previous sections to explore the convergence of cluster statistics on bulk values as system size is increased. As in the previous section, I consider a reference system of n particles in a volume v , where n is the size of the largest cluster, and proceed using the partition functions z_i defined in this volume.

The statistical weight of the macrostate $i, j, k\dots$, containing a total of Dn monomers in a volume D , with all degrees of freedom treated indistinguishably, is given by:

$$Q_{\{\eta_m\}}(D) = \prod_l^n \frac{(Dz_l)^{\eta_l}}{\eta_l!(l!)^{\eta_l}}, \quad (\text{D.10})$$

Defining $\psi_l(D) = z_l/(z_1^l D^{l-1})$, I obtain an expression for the fractional yield of a cluster of size c in a system of size D :

$$f_c(D) = \frac{c \sum_{\{\eta_m\}} \eta_c \prod_l^n \frac{1}{(\eta_l)!} \left(\frac{\psi_l}{l!}\right)^{\eta_l}}{Dn \sum_{\{\eta_m\}} \prod_l^n \frac{1}{(\eta_l)!} \left(\frac{\psi_l}{l!}\right)^{\eta_l}}. \quad (\text{D.11})$$

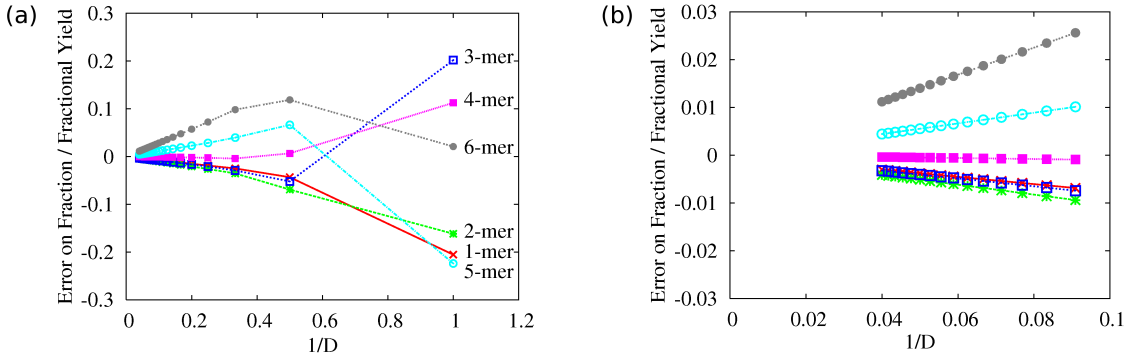


Figure D.2: (a) Relative fractional error on the yield of various cluster sizes as a function of $1/D$ in a system with a maximum cluster size of six. All clusters show convergence with $1/D$ scaling in the large D regime (as highlighted in (b)).

This expression for $f_c(D)$ can be used to check that the extrapolation scheme is valid for the system in question, by simulating a system with $D = 2$ and comparing to the results expected from a $D = 1$ simulation. Such a simulation would also allow the examination of clusters larger than the target structure, and their incorporation into the statistics if their presence is significant.

In all cases that I have been able to study to high D (the meaning of ‘high’ will be clarified later), $f_c(D) - f_c(\infty)$ is observed to scale as $1/D$ in the large D limit (see Figure D.2). The question of convergence speed then reduces to how large D must be for this scaling to hold, and the value of $f_c(D) - f_c(\infty)$ at this point. In general there are two distinct regimes of convergence, determined by the yield of target structures. I shall illustrate these regimes by considering completely cooperative transitions (in which only the target cluster and monomer concentrations are non-negligible), before commenting on the effects of other cluster sizes.

D.2.1 Convergence at low yield

Section D.1 indicates that simulations of a single cluster underestimate the transition width and hence underestimate the number of clusters at low yield. In effect, in order to have a high isolated monomer fraction in bulk despite the effects of volume fluctuations, the fraction of monomers in a single target simulation must be even higher. As the system size

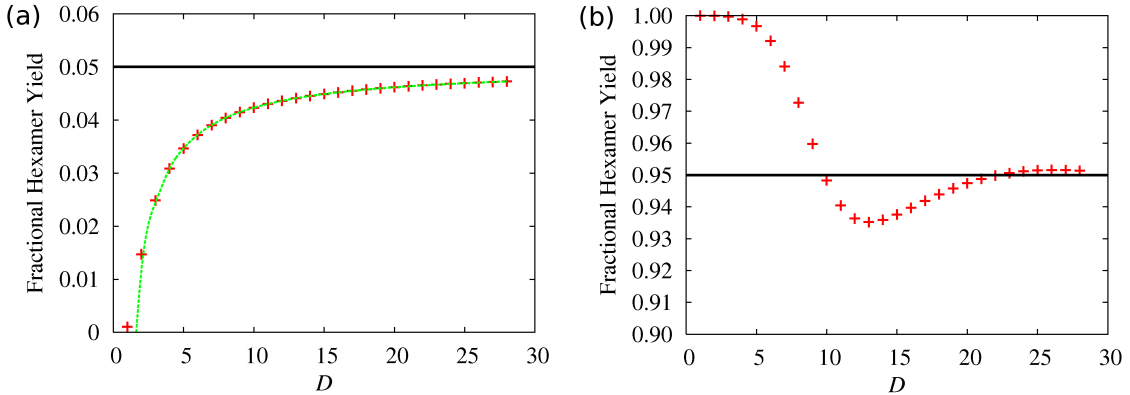


Figure D.3: Fractional yield of hexamers in a perfectly cooperative model as a function of system size D for (a) low yield (5% hexamers) and (b) high yield (95% hexamers). The ‘+’ symbols are the calculated points and the flat line the bulk value. The curve in (a) is a fit to a $1/D$ convergence.

is increased, concentration fluctuations tend to transfer statistical weight from the extreme state favoured at $D = 1$ towards a more balanced cluster size distribution.

At low yield this effect produces a steady increase in the proportion of clusters with D , with the deviation from the bulk fraction scaling as approximately $1/D$ from low D (see Figure D.3(a)). At very low yield, initial convergence becomes noticeably slower than $1/D$ —this effect increases with target size. As a consequence, relative errors remain significant at increasingly large values of D as the yield is decreased or the target size increased.

D.2.2 Convergence at high yield

At high yield, single target simulations overestimate the monomer fraction, for reasons similar to the underestimate at low yield. Convergence, however, does not initially show a $1/D$ behaviour, as illustrated in Figure D.3(b). Instead, a period of slow convergence is followed by a rapid drop to a target yield just below the bulk value, leading eventually to an oscillation in the vicinity of the bulk yield. These oscillations persist for approximately $n - 1$ half-cycles, before settling in to a $1/D$ convergence (n being the target cluster size).

These oscillations result from certain configurations disproportionately biasing the ensemble, due to the inherently discrete nature of a small system. At $D = 1$, the system is restricted to two states: one cluster or n monomers. At high cluster yield, n monomers are extremely unfavourable and hence the single cluster state is overwhelmingly observed,

causing $f_n(1)$ to exceed $f_n(\infty)$. As the system size is increased, the zero monomer state continues to exert a disproportionate influence on the ensemble, keeping $f_n(D)$ well above $f_n(\infty)$. Eventually, however, the system becomes sufficiently large that the state with $D - 1$ clusters is most favourable. Due to the discreteness of the system, this occurs before $(D - 1)/D = f_n(\infty)$. As a consequence, $f_n(D)$ is then underestimated (or equivalently the number of monomers is overestimated), resulting in the observed drop of $f_n(D)$. At still larger values of D , the state with $D - 1$ clusters remains most favourable but now constitutes an overestimate of $f_n(D)$, resulting in the observed rise in $f_n(D)$. This process is repeated for increasing number of monomers, leading to oscillations which are eventually overwhelmed by the $1/D$ convergence at large system size.

The question is then why oscillations are observed at high but not low yield, where the discreteness of the system is still present. To answer this, it is illuminating to allow D to take non-integer values so that the system size $n' = Dn$ can take any integer value. At high yield, as shown in Figure D.4(b), one sees that the system is extremely sensitive to the exact number of particles, because if D is not an integer there are necessarily excess monomers. This results in the rapid oscillation of $f_n(n')$ with a period of approximately n . Closer inspection, however, reveals that the period is longer than n , due to the fact that states with no monomers present become increasingly unfavourable as D gets larger. The region in which the $f_n(n')$ peaks transfer from $n' \bmod n = 0$ to $n' \bmod n = 1$ corresponds to the region in which $f_n(D)$ drops off rapidly. By contrast, $f_n(n')$ increases monotonically with n' at low yield (Figure D.4(a)). In this regime, the fraction of clusters is not high enough for the value of $n' \bmod n$ to be significant, and so the general tendency to transfer statistical weight to states with a greater mix of cluster sizes is dominant, and smooth convergence is observed.

As a consequence of this behaviour, convergence at high cluster yield is extremely poor until D is sufficiently large that the state with $D - 1$ clusters has approximately the same weight as the state with D clusters:

$$\frac{(\psi_n)^{D-1}\psi_1}{(D-1)!(n!)^{D-1}n!} \approx \frac{(\psi_n)^D}{(D)!(n!)^D}. \quad (\text{D.12})$$

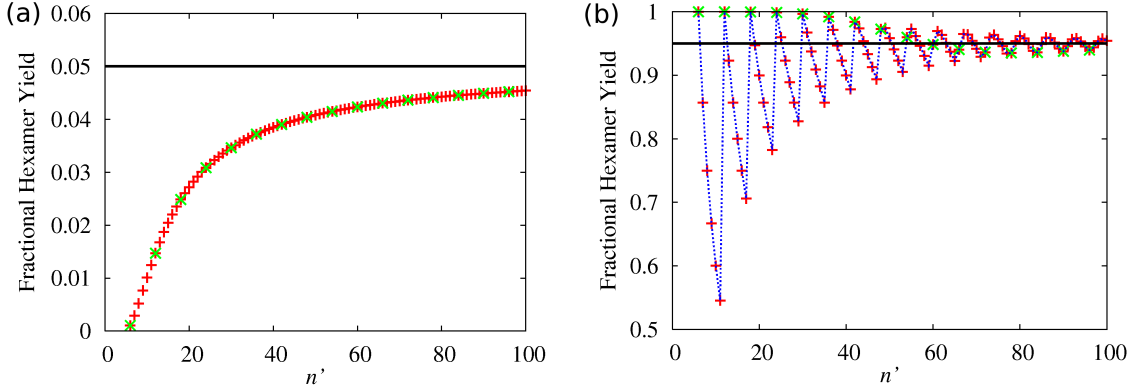


Figure D.4: Fractional yield of hexamers in a perfectly cooperative model as a function of n' for (a) low yield (5% hexamers) and (b) high yield (95% hexamers). The ‘+’ symbols are the calculated points and the flat line the bulk value. ‘x’ symbols indicate system sizes for which $D = n/n'$ is integral. The dashed line in (b) is added as a guide to the eye.

Substituting using the definition of $\psi_i(D)$ gives:

$$D_{\text{crossover}} \approx \left(\frac{z_n}{z_1^n} \right)^{1/n}. \quad (\text{D.13})$$

The quantity z_n/z_1^n corresponds to the ratio of cluster to monomers at $D = 1$, and consequently increases with n at fixed bulk yield. This increase is offset by the $1/n$ exponent, meaning that the value of $D_{\text{crossover}}$ is relatively independent of target size, but increases with the target yield. It should also be noted that the oscillations persist until a system size of approximately $nD_{\text{crossover}}$, although they are generally reasonably small. It is this value, $D \approx nD_{\text{crossover}}$, that defines the large D limit.

In the intermediate yield regime near to the midpoint of the transition, the initial error is small and z_n/z_1^n is not large, hence convergence is fast (whether it proceeds by the first or second method). Away from the midpoint, however, significant relative discrepancies can persist up to surprisingly large system sizes.

D.2.3 Intermediate cluster sizes

An additional complication for $n > 2$ is the fact that intermediate cluster sizes may be relevant to the system, which can affect convergence. I shall analyze the effects of the presence of intermediate cluster sizes under the assumption that the majority of particles are found either as isolated monomers or in the target cluster size: for the purposes of

this section, the term ‘majority species’ applies to the most prevalent of either the target cluster or monomers, and ‘minority species’ to the less common of these two. Note that our discussions will compare the effects of intermediate cluster sizes in systems with a certain yield of the majority species, as it is the tendency of one species to dominate in bulk despite concentration fluctuations that causes the large discrepancies at $D = 1$. Firstly, I consider the low yield case. Here, the presence of clusters of intermediate sizes with bulk yields comparable to the target cluster has little effect on the relative error of the target yield at $D = 1$, which is largely determined by the bulk fraction of monomers. By contrast, if the relevant intermediate cluster size is small (for instance a dimer in a system forming a dodecahedron), the relative error between dimer and isolated monomers is comparatively small, meaning that $f_2(1) \approx f_2(\infty)$, because from the perspective of the monomer/dimer equilibrium the system has an effective size of $D_{\text{eff}} = n/2$. As a consequence, states including dimers are common and so the entropic penalty associated with having no target clusters is reduced, meaning that statistical weight is transferred to larger clusters more slowly as the system size is increased. The effect manifests itself as a poor convergence in the first few steps, as shown in Figure D.5(a). Also shown is the effect of having a significant presence of large intermediate clusters, which is smaller as they do not relieve the entropic penalty of having many monomers as swiftly as dimers do (the relative error is seen to behave similarly to a completely cooperative system with the same monomer yield).

I now consider the effect of a significant presence of intermediate clusters on the convergence of the yield of *isolated monomers* at high cluster fraction. If the relevant intermediate clusters are large, the initial error is significantly reduced as the relative error between two large clusters of similar size is much smaller than for a large cluster and a monomer, and in forming intermediate clusters some monomers are ‘spare’. Convergence, however, is not improved as instead of the state with $D - 1$ target clusters and D monomers coming to dominate the ensemble, as in the completely cooperative case, states containing intermediate clusters become most prevalent (in effect, they reduce the ‘entropy cost’ associated with having few monomers in the system). If the intermediate clusters are large, there will be few monomers in these states and as a consequence, statistical weight is transferred to

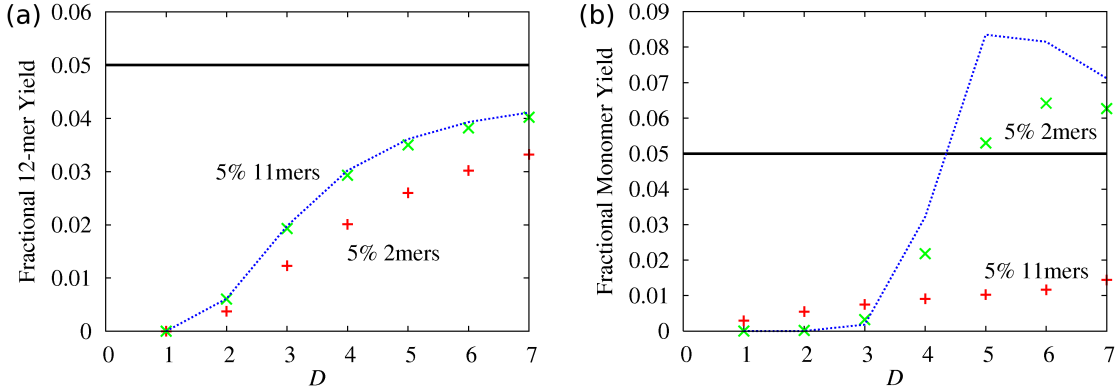


Figure D.5: (a) Fractional yield of dodecamers in a dodecamer forming system at low yield (90% isolated monomers, 5% dodecamers in bulk). (b) Fractional yield of isolated monomers in a dodecamer forming system at high yield (90% dodecamers, 5% monomers in bulk). Plotted are points for systems in which the other 5% is assumed to consist entirely of either 2-mers or 11-mers. Also shown (dashed curves) are the results for completely cooperative systems with the same 90% majority species yield. These have been scaled by a factor of 0.5 so that the relative errors can be directly compared.

isolated monomers more slowly. This effect is illustrated in Figure D.5 (b): also plotted is a case with a significant presence of small intermediate clusters. In this case convergence is not dramatically slowed (relative to a completely cooperative system with the same target cluster yield), as the states which become prevalent contain $D - 1$ target clusters and a mix of smaller species, including several monomers.

In summary, for monodisperse clusters, the significant presence of intermediate cluster sizes tends to reduce the rate of convergence of the fractional yield of the minority species relative to a completely cooperative system (at a fixed yield of the majority species), particularly if the relevant intermediate clusters are closer in size to the majority species, by reducing the entropic penalty associated with having few of the minority species in the system. In several cases, authors have studied systems capable of forming approximately 10 to 20 clusters [49, 220, 223]. It is probable that the finite-size effects illustrated here are relevant to these systems in the regimes dominated by one cluster size.

Small-system simulations on the formation of micelles have also been performed in the canonical ensemble (examples are given in Reference [224]). Micelles are not monodisperse, but show some variation about a typical finite size. The techniques shown in this section can be equally well applied to such a process, with qualitatively similar results – in particular,

convergence to bulk yields appears to be extremely slow when the micellar concentration is low. The details are discussed in Reference [224].

D.3 Simulations in the grand canonical ensemble

As an alternative to *NVT* simulations, it is possible to use the grand canonical ensemble. Here, instead of fixing the number of particles absolutely, one simulates a system in a volume v such that configurations containing n particles are sampled with the relative probability [225]:

$$P(n) \propto \frac{e^{\beta\mu n} Z(n)}{n!}, \quad (\text{D.14})$$

where $Z(n)/n!$ is the partition function of an n -particle system, calculated using distinguishable statistics. Configurations of the same n have the same relative Boltzmann weighting as in the canonical ensemble. Thus a configuration containing n_j clusters of size j is observed in a simulation with probability:

$$P(\{n_j\}) = \prod_{j>0} \frac{z_j^{n_j}}{n_j! j^{n_j}} e^{\beta\mu_j n_j} \quad (\text{D.15})$$

To date, grand canonical techniques (and related semi-grand canonical approaches) have primarily been used to study the formation of micellar structures [226, 227, 228, 229, 230, 231, 232], as opposed to monodisperse target structures. In many of these cases, monomer concentrations are taken to be so low that the probability of finding more than one cluster in a simulation volume is neglected, and the number of monomers in the simulation is taken as a proxy for cluster size [226, 227, 228, 229]. There has been considerable debate on the details of inferring cluster probabilities from simulations in which one monomer is fixed at the centre of the simulation volume, under this extremely dilute assumption [226, 233, 234]. Other workers have explicitly simulated the formation of multiple micellar clusters [230, 231, 232]. The advantage of the grand canonical formalism for large micelles is obvious, as it allows a small simulation volume to contain a large number of particles without requiring a high average concentration. There is no reason, however, why the technique could not also be applied to smaller, monodisperse self-assembling systems.

In principle, grand-canonical simulations are capable of capturing the concentration fluctuations which must be considered to extract the bulk thermodynamics from a small simulation. This is only true, however, if the simulation samples the formation of *multiple* clusters. Under the assumptions of ideal behaviour of separate species, the errors are simple to quantify. In bulk equilibrium:

$$[N_1] = \frac{z_1 e^{\beta\mu}}{v}, \quad (\text{D.16})$$

where concentrations are measured relative to the simulation volume v , and:

$$[N_j] = [N_1]^j (Dv)^{j-1} \frac{q_j}{q_1^j} = \frac{z_j}{vj!} e^{\beta\mu j}. \quad (\text{D.17})$$

I assume that a simulation samples states which contain at most a single cluster of more than one particle. In this case, the probability of observing a cluster of size $j > 1$ is given by:

$$p(j) = \frac{\frac{z_j}{j!} e^{\beta\mu j} \sum_{n \geq 0} \frac{z_1^n e^{\beta\mu n}}{n!}}{(1 + \sum_{k>1} \frac{z_k}{k!} e^{\beta\mu k}) \sum_{n \geq 0} \frac{z_1^n e^{\beta\mu n}}{n!}}, \quad (\text{D.18})$$

Thus the simulation concentration (measured relative to the simulation volume) is:

$$[N_j]^{\text{sim}} = \frac{\frac{z_j}{vj!} e^{\beta\mu j}}{(1 + \sum_{k=2}^{\infty} \frac{z_k}{k!} e^{\beta\mu k})} = \frac{[N_j]}{1 + \sum_{k>1} [N_k]v} \quad (\text{D.19})$$

Note that all $[N_j]$ are inversely proportional to the simulation volume v – hence relative errors initially grow with the simulation volume, before plateauing in the limit of $\sum_{k>1} [N_k] \gg 1$ (when the relative error is approximately unity).

$$\frac{[N_j] - [N_j]^{\text{sim}}}{[N_j]} = \frac{\sum_{k>1} [N_k]v}{1 + \sum_{k>1} [N_k]v}. \quad (\text{D.20})$$

Extracting $[N_j]$ from the measured $[N_j]^{\text{sim}}$ is simple. It can be shown that, under our assumptions of ideality, $[N_1] = [N_1]^{\text{sim}}$ for such a simulation.

Equation D.19 can be rearranged to give $j_{\max} - 1$ linear simultaneous equations for $[N_j]$, where j_{\max} is the largest cluster considered ($[N_1]$ is trivially given by $[N_1] = [N_1]^{\text{sim}}$). The validity of the extrapolation can be simply checked by simulating the system in a different volume and observing whether the $[N_j]^{\text{sim}}$ are predicted by the $[N_j]$ inferred from the original simulation.

In reality, it is unlikely that simulations are explicitly forbidden from sampling states with multiple large clusters. If, however, cluster formation involves a significant free energy barrier, and only the formation of a single cluster is actively biased by the simulation, multiple large clusters will not be observed. In these cases, Equation D.19 can be used, but any rare instances where multiple clusters do occur (for example, two dimers) must not be included in evaluating $[N_j]^{\text{sim}}$. In the extremely dilute limit, the results obtained from this method will coincide with the assumption that at most one cluster exists in a simulation volume. Given the simplicity of the extrapolation scheme presented here, however, this assumption seems unnecessary.

D.4 Monodisperse large heteroclusters

Interesting structures are not only formed from identical subunits. DNA nanostructures, such as the Turberfireld group's tetrahedra [24], often involve one each of several different single strands. Virus capsids can require more than one type of coat protein, and some work has been undertaken to simulate models of such structures [148]. Simulators have also considered templated assembly, in which distinct shells of particles form cooperatively [235].

To discuss finite size effects present in large heterocluster formation, it is necessary to extend the definitions of Section D.1.

- y is the number of distinct monomer species present.
- n_x is the number of monomers of species x required to form one target structure.

$$n = \sum_x n_x$$

- $z_{(i_1, i_2 \dots i_y)}$ is the partition function for species $(i_1, i_2 \dots i_y)$ (a cluster of i_1 monomers of type 1, i_2 monomers of type 2 *etc.*), in the simulation volume v , with the internal degrees of freedom treated *distinguishably*.
- let $\{i\}$ denote (i_1, i_2, i_3) to conserve space.

- $Z_{(i_1, i_2 \dots i_y), (j_1, j_2 \dots j_y) \dots} = Z_{\{i\}, \{j\} \dots}$ is the partition function of a system of volume v when in a state which contains one molecule of species $(i_1, i_2 \dots i_y)$, one of species $(j_1, j_2 \dots j_y)$ etc.. This partition function is calculated using *distinguishable statistics*.
- $Z(n)$ is the total partition function of the n -particle system in a volume v , calculated using *distinguishable statistics*.
- $q_{(i_1, i_2 \dots i_y)} = q_{\{i\}}$ is the partition function for species $(i_1, i_2 \dots i_y)$ in a volume Dv , with the internal degrees of freedom treated indistinguishably.
- $\eta_{\{m\}}$ is the number of times that the cluster $(m_1, m_2, \dots m_y)$ appears in a macrostate. The macrostate is therefore completely specified by the set $\{\eta_{\{m\}}\}$, or alternatively the list of clusters $\{i\}, \{j\}, \{k\} \dots$ – both alternatives will prove useful.

As in Section D.1, the important quantities to estimate are:

$$(Dv)^{i_0-1} \frac{q_{(i_1, i_2 \dots i_y)}}{q_{(1,0,\dots,0)}^{i_1} q_{(0,1,\dots,0)}^{i_2} \dots q_{(0,0,\dots,1)}^{i_y}}, \quad (\text{D.21})$$

where $i_0 = \sum_x i_x$. Given these ratios, the bulk concentrations are found by solving:

$$\sum_{\{i_1, i_2 \dots i_y\}} i_x [N_{(i_1, i_2 \dots i_y)}] = \frac{n_x}{v} \quad (\text{D.22})$$

for every x , using:

$$\frac{[N_{(i_1, i_2 \dots i_y)}]}{[N_{(1,0,\dots,0)}]^{i_1} [N_{(0,1,\dots,0)}]^{i_2} \dots [N_{(0,0,\dots,1)}]^{i_y}} = (Dv)^{i_0-1} \frac{q_{(i_1, i_2 \dots i_y)}}{q_{(1,0,\dots,0)}^{i_1} q_{(0,1,\dots,0)}^{i_2} \dots q_{(0,0,\dots,1)}^{i_y}}. \quad (\text{D.23})$$

One can relate the $q_{(i_1, i_2 \dots i_y)}$ to $z_{(i_1, i_2 \dots i_y)}$, which are closer to the quantities directly measurable in simulation, via:

$$q_{(i_1, i_2 \dots i_y)} = D \frac{z_{(i_1, i_2 \dots i_y)}}{\prod_x^y i_x!}. \quad (\text{D.24})$$

Clearly, this is a more involved problem than the homocluster case. Having obtained $z_{(i_1, i_2 \dots i_y)}$, one must then solve y nonlinear simultaneous equations, rather than just a single polynomial equation. A more subtle problem, however, is how to extract the relative values of $z_{(i_1, i_2 \dots i_y)}$ from the simulation data. For the homocluster case, the method presented in Section D.1 is a natural one. The size of the largest cluster is a good order parameter, and

it is likely that any sampling scheme that forces the system between monomers and the target will sample all of the intervening states accurately.

By contrast, the order parameters for heterocluster formation are naturally multi-dimensional (simply sampling according to the largest cluster size is insufficient to isolate $z_{\{i\}}$). The question of how to best disentangle the information is then somewhat system dependent. One simple approach would be to obtain an estimate for each individual $Z_{\{i\},\{j\},\{k\}\dots}/Z(n)$ from simulation, and then fit $z_{\{i\}}/Z(n)^{i_0/n}$ to reproduce these estimates, using:

$$Z_{\{\eta_{\{m\}}\}}/Z(n) = \prod_x^y n_x! \prod_{\{l\}} \left(\frac{1}{\eta_{\{l\}}!} \right) \left(\frac{z_{\{l\}}/Z(n)^{l_0/n}}{\prod_x^y l_x!} \right)^{\eta_{\{l\}}}, \quad (\text{D.25})$$

where $i_0 = \sum_x i_x$, and the product over $\{l\}$ runs over all possible clusters. Note that the presence of $Z(n)$, which arises because only relative measurements are possible, is not important as it will always cancel when values are substituted into quotients such as on the RHS of Equation D.23.

Alternatively, one could construct a set of simultaneous equations to obtain a subset of $z_{\{i\}}/Z(n)^{i_0/n}$, and then use this subset to obtain all $z_{\{i\}}/Z(n)^{i_0/n}$. This approach is similar to the one used in Section D.1, but is complicated by the fact that there is no obvious order parameter which immediately decouples all $z_{\{i\}}/Z(n)^{i_0/n}$. This approach is best illustrated with a basic example – consider the assembly of two species A and B into a structure which contains two of each type of particle.

With a system this simple, it is trivial to enumerate all possible macrostates of a small simulation and quantify their probability of occurring.

1. No clusters of more than one particle:

$$Z_{(1,0),(1,0),(0,1),(0,1)}/Z(4) = (2!)^2 \left(\frac{z_{(1,0)}/Z(4)^{1/4}}{1!} \right)^2 \left(\frac{z_{(0,1)}/Z(4)^{1/4}}{1!} \right)^2 \left(\frac{1}{2!} \right)^2.$$

2. One cluster of two A particles:

$$Z_{(2,0),(0,1),(0,1)}/Z(4) = (2!)^2 \left(\frac{z_{(2,0)}/Z(4)^{1/2}}{2!} \right) \left(\frac{z_{(0,1)}/Z(4)^{1/4}}{1!} \right)^2 \left(\frac{1}{2!} \right).$$

3. One cluster of two A particles, one cluster of two B particles:

$$Z_{(2,0),(0,2)}/Z(4) = (2!)^2 \left(\frac{z_{(2,0)}/Z(4)^{1/2}}{2!} \right) \left(\frac{z_{(0,2)}/Z(4)^{1/2}}{2!} \right).$$

4. One cluster of two B particles:

$$Z_{(1,0),(1,0),(0,2)}/Z(4) = (2!)^2 \left(\frac{z_{(1,0)}/Z(4)^{1/4}}{1!} \right)^2 \left(\frac{z_{(0,2)}/Z(4)^{1/2}}{2!} \right) \left(\frac{1}{2!} \right).$$

5. One cluster of one A and one B particle:

$$Z_{(1,1),(1,0),(0,1)}/Z(4) = (2!)^2 \left(\frac{z_{(1,1)}/Z(4)^{2/4}}{1!1!} \right) \left(\frac{z_{(1,0)}/Z(4)^{1/4}}{1!} \right) \left(\frac{z_{(0,1)}/Z(4)^{1/4}}{1!} \right).$$

6. Two cluster of one A and one B particle:

$$Z_{(1,1),(1,1)}/Z(4) = (2!)^2 \left(\frac{z_{(1,1)}/Z(4)^{2/4}}{1!1!} \right)^2 \left(\frac{1}{2!} \right).$$

7. One cluster of three particles (two A and one B):

$$Z_{(2,1),(0,1)}/Z(4) = (2!)^2 \left(\frac{z_{(2,1)}/Z(4)^{3/4}}{2!1!} \right) \left(\frac{z_{(0,1)}/Z(4)^{1/4}}{1!} \right)^2.$$

8. One cluster of three particles (two B and one A):

$$Z_{(1,2),(1,0)}/Z(4) = (2!)^2 \left(\frac{z_{(1,2)}/Z(4)^{3/4}}{2!1!} \right) \left(\frac{z_{(0,1)}/Z(4)^{1/4}}{1!} \right)^2.$$

9. One cluster of four particles (two B and two A):

$$Z_{(2,2)}/Z(4) = (2!)^2 \left(\frac{z_{(2,2)}/Z(4)}{2!2!} \right).$$

It is possible to take the simulation results for macrostates 1, 5 and 6 above and use them to solve for $z_{(0,1)}/Z(4)^{1/4}$, $z_{(1,0)}/Z(4)^{1/4}$ and $z_{(1,1)}/Z(4)^{1/2}$. Given these values, one can simply find the remaining $z_{(i,j)}$ by substituting them into expressions for macrostates 2, 4, 7 and 8 and 9.

Although the method is fairly simple in this case, it is not obvious that a systematic process (that will work well for all situations) exists. For example, in this case macrostate 3 (which consists of two homoclusters) is neglected in the calculation. This is wasteful of information, particularly if macrostate 3 is frequently sampled in simulations. It is possible to combine macrostates 3 and 4, and use the aggregated data to find $z_{(i,j)}$, but there is no reason why this particular aggregation should be made rather than, for instance, 2 and 4. As systems get larger and more complex, further arbitrary decisions will have to be made.

D.5 Heterocluster convergence

If a simulation of the formation of a single heterocluster has been performed, and the quantities $z_{(i_1, i_2 \dots i_y)}$ extracted, one can use the formalism of the previous section to extrapolate to bulk, under the assumption that distinct species behave ideally. As with homoclusters and dimers, it is useful to be able to predict not only the bulk yield, but how fractional yields will change as the system is increased in size, so that the assumption of ideality can be checked.

Proceeding by analogy with Section D.2, consider a system of size D containing Dn particles. Defining the fraction of particles of type a in cluster $(c_1, c_2, \dots c_y)$ as $f_{(c_1, c_2, \dots c_y)}^a(D)$:

$$f_{(c_1, c_2, \dots c_y)}^a(D) = \frac{\sum_{\{\eta_{\{m\}}\}} c_a \eta_{(c_1, c_2, \dots c_y)} \prod_{\{l\}} \frac{(\psi_{\{l\}} / (\Pi_x^y l_x!))^{\eta_{\{l\}}}}{(\eta_{\{l\}})!}}{Dn_a \sum_{\{\eta_{\{m\}}\}} \prod_{\{l\}} \frac{(\psi_{\{l\}} / (\Pi_x^y l_x!))^{\eta_{\{l\}}}}{(\eta_{\{l\}})!}}, \quad (\text{D.26})$$

where:

$$\psi_{\{l\}}(D) = \frac{z(l_1, l_2, \dots l_y)}{D^{l_0-1} z_{(1, 0, \dots 0)}^{l_1} z_{(0, 1, \dots 0)}^{l_2} \dots z_{(0, 0, \dots 1)}^{l_y}}. \quad (\text{D.27})$$

To illustrate this convergence, the behaviour of a tetramer formed from two particles each of two different types is shown in Figure D.6.

D.6 Immobilized species

In some experimental systems, the particles which associate are not all free to diffuse. For example, DNA microarray assays consist of DNA ‘probes’ which are tethered to a surface and ‘target’ molecules which diffuse through solution [139, 210]. Another possibility would be DNA walkers, which are generally stuck to a track and may in some applications be effectively localized.

It is not *a priori* obvious whether tethering one reactant will change the result of the previous sections, in which local concentration fluctuations were invoked to explain the difference between bulk and small-system statistics. Note that here I am not concerned with whether the mechanism of tethering interacts with the particles, either destabilizing

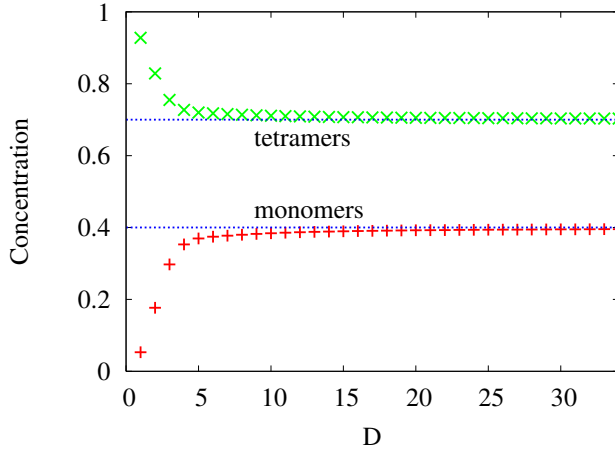


Figure D.6: Convergence of the concentration (measured in clusters per unit simulation volume) in a tetramer forming system, as the system size D is increased. Yields are calculated for a system with bulk concentrations of $[N_{10}] = [N_{01}] = 0.4$, $[N_{20}] = [N_{02}] = 0.005$, $N_{11} = 0.04$, $[N_{21}] = [N_{12}] = 0.05$ and $[N_{22}] = 0.7$.

or stabilizing the bound state, but rather whether extrapolation to bulk for a given set of $z_{\{i\}}$ differs from Section D.4.

To analyze this problem, it is instructive to consider how the standard result of Equation 4.3 arises directly from the partition function. Consider the contribution to the partition function of a large system of a state which has $N_{\{i\}}$ clusters of type $\{i\}$:

$$Q_{\{N_{\{i\}}\}}(D) = \prod_{\{i\}} \frac{q_{\{i\}}^{N_{\{i\}}}}{N_{\{i\}}!} = \prod_{\{i\}} \frac{(Dz_{\{i\}} / \prod_x^y i_x!)^{N_{\{i\}}}}{N_{\{i\}}!}. \quad (\text{D.28})$$

In this expression, the $N_{\{i\}}!$ term arises to avoid double counting of states, and must be included because each $q_{\{i\}}$ includes a separate integral for each cluster over the whole of the system volume. Maximizing $Q_{\{N_{\{i\}}\}}$ with respect to $N_{\{i\}}$ yields Equation 4.3.

I now consider a system in which one of the reactants is immobilized (let this be particle type 1).¹ The first consequence is that there is no need to divide by $N_{\{i\}}!$ when the cluster $\{i\}$ includes the immobilized species, as there is no tendency to double count when the clusters cannot move over all space. One must, however, still deal with combinatorial effects. In particular, I now have to calculate the combinatorial factor associated with the number of different choices of immobilized particles that are involved in cluster formation. This

¹I will also assume that *all* particles of type 1 are localized.

introduces a factor:

$$\frac{N_1^{\text{total}!}}{\prod_{\{i\}, i_1 \neq 0} N_{\{i\}}!} \quad (\text{D.29})$$

The second effect is that $q_{\{i\}}$ does not scale with system volume for $i_1 \neq 0$, so that:

$$q_{\{i\}} = \frac{z_{\{i\}}}{\prod_x^y i_x!} \quad \text{for } i_1 \neq 0. \quad (\text{D.30})$$

Combining these two differences, I obtain:

$$\begin{aligned} Q_{\{N_{\{i\}}\}}^{\text{immob.}}(D) &= \left(\prod_{\{i\}, i_1=0} \frac{q_{\{i\}}^{N_{\{i\}}}}{N_{\{i\}}!} \right) \left(\prod_{\{i\}, i_1 \neq 0} q_{\{i\}}^{N_{\{i\}}} \right) \left(\frac{N_1^{\text{total}!}}{\prod_{\{i\}, i_1 \neq 0} N_{\{i\}}!} \right). \\ &= \left(\prod_{\{i\}, i_1=0} \frac{(Dz_{\{i\}} / \prod_x^y i_x!)^{N_{\{i\}}}}{N_{\{i\}}!} \right) \left(\prod_{\{i\}, i_1 \neq 0} \frac{(z_{\{i\}} / \prod_x^y i_x!)^{N_{\{i\}}}}{N_{\{i\}}!} \right) N_1^{\text{total}!} \end{aligned} \quad (\text{D.31})$$

It is trivial to check that for two sets of clusters $\{N_{\{i\}}\}$ and $\{N'_{\{i\}}\}$, $Q_{\{N_{\{i\}}\}}^{\text{immob.}}/Q_{\{N'_{\{i\}}\}}^{\text{immob.}}$ has exactly the same functional dependence on $z_{\{i\}}$, $N_{\{i\}}$ and $N'_{\{i\}}$ as $Q_{\{N_{\{i\}}\}}/Q_{\{N'_{\{i\}}\}}$. Therefore there are no statistical consequences of tethering one of the reactants – a given set of $z_{\{i\}}$ will produce identical bulk yields to the case when all species are free to diffuse.

Physically, this is because in this idealized limit the only important coordinates are the relative separation of cluster-forming particles. Consequentially, it is irrelevant that particles of type 1 are tethered, as the concentration fluctuations of the other particles in the vicinity of type 1 provide the same statistical correction as the untethered case.

Such an argument does not hold if two particles involved in an assembly are localized. For a trivial counter-example, one could take heterodimer formation. Localizing both species will give thermodynamics identical to the small system limit.

When performing simulations of this kind, it is possible that non-tethered particles will interact with the tethering mechanism in the unbound state. If this has a significant effect on the unbound partition function, it will lead to errors in the extrapolation. Such effects can be checked by changing the simulation volume and observing whether the statistics of bound states change in the expected way.

Appendix E

Details of sampling methods for DNA tweezers

E.1 Sequences

The strand sequences used in simulations are given below in the 5'-3' convention.

Hinge (h): AGCT TCGA CCTT TTAG GGCC TTAT.

Tweezer arm 1 (α): GTCA GCCA ATAA GGCC CT.

Tweezer arm 2 (β): GGTC GAAG CTT C GAAG CT.

Fuel (f): AGCT TCGA TGGC TGAC CTTA TTCA.

Antifuel (\bar{f}): TGAA TAAG GTCA GCCA TCGA AGCT.

E.2 Definition of Q_5

As discussed in Chapter 7, a reaction coordinate \mathbf{Q} was used to bias the ensemble of the tweezers to facilitate sampling. The first four dimensions of \mathbf{Q} were simply given by:

- Q_1 : the number of correct base pairs between α and f.
- Q_2 : the number of correct base pairs between β and f.
- Q_3 : the number of correct base pairs between f and \bar{f} , restricted to the bases of f that bind to α or are in the toehold.

Q_5 value	bp	Separation
0	$Q_1 = Q_2 = Q_3 = Q_4 = 0$	$f.\alpha > 4 \text{ AND } f.\beta > 4$
1	$(Q_1 = 0 \text{ OR } Q_2 = 0) \text{ AND } Q_3 = Q_4 = 0$	$f.\alpha > 4 \text{ XOR } f.\beta > 4$
2	$(Q_1 = 0 \text{ OR } Q_2 = 0) \text{ AND } Q_3 = Q_4 = 0$	$f.\alpha < 4 \text{ AND } f.\beta < 4$
3	$Q_1 > 0 \text{ AND } Q_2 > 0 \text{ AND } Q_3 = Q_4 = 0$	$f.\bar{f}_{\text{toehold}} > 4$
4	$Q_1 > 0 \text{ AND } Q_2 > 0 \text{ AND } Q_4 = 0$	$f.\bar{f}_{\text{toehold}} < 4$
5	$Q_2 > 0 \text{ AND } Q_3 > 0 \text{ AND } Q_1 = Q_4 = 0$ AND No bp between f_8 and \bar{f}_{15}	$f_8.\alpha_7 < 3$
6	$Q_2 > 0 \text{ AND } Q_3 > 0 \text{ AND } Q_1 = Q_4 = 0 \text{ AND}$ $((\text{No bp between } f_8 \text{ and } \bar{f}_{15} \text{ AND } 3 < f_8.\alpha_7 < 5)$ $\text{OR } (\text{bp between } f_8 \text{ and } \bar{f}_{15} \text{ AND } f_8.\alpha_7 < 3))$	
7	$Q_2 > 0 \text{ AND } Q_3 > 0 \text{ AND } Q_1 = Q_4 = 0 \text{ AND}$ (bp between f_8 and \bar{f}_{15} AND $3 < f_8.\alpha_7 < 5$)	
8	$Q_2 > 0 \text{ AND } Q_3 > 0 \text{ AND } Q_1 = Q_4 = 0$	$f_8.\alpha_7 > 5$
9	$Q_2 > 0 \text{ AND } Q_3 > 0 \text{ AND } Q_1 = 0 \text{ AND } Q_4 > 0$	none
10	$Q_3 > 0 \text{ AND } Q_4 > 0 \text{ AND } Q_1 = Q_2 = 0$ AND No bp between f_0 and \bar{f}_{23}	$f_0.\beta_{17} < 3$
11	$Q_3 > 0 \text{ AND } Q_4 > 0 \text{ AND } Q_1 = Q_2 = 0 \text{ AND}$ $((\text{No bp between } f_0 \text{ and } \bar{f}_{23} \text{ AND } 3 < f_0.\beta_{17} < 5)$ $\text{OR } (\text{bp between } f_0 \text{ and } \bar{f}_{23} \text{ AND } f_0.\beta_{17} < 3))$	
12	$Q_3 > 0 \text{ AND } Q_4 > 0 \text{ AND } Q_1 = Q_2 = 0 \text{ AND}$ (bp between f_0 and \bar{f}_{23} AND $3 < f_0.\beta_{17} < 5$)	
13	$Q_3 > 0 \text{ AND } Q_4 > 0 \text{ AND } Q_1 = Q_2 = 0$	$f_0.\beta_{17} > 5$
14	Any forbidden state (see Section E.3)	none

Table E.1: Table showing the definition of Q_5 . In this table, the “.” symbol indicates the minimum distance, measured in simulation units, between any two bases which can form a correct base pair for the strands indicated. For example, $f.\beta < 4$ means that at least two of the fuel and β -arm bases that can be paired in the closed tweezer state are within 4 simulation units. Subscript indicates that a specific base or bases (counting from the 5' end, starting from 0), rather than the entire strand, should be considered.

- Q_4 : the number of correct base pairs between f and \bar{f} , restricted to the bases of f that bind to β .

Here, a base pair was considered to be formed if the H-bond energy between nucleotides was below -0.1 in reduced units. A fifth coordinate, which explicitly depended on the separation of strands as well as the number of base pairs, was also defined. Its definition is given in Table E.1. The definition is fairly complex, but the general idea is to favourably bias states in which strands are close but not yet paired. The most difficult transitions to sample involve the completion of displacement of either arm, as it is necessary to sample

Window	Q_5														Other	
	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14	
α	✓	✓	✓													
β		✓	✓	✓	✓											$Q_3 = 0$
γ				✓	✓											$Q_3 \leq 8$
δ					✓											$8 \geq Q_1 \geq 5$
ϵ						✓										$6 \geq Q_1 \geq 3$
ζ							✓									$4 \geq Q_1 \geq 1$
η								✓	✓	✓	✓	✓				$Q_1 \leq 3$
θ								✓	✓	✓	✓	✓				$8 \geq Q_2 \geq 5$
ι								✓	✓	✓	✓	✓				$6 \geq Q_2 \geq 3$
κ								✓	✓	✓	✓	✓				$4 \geq Q_2 \geq 1$
λ											✓	✓	✓	✓	✓	$Q_2 \leq 3$

Table E.2: Values of the order parameter to which sampling was restricted in each simulation window.

the rebinding of the arms to the fuel (which requires fraying of the f/\bar{f} duplex).

The particularly complex definitions for $Q_5 = 5 \rightarrow 8$ and $Q_5 = 10 \rightarrow 13$ are designed to favour the intermediate states of these rebinding processes, by encouraging the fraying of the f/\bar{f} duplex and increasing the proximity of the final displaced base of the relevant tweezer arm to the base from which it was displaced.

The landscape was split into several sampling windows for convenience. The range of each window is shown in Table E.2. Each window has significant overlap with the windows on either side, allowing the entire landscape to be recovered using the WHAM technique [157].

E.3 Restrictions to the ensemble

To simplify the sampling and facilitate the use of a windows, several restrictions were imposed on the system during the umbrella sampling simulations.

1. In all umbrella simulations, dissociation of the tweezer unit itself was forbidden.
2. In simulations involving \bar{f} , the complete dissociation of f and β was forbidden before the dissociation of f and α .

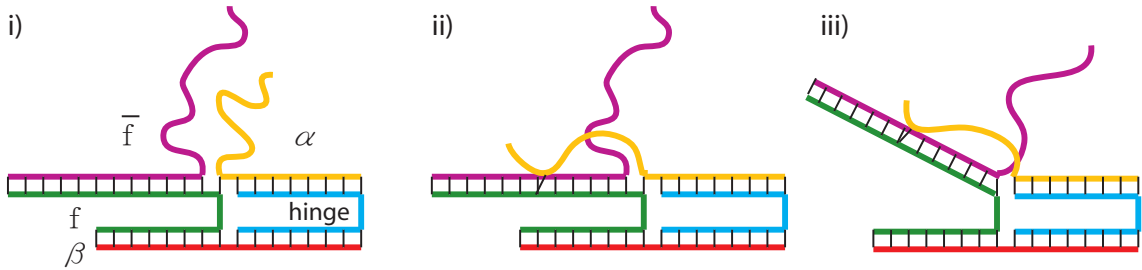


Figure E.1: Illustration of the formation of a pathological state in windowed sampling.

3. Letting X denote the base furthest from the toehold along f to which \bar{f} is bound, states with tweezer strands bound to f at bases between X and the toehold were explicitly forbidden.

To sample states violated by restrictions 1 and 2, it would have been necessary to drive the relevant dissociation/hybridization reactions, thereby complicating the sampling. I decided that the most honest approach was to explicitly forbid these possibilities, rather than simply fail to sample them.

Restriction 3 was based on the need to split the umbrella sampling into windows. I used umbrella windows designed specifically to sample the progression of the displacement of the α arm (windows γ to ζ), in which complete detachment from the fuel was forbidden, and a window designed to sample the release and reattachment equilibrium for the final few base-pairs of the α/f duplex (window η).

In an early simulation of a window in which the dissociation of α and f was forbidden, the situation illustrated in Figure E.1 occurred. (i) The α/f duplex was reduced to one base pair. (ii) A weak bond then formed further up the α arm to a base of f already bound to \bar{f} (a bound base can form transient, weak bonds to other bases in our model). (iii) In this instance, the original α/f base pair broke first, leaving α and f weakly attached. This state actually has a low free energy compared to others in the simulation window, as it maximizes the number of strong base pairs, but the bond would have quickly broken and allowed the tweezers to open in a simulation in which dissociation was possible. In this case, this easy escape was prohibited, and so the simulation was essentially stuck (although low in free energy, the state was hard to access, requiring a peculiar sequence of events - it was

Windows	\mathbf{Q}	test statistic	Windows	\mathbf{Q}	test statistic
α/β	(8,0,0,0,1)	0.0413	ζ/η	(2,8,14,0,4)	0.8182
α/β	(0,8,0,0,1)	0.6610	ζ/η	(1,8,15,0,4)	0.3882
β/γ	(8,8,0,0,3)	0.8951	η/θ	(0,8,16,0,6)	0.5098
β/γ	(7,7,0,0,3)	0.9349	η/θ	(0,8,16,0,8)	0.6719
γ/δ	(8,8,8,0,4)	0.9435	θ/ι	(0,6,16,2,9)	0.8249
γ/δ	(7,7,7,0,4)	0.9316	θ/ι	(0,5,16,3,9)	0.3024
δ/ϵ	(6,8,10,0,4)	0.7710	ι/κ	(0,4,16,4,9)	0.2573
δ/ϵ	(5,8,11,0,4)	0.7278	ι/κ	(0,3,16,5,9)	0.9430
ϵ/ζ	(4,8,12,0,4)	0.6722	κ/λ	(0,2,16,6,9)	0.6473
ϵ/ζ	(3,8,13,0,4)	0.7310	κ/λ	(0,1,16,7,9)	0.9651

Table E.3: Test statistics obtained from comparing the distribution of $\langle E(\mathbf{Q}) \rangle$ in overlapping windows.

consequentially also difficult to escape from). Rather than attempting to force the sampling of these states, it was decided to impose restriction 3 to prevent them forming.

To ensure that the neglected states were not significant in the operational cycle of the tweezers, simulations were also performed in which no artificial weighting or bias was imposed. Many complete (irreversible) cycles were recorded: no violations of restrictions 1 and 2 were observed. Rare, transient violations of the restriction 3 were observed, but they had no noticeable affect on the displacement processes.

E.4 Comparing $\langle E(\mathbf{Q}) \rangle$ from different windows

To be certain that windowing of simulations did not affect sampling accuracy by limiting ergodicity, as outlined in Chapter 7, values of $\langle E(\mathbf{Q}) \rangle$ were compared for two commonly sampled values of \mathbf{Q} for each overlap region between adjacent windows. The distributions of $E(\mathbf{Q})$ measured in the simulation were tested for significance using Welch's unpaired t-test [214]. The results, shown in Table E.3, are as would be expected for consistent sampling (only one order parameter appears to show significant difference at the 95% level, as would be expected from 20 tests).

Appendix F

Details of sampling methods for the DNA walker

Three types of simulation were performed on the DNA walker system presented in Chapter 8. Firstly, simple unbiased Langevin and VMMC simulations were performed to study the kinetics of foot-binding, and also to explore the possibility of lifting of the wrong foot. To obtain thermodynamic data, VMMC simulations were performed with biases that are discussed in the following section. Finally, Langevin simulations using forward flux sampling were used to study the possible mechanism of lifting the wrong foot from the track.

F.1 Order parameters used in thermodynamic simulations

Generally, order parameters were primarily constructed using the number of hydrogen-bonding interactions with energies below -0.1 (in reduced units). An additional dimension, analogous to the Q_5 introduced for the tweezers (see Appendix E), was sometimes used to favour bringing duplexes into close proximity. For all simulations considered in this section, base pairs were deemed to be formed if hydrogen-bonding energy was < -0.1 simulation units.

- **Front site binding:** 3-dimensional order parameter.
 - Q_1 : the number of correct bp between foot 2^1 and the front site.

¹the initially raised foot

- Q_2 : the number of misbonded bp between foot 2 and the front site.
- Q_3 : based on the separation of foot 2 from the front site when in the unbound state.

In these simulations, binding to the back site was forbidden, as was the dissociation of the initially attached foot from the track. Correct bonds and misbonds with the front site were limited to 14 and 2 bp respectively.

- **Back site binding** simulations were performed exactly analogously to the front site calculations.
- **Competition between toehold domains:** 3-dimensional order parameter.

- Q_1 : the number of correct bp between foot 1 and the middle site.
- Q_2 : the number of correct bp between foot 2 and the front site.
- Q_3 : if either of the bp at the ends of the duplexes away from the competition region had frayed, $Q_3 = 1$, otherwise $Q_3 = 0$.

To encourage migration of the displacement junction, states with one fewer than the maximum number of bonds were favourably biased. The unintended result of this bias (when first applied) was that additional fraying was observed at the far ends of the walker/track duplexes, rather than fraying near the junction point. Q_3 was therefore included to distinguish between states with and without frayed ends, allowing fraying at the displacement junction to be selectively favoured. In these simulations, detachment of either foot was prohibited, as was any binding to the back site.

- **Fuel binding:** 6-dimensionnal order parameter.
- Q_1 : the number of correct bp between foot 1 and the middle site.
- Q_2 : the number of correct bp between foot 2 and the front site.
- Q_3 : if either of the bp at the ends of the duplexes away from the competition region had frayed, $Q_3 = 1$, otherwise $Q_3 = 0$.

- Q_4 : the number of correct bp with the competing toehold domain and the non-toehold domain of the back foot.
- Q_5 : the number of correct bp with the competing toehold domain of the front foot.
- Q_6 : based on the separation of the fuel from the toehold domain of the feet.

Both fuel-binding dimensions were limited to a maximum of 6 bp, and detachment from the track of either foot was prohibited. An additional dimension was included to bias the fuel towards approaching the toehold domains. In these simulations, binding to the back site was prohibited and incorrect pairing involving the fuel or foot 2 was limited to 2 bp.

- **Displacement:** 3-dimensional order parameter.

- Q_1 : the number of correct bp between foot 1 and the middle site.
- Q_2 : the number of correct bp between foot 1 and the fuel.
- Q_3 : if either of the bp at the ends of the foot 1/fuel or foot 1/track duplexes away from the competition region had frayed, $Q_3 = 1$, otherwise $Q_3 = 0$.

Migration of the branch point was accelerated in a manner analogous to the simulation of the competition domain. Once again, binding to the back site and total detachment of either foot from the track was prohibited, as was the detachment of fuel from the back foot. Incorrect pairing involving the fuel or the front foot was limited to 2 bp. In order to sample the two mismatch repairing steps, it was found to be advantageous to split the displacement into two windows that were combined using the WHAM algorithm [157].

- **Foot lifting:** 3-dimensional order parameter.

- Q_1 : the number of correct bp between foot 1 and the middle site.
- Q_2 : the number of correct bp between foot 1 and the fuel.

- Q_3 : based on the separation of foot 1 and the back site when in the unbound state.

In these simulations, detachment of foot 2 from the track, detachment of the fuel from foot 1 and any binding to the back site were forbidden. Bonds between foot 1 and the middle site were limited to 7 or fewer, and incorrect pairing involving the fuel or the front foot was limited to 2 bp. As the dimensions used for foot lifting were not identical with the displacement simulations, WHAM cannot be used to combine the two sets of data. Instead, the data was condensed onto a 1-dimensional axis (the number of bp between back foot and the middle site of the track) for both windows, and a near-perfect agreement in the overlap region used to combine the data to produce Figure 8.8 (b).

As with the tweezers, the restrictions to the system were imposed to ensure that accurate and well-defined sampling could be performed. In this case, unbiased simulations have revealed a number of effects not visible in the restricted thermodynamics simulations – such as the possibility of binding proceeding via misbonds. Nonetheless, the thermodynamic simulations provide a good starting point in understanding the kinetic results, such as the bias for binding correctly to the front site under tension.

F.2 Forward flux sampling of lifting the front foot

In Chapter 8, a possible mechanism for lifting the front foot after binding to its raised toehold is suggested. Once bound to the raised toehold, the fuel can either detach or reach round and begin to displace the track from the front foot. I estimated the relative probability of these two events occurring using forward flux sampling. Forward flux sampling is discussed in detail in Chapter 3.

Unlike umbrella sampling, forward flux sampling does not easily generalize to a transition which can only be well described by a two-dimensional order parameter. In this case, I am essentially interested in finding the relative probability of fuel dissociating normally from the front toehold compared to fuel unbinding from the toehold whilst bound to another

part of the foot, leading to displacement of the track. A natural order parameter, therefore, would have one dimension related to the binding of the fuel to the toehold and another describing the binding of the fuel to the rest of the foot.

To simplify the requirements of the algorithm, I used forward flux sampling to measure the flux from a state in which the fuel is bound to the toehold to one in which it has dissociated, and separately to one in which it has formed 8 bp with the non-toehold region (but still attached to the toehold). Such a partially-displaced state was observed to be a metastable minimum of free energy, corresponding to a ‘double-X’ configuration. Individually, both processes are relatively simple to describe using a one-dimensional order parameter. I then assumed that trajectories that reached the metastable minimum of partial displacement tend to equilibrate within the local minimum before either the fuel detaches from the toehold or the non-toehold region. Using this assumption, I ran separate simulations initiated from states with 8 bp between the non-toehold region and the fuel to calculate the probability of successful foot-raising after such a state has been reached.²

F.2.1 Measuring the melting flux

I measured the flux from a state in which the fuel is bound to the raised toehold of the front foot to a state in which the hydrogen-bonding sites of the fuel are separated by at least 3 simulation units from the hydrogen-bonding sites of the toehold. The order parameter definitions and simulation results are given in Table F.1.

Multiplying the initial flux by the probabilities gives a melting flux of $\Phi_{\text{melt}} = 1.24 \times 10^{-7}$ per unit simulation time.³ An error on this quantity can be estimated in the following manner. 10 simulations were performed to estimate the initial flux, with a standard error on the mean of 1.43×10^{-4} per unit time (compared to an average of 7.33×10^{-3} per unit time). The later stages can be modelled as Bernoulli trials – the probability of success

²It should be noted that the estimate of displacement only includes trajectories that reach 8 bp between the non-toehold region and the fuel prior to dissociation of the fuel from the toehold. It is possible that displacement may occur without such a state being reached, and also that there are completely different pathways to displacement that are not sampled due to the chosen order parameter.

³Due to the caveats with comparing simulation time to absolute time, I will report all fluxes in terms of the simulation time, and only compare relative values.

Q	Definition	Simulation results	
$Q = -2$	6 bp		
$Q = -1$	5 bp	crossing events:	steps:
$Q = 0$	3 or 4 bp	10618	4.82×10^8
		attempts:	successes:
$Q = 1$	1 or 2 bp	20000	1464
$Q = 2$	no bp and ≥ 1 nearly formed bp	9800	360
$Q = 3$	no bp and no nearly formed bp	2000	1724
$Q = 4$	separation ≥ 3 units	2000	1451

Table F.1: Order parameter definitions and simulation results for the melting detachment of the fuel from the front-foot toehold domain. For these simulations, only correct bp between toehold and fuel were considered and were counted as being formed if their energy was less than 0. “Nearly formed” bp were recorded if a pair of these nucleotides had hydrogen-bonding sites within 0.9 simulation units and would have had negative energy but for a single factor in the hydrogen-bonding term being zero. The time step used was 0.003 simulation units.

measured after N attempts has a variance of $p(1 - p)/N$, where p is the true probability of success. The measured p can then be used to estimate the standard error on p for each stage. Adding the relative errors in quadrature gives a standard error of around 6.4%.

F.2.2 Measuring the flux to a partially displaced state

The alternative to directly melting is for the fuel strand to begin displacement of the track from the foot, by reaching back to form a double-X-like structure. 8 bp with the track was observed to be a metastable minimum of free energy, corresponding to such a configuration. I measured the flux to this state from a state in which the fuel is only bound to the toehold, with the order parameter and results outlined in Table F.2.

Multiplying the initial flux by the probabilities for each stage gives a total flux to the partially displaced state of $\Phi_{\text{partial}} = 8.77 \times 10^{-10}$ per unit simulation time. From this point, either of the fuel/toehold or the fuel/non-toehold duplexes can melt. Whenever the first case occurred, full displacement of the track and foot lifting was always observed. In the other case, the system had returned to the original state of fuel binding and displacement had

Q	Definition	Simulation results	
$Q = -2$	separation between correct bp of fuel and non-competition domain > 4 units		
$Q = -1$	$4 \geq$ separation > 2 units	crossing events:	steps:
$Q = 0$	$2 \geq$ separation > 1 unit	3850	3.84×10^9
		attempts:	successes:
$Q = 1$	separation ≤ 1 and no bp or nearly-formed bp	20000	1798
$Q = 2$	no bp and ≥ 1 nearly formed bp	4500	1343
$Q = 3$	1 or 2 bp	20000	824
$Q = 4$	3 or 4 bp	20000	956
$Q = 5$	5,6 or 7 bp	3573	462
$Q = 6$	≥ 8 bp	5000	2020

Table F.2: Order parameter definitions and simulation results for the initiation of wrong-foot displacement after the fuel has bound to the front-foot toehold domain. For these simulations, only correct bp were considered and were counted as being formed if their energy was less than -0.5 in simulation units. ‘Nearly formed’ bp were recorded if a pair of these nucleotides had a hydrogen-bonding energy of $0 > E \geq -0.5$, or had hydrogen-bonding sites within 0.9 simulation units and would have had negative energy but for a single factor in the hydrogen-bonding term being zero. The time step used was 0.003 simulation units.

failed. Both processes are reasonably fast (being in the double-X configuration destabilizes the fuel/toehold duplex), so the relative rates could be estimated simply by running unbiased simulations starting from the intermediate displacement state. Of 50 simulations initiated from the partially displaced state, 25 completed the displacement and 25 returned to the original state of having the fuel bound to the track only (one failed to do either in the simulation time). Combining this estimate with the flux to the partially displaced state, I obtain a flux to successful displacement from a state with fuel bound to the walker toehold of $\Phi_{\text{displacement}} = 4.39 \times 10^{-10}$ per unit simulation time, with an error of around 16.2%. Thus,

$$\frac{\Phi_{\text{displacement}}}{\Phi_{\text{melt}}} = 0.0035, \quad (\text{F.1})$$

with an error of around 17.2%.

Thesis errata

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27/4/2012

For anyone intending to use the description of the model in this text to simulate said model, the following corrections are important.

1. The final line of Equation 2.16 should have additional minus signs, giving:

$$\begin{aligned}
 V_{\text{stack}} = & f_1(\delta r_{\text{stack}}, \epsilon_{\text{stack}}, a_{\text{stack}}, \delta r_{\text{stack}}^0, \delta r_{\text{stack}}^{c,\text{low}}, \delta r_{\text{stack}}^{c,\text{high}}, \delta r_{\text{stack}}^{\text{low}}, \delta r_{\text{stack}}^{\text{high}}) \\
 \times & f_4(\theta_4, a_{\text{stack},4}, \theta_{\text{stack},4}^0, \Delta\theta_{\text{stack},4}^*) \\
 \times & f_4(\theta_5', a_{\text{stack},5}, \theta_{\text{stack},5}^0, \Delta\theta_{\text{stack},5}^*) f_4(\theta_6', a_{\text{stack},6}, \theta_{\text{stack},6}^0, \Delta\theta_{\text{stack},6}^*) \\
 \times & f_5(-\cos(\phi_1), a_{\text{stack},1}, -\cos(\phi_1)_{\text{stack}}^*) f_5(-\cos(\phi_2), a_{\text{stack},2}, -\cos(\phi_2)_{\text{stack}}^*).
 \end{aligned}$$

2. The V_{stack} entry into Table 2.1 should have additional minus signs, giving:

Interaction		Parameters		
V_{stack}	$f_1(\delta r_{\text{stack}})$	$\epsilon_{\text{stack}} = 1.3448$ $+2.6568 kT$	$a_{\text{stack}} = 6$	$\delta r_{\text{stack}}^0 = 0.4$
		$\delta r_{\text{stack}}^c = 0.9$	$\delta r_{\text{stack}}^{\text{low}} = 0.32$	$\delta r_{\text{stack}}^{\text{high}} = 0.75$
	$f_4(\theta_4)$	$a_{\text{stack},4} = 1.30$	$\theta_{\text{stack},4}^0 = 0$	$\Delta\theta_{\text{stack},4}^* = 0.8$
	$f_4(\theta_5')$	$a_{\text{stack},5} = 0.90$	$\theta_{\text{stack},5}^0 = 0$	$\Delta\theta_{\text{stack},5}^* = 0.95$
	$f_4(\theta_6')$	$a_{\text{stack},6} = 0.90$	$\theta_{\text{stack},6}^0 = 0$	$\Delta\theta_{\text{stack},6}^* = 0.95$
	$f_5(-\cos(\phi_1))$	$a_{\text{stack},1} = 2.00$	$-\cos(\phi_1)_{\text{stack}}^* = -0.65$	
	$f_5(-\cos(\phi_2))$	$a_{\text{stack},2} = 2.00$	$-\cos(\phi_2)_{\text{stack}}^* = -0.65$	

3. Equation (A.7) should contain additional minus signs, giving:

$$\begin{aligned}
 & \frac{dV_{\text{FENE}}(\delta r_{\text{backbone}})}{d\delta r_{\text{backbone}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{backbone}}) + \sum_{x,y*} \frac{df_3(\delta r_{xy})}{d\delta r_{xy}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{xy}) \\
 & + \frac{V_{\text{stack}}}{f_1(\delta r_{\text{stack}})} \frac{df_1(\delta r_{\text{stack}})}{d\delta r_{\text{stack}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{stack}}) + \sum_{i=4,5',6'} \frac{V_{\text{stack}}}{f_4(\theta_i)} \frac{df_4(\theta_i)}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
 \nabla_{\mathbf{q},\mathbf{r}} V_{\text{neighbour}} = & + \frac{V_{\text{stack}}}{f_5(-\cos(\phi_1))} \frac{df_5(-\cos(\phi_1))}{d\cos(\phi_1)} \nabla_{\mathbf{q},\mathbf{r}}(\cos(\phi_1)) \\
 & + \frac{V_{\text{stack}}}{f_5(-\cos(\phi_2))} \frac{df_5(-\cos(\phi_2))}{d\cos(\phi_2)} \nabla_{\mathbf{q},\mathbf{r}}(\cos(\phi_2)),
 \end{aligned}$$

4. Equation (A.8) contains a mistake: some f_4 functions should be replaced by f_5 , giving:

$$\begin{aligned}
& \sum_{x,y} \frac{df_3(\delta r_{xy})}{d\delta r_{x,y}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{xy}) + \frac{V_{\text{HB}}}{f_1(\delta r_{\text{HB}})} \frac{df_1(\delta r_{\text{HB}})}{d\delta r_{\text{HB}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{HB}}) \\
& + \sum_{i=1-4,7,8} \frac{V_{\text{HB}}}{f_4(\theta_i)} \frac{df_4(\theta_i)}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
& + \frac{V_{\text{cross_stack}}}{f_2(\delta r_{\text{HB}})} \frac{df_2(\delta r_{\text{HB}})}{d\delta r_{\text{HB}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{HB}}) + \sum_{i=1}^3 \frac{V_{\text{cross_stack}}}{f_4(\theta_i)} \frac{df_4(\theta_i)}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
& + \sum_{i=1,7,8} \frac{V_{\text{cross_stack}}}{f_4(\theta_i)+f_4(\pi-\theta_i)} \frac{d(f_4(\theta_i)+f_4(\pi-\theta_i))}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
\nabla_{\mathbf{q},\mathbf{r}} V_{\text{non-neighbour}} = & + \frac{V_{\text{coax_stack}}}{f_2(\delta r_{\text{stack}})} \frac{df_2(\delta r_{\text{stack}})}{d\delta r_{\text{stack}}} \nabla_{\mathbf{q},\mathbf{r}}(\delta r_{\text{stack}}) + \frac{V_{\text{coax_stack}}}{f_4(\theta_4)} \frac{df_4(\theta_4)}{d\theta_4} \nabla_{\mathbf{q},\mathbf{r}}(\theta_4) \\
& + \frac{V_{\text{coax_stack}}}{f_4(\theta_1)+f_4(2\pi-\theta_1)} \frac{d(f_4(\theta_1)+f_4(2\pi-\theta_1))}{d\theta_1} \nabla_{\mathbf{q},\mathbf{r}}(\theta_1) \\
& + \sum_{i=5}^6 \frac{V_{\text{coax_stack}}}{f_4(\theta_i)+f_4(\pi-\theta_i)} \frac{d(f_4(\theta_i)+f_4(\pi-\theta_i))}{d\theta_i} \nabla_{\mathbf{q},\mathbf{r}}(\theta_i) \\
& + \sum_{i=3}^4 \frac{V_{\text{coax_stack}}}{f_5(\cos(\phi_i))} \frac{df_5(\cos(\phi_i))}{d\cos(\phi_i)} \nabla_{\mathbf{q},\mathbf{r}} \cos(\phi_i),
\end{aligned}$$