Sequence specificity of single-stranded DNA-binding proteins: a novel DNA microarray approach

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ABSTRACT
We have developed a novel DNA microarray-based approach for identification of the sequence-specificity of single-stranded nucleic-acid-binding proteins (SNABPs). For verification, we have shown that the major cold shock protein (CspB) from Bacillus subtilis binds with high affinity to pyrimidine-rich sequences, with a binding preference for the consensus sequence, 5'-GTCTTTG/T-3'. The sequence was modelled onto the known structure of CspB and a cytosine-binding pocket was identified, which explains the strong preference for a cytosine base at position 3. This microarray method offers a rapid high-throughput approach for determining the specificity and strength of ss DNA–protein interactions. Further screening of this newly emerging family of transcription factors will help provide an insight into their cellular function.

INTRODUCTION
We show that microarray technology can provide rapid high-throughput assays for the identification of sequence-specific ss DNA–protein interactions. The majority of transcription factors recognize target sequences in duplex form. However, single-stranded regions can be induced by torsional stress of double-stranded DNA, allowing single-stranded nucleic-acid-binding proteins (SNABPs) access to their binding sites (1). SNABPs have been shown to bind with high affinity, non-specifically (2) and specifically (3), to ss DNA, which has been shown to regulate gene expression both positively and negatively (1,4). Gene expression can also be regulated on a translational level, by SNABP binding to mRNA (5).

Genome sequencing has allowed SNABPs to be identified and characterized for a range of eukaryotic and prokaryotic organisms. To understand how binding of SNABPs to ss nucleic acids regulates transcription and translation, the regions of sequence specificity must be identified. Many techniques including electrophoretic mobility shift assay (EMSA) (6), nitrocellulose-binding assays (7), Southern blotting (8), phage display (9), UV cross-linking (10) and X-ray crystallography (11) were developed to study sequence-specific ss DNA–protein interactions. Available techniques including, fluorescence measurements (12), polymerase chain reaction (PCR), fluorescence resonance energy transfer (FRET) combined with a DNA foot-printing assay (13), surface plasmon resonance (SPR) and fluorescence polarization (14) have all been used to study effectively specific ss DNA–protein interactions. The most frequent approach used to study the sequence specificity of DNA-binding molecules is by systematic evolution of ligands by exponential enrichment (SELEX), this method allows for the identification of sequences which bind with high affinity to the molecule of interest (15). This method has been used mostly to select for double-stranded DNA molecules that bind to the target but it has been also used to screen ss DNA molecules (16,17). SELEX has advantages to the previous methods but still lacks in its ability for high-throughput analysis as numerous microarray experiments can be completed in a single day, thus providing a detailed analysis of binding-site recognition at an unparalleled rate.

These techniques made use of non-immobilized ss DNA in liquid phase to probe ss DNA interactions with other molecules such as proteins, drugs and ligands, all of which suffered from being time-consuming, laborious, expensive and incapable of high-throughput screening. Therefore, oligonucleotides immobilized to solid supports provide an
important tool for the rapid high-throughput examination of sequence-specific DNA–protein interactions.

Two current studies (18,19) have used microarrays displaying all possible 8-mer and 10-mer DNA duplexes to study effectively the sequence-recognition of both transcription factors and small molecules. These methods illustrate the potential high-throughput use of k-mer arrays in examining the DNA–binding properties of duplex-binding molecules but leave the area of SNABP specificity unexplored.

The innovative high-throughput assay described here provides a parallel screening system for identifying the specificity of SNABP binding. The major cold shock protein from *Bacillus subtilis* (CspB) was used to develop this microarray-based assay. This protein influences transcription and translation *in vitro* (20) by binding to stretches of 6–7 nucleotides (21) of ss DNA with a high degree of specificity (21,22).

We have used an oligonucleotide chip, for the identification of high-affinity 6-mer binding motifs for CspB. The chip contains all possible 4096 ss hexadeoxynucleotides incorporated onto a standardized anchor. The oligonucleotides on the array were originally designed to hybridize to folded mRNA which requires a significant spacer between the array surface and the recognition hexamer (Figure D, supplementary).

The use of a competitor protein in this assay allowed the identification of high-affinity DNA-binding sites. The binding affinity of a competitor protein will limit the amount of ss DNA-binding sites available to the CspB. The competitor protein chosen was a single-stranded DNA-binding protein from the crenarchaeote *Sulfolobus solfataricus* (SsoSSB). SsoSSB has a molecular weight of 16 kDa and binds non-specifically to ss DNA with a binding density of 5 nt per monomer and an apparent dissociation constant ($K_d$) of $\sim 90$ nM (2). Thus, the competitive binding of the SsoSSB protein provided a means of identifying high-affinity consensus binding motifs for CspB by reducing non-specific and weak CspB-ss DNA binding.

**MATERIALS AND METHODS**

**Microchip manufacture**

Oligonucleotide chips were supplied by Nyrion Ltd and contained all possible 4096 ss hexadeoxynucleotides incorporated into a general structure, 5'-NH2-C12-Spacer-AAAAAAA-NNNNNNNNN-XXXXXX-3', where $N$ was one of four bases and $X$ was a specific hexadeoxynucleotide. Each chip is made up of a 4 x 4 meta-grid and each of these sub grids contains 18 columns x 15 rows of spots, which are 135 µm ± 15% in diameter. Oligonucleotides were immobilized to the chip surface using standard Exiqon amino-link chemistry. All arrays were manufactured by pin spotting, according to complete standard commercial practices. This was all done under contract by MWG Biotech custom arrays. For control purposes, arrays are batch tested using a standard mRNA template and a standard QC procedure expected to give a standard signal. This standard signal serves as a positive and negative control for all arrays. MWG Biotech spot biotin on the surface of the array, which also serves as a negative standard control (generates zero signal).

**Expression and purification of recombinant SsoSSB**

A mutant version of the SsoSSB protein from the crenarchaeote *Sulfolobus solfataricus* was constructed by changing the C-terminal glutamate residue to a cysteine (E145C mutant), allowing for the incorporation of spin labels and fluorescent probes on the C-terminal tail. This mutation minimizes the effect of labelling on DNA-binding activity as the C-terminal glutamate is not involved in ss DNA binding. The E145C mutant was constructed only as a precaution if the amine-reactive labelling methods were unsuccessful. Protein expression was induced by addition of 0.2 mM IPTG at 37°C for 3 h, after which cells were pelleted and frozen until required.

Cell lysis, centrifugation and chromatography steps were carried out at 4°C. Cells (20 g) were thawed in 30 ml lysis buffer (50 mM Tris–HCl pH 7.5, 500 mM NaCl, 1 mM EDTA, 1 mM DTT) and immediately sonicated for 5 x 1 min with cooling. The lysate was centrifuged at 40 000 g for 45 min. DNase I [40 µg/ml] and RNase A [10 µg/ml]) were then added to the cell lysate and incubated at room temperature for 30 min with gentle agitation. The supernatant was heated to 70°C for 30 min in a water bath, and denatured proteins were precipitated by centrifugation at 40 000 g at 4°C. The supernatant was analysed by SDS–PAGE, and shown to contain recombinant SSB, which migrated as a band of $\sim 16$ kDa as expected. The supernatant was diluted 5-fold with buffer A (50 mM Tris–HCl pH 7.5, 1 mM EDTA, 1 mM DTT) and applied to a Heparin-Sepharose (Amersham) column equilibrated with buffer A. SsoSSB was eluted over a linear gradient comprising 0–1 M NaCl. Fractions corresponding to a distinct absorbance peak were analysed by SDS–PAGE, pooled and concentrated. Subsequent gel filtration step (HR 10/30 Superdex-200) in a buffer containing, 10 mM Tris/HCl pH 7.5, 150 mM NaCl, 1 mM EDTA and 1 mM DTT, resulted in essentially homogeneous SsoSSB, as determined by SDS–PAGE analysis. This method is an adaptation of the previously published method (23). SsoSSB was concentrated using a Viva Spin column (MWCO = 5 kDa) and quantified using both the Bradford method and the theoretical extinction coefficient, $\varepsilon_{280} = 12660$ M$^{-1}$. cm$^{-1}$.

**Cloning, expression and purification of recombinant His$_6$-CspB**

Primers B.S_CSP fwd (5′-dAGCCATATG TTA GAA GGT AAA GTA AAA TAA -3′) and B.S_CSP rev (5′-dCGGATCC TAA CGC TTC TTT AGT AAC GTT AGC-3′) were used in a PCR with plasmid DNA (pET11-CspB vector provided by Michael Wunderlich (University of Bayreuth)) containing the CspB gene. Bases were added to the primers to introduce the NdeI and BamHI restriction sites (underlined). These sites were used to clone the PCR product in NdeI-BamHI-digested pET28a vector, resulting in the plasmid pET28a_B.S_CspB.
BL21 (DE3)pLysS *Eschericia coli* (E. coli) was transformed with pET28_B.S._CspB and transformants were grown in Luria-Bertani medium, containing 50 µg/ml kanamycin at 37°C with agitation. One-litre cultures were grown to an OD₆₀₀ of 0.5–0.7 and IPTG (isopropyl-β-D-thiogalactosidase) was then added to a final concentration of 1 mM. Incubation was then continued for an additional 5–6 h and the cells were harvested at 8000 g in a JLA-9.1000 rotor for 12 min at 10°C. Pellets were then frozen in liquid nitrogen and stored at −80°C. Cell pellets were resuspended in lysis buffer (20 mM Tris HCl pH 8.0, 500 mM NaCl, 0.1% Triton X-100, 0.1 mM phenylmethylsulphonyl fluoride (PMSF), 1 mM EDTA and protease inhibitors) to a final volume of 30 ml/5 g of cells and then lysed using a French-press. DNase I [40 µg/ml] and RNase A [10 µg/ml] were then added to the cell lysate and incubated at room temperature for 30 min with gentle agitation. As an initial step, the fusion protein was purified using a Ni-NTA resin affinity column, as per manufacturer’s instructions and then purified to homogeneity as described previously (24). Briefly, to remove minor contaminants the fractions containing His₆-CspB were pooled and dialysed overnight into a buffer containing 20 mM Tris/HCl pH 6.8, 1 mM DTT. The solution was applied to an HR 5/5 Mono-Q (1 ml) anion exchange column. Bound protein was eluted with a NaCl gradient ranging from 0–1 M. CspB eluted at a concentration of 250 mM NaCl. A subsequent gel filtration step (HR 10/30 Superdex-75) in a buffer containing 10 mM Tris/HCl pH 7.5 and 100 mM NaCl resulted in visually pure His₆-CspB, as determined by SDS-PAGE analysis. His₆-CspB was concentrated using a Viva Spin column (MWCO = 3.5 kDa) and quantified using both the Bradford method and the theoretical extinction coefficient, ε₂₈₀ nm = 5690 M⁻¹ cm⁻¹.

**Electrophoretic mobility shift assay (EMSA)**

Three hundred picomoles of each ss DNA templates were 5'-end labelled by incubating templates with 0.03 mCi of [γ-³²P]ATP, T4 polynucleotide kinase and T4 polynucleotide kinase buffer in a total volume of 60 µl at 37°C for 2.5 h. The reaction was stopped by heat inactivation (30 min at 65°C). Unincorporated [γ-³²P]ATP was removed with QIAquick Nucleotide Removal Kit (Qiagen).

For a standard EMSA, 20 pmol of labelled ss DNA was mixed with increasing amounts of protein (total volume, 18 µl) at 4°C for 20 min in binding buffer (50 mM Tris, pH 8.0, 100 mM NaCl) unless stated otherwise. Two microlitres of dye solution (20% glycerol, 0.034% bromophenol blue) was added to the samples prior to gel electrophoresis.

Electrophoresis preformed in TBE (89 mM Tris, 89 mM boric acid, 2 mM EDTA, pH 8.0) buffer through a non-denaturing acrylamide gel (a 10% or 20% gel was used depending on the protein; e.g. for 50 ml of 20% gel; 25 ml of 40% acrylamide, 2.5 ml of 10 × TBE, 0.5 ml APS and 50 µl Temed) at 75 V until the samples had entered the gel and then at 100 V at 4°C (overnight for a 20% gel and 12 h for a 10% gel). Autoradiographs were obtained by exposing gels to Kodak BioMax MS film for 1–3 h at room temp.

**Labelling of ss DNA-binding proteins with Cy5 dye**

In the standard procedure, the contents of 1 vial (to label 1 mg of protein) of cyanine 5 (Cy5) mono-functional dye the contents (Amersham) were dissolved in 50 µl of anhydrous DMSO. Proteins were dialysed into buffer B (150 mM, Na₂CO₃ (pH 9.3, pH was adjusted with H₃PO₄)) and concentrated using a Viva Spin column (MWCO = 5 kDa). Typical working concentrations of proteins were 1 mg/ml, unless stated otherwise. Ten microlitres of dye/DMSO was pipetted into 200 µl protein solution under slow vortexing. After a 30-min incubation at 25°C in the dark, the reaction was terminated by the addition of 300 µl of 100 mM Na₂H₃PO₄ (to suppress further labelling) to the sample. To separate the unbound dye, the sample was loaded onto a PD-10 column (10 ml bed of Sephadex G-25M), which had been pre-equilibrated in a buffer A (100 mM NaCl, 50 mM Na₂H₃PO₄ and 1 mM EDTA (pH 7.5, pH adjusted with NaOH). The column was then washed with buffer A (2 × 1 ml) and the labelled protein was then eluted by adding 2 ml of water to the column. The extent of the modification was assessed using MALDI-TOF mass-spectrometry. Protein concentration was determined before and after labelling, Cy5-protein concentration was calculated as per manufacturer’s instructions.

**Protein hybridization**

Microarrays were pre-wet with phosphate-buffered saline (PBS) and 0.01% Triton X-100 and then blocked in 2% non-fat dried milk for 1 h. Blocked microarray slides were washed once with PBS and 0.1% Tween 20, and once with PBS and 0.01% Triton X-100. Protein (Cy5 labelled and unlabelled) binding to ss 25-mers (containing all possible [4096] 6-mer sequences) on a generic microchip was carried out in a hybridization chamber (Camlab, RTP/7870). Protein binding was performed in a humid chamber at 4°C with 80 µl of protein-binding reaction mixture containing: 50 mM KCl, 20 mM Tris (pH 8.0), 2% (w/v) non-fat dried milk, 0.2 µg/µl bovine serum albumin (BSA) and 40 µM of test protein. Slides were covered with a siliconized cover-slip (BDH cover glass 22 × 50 mm, Cat. No. 406/0188/42, Borosilicate Glass) and incubated for 1 h at 4°C. The cover-slip was removed and the slide was washed (3x) in a slide chamber filled with PBS and 0.05% Tween-20, with PBS and 0.01% Triton X-100 (3 ×) and once with PBS for 3 min each. Excess water was removed from the slide surface (by flicking), which was allowed to dry before scanning. This method is an adaptation of the previously published method (25). Various methods (denaturing conditions; including high temperatures, various concentrations of detergents and pH range in combination with high NaCl concentration) were used to remove bound protein from the chip surface in order to reuse the array but were all unsuccessful as they either had detrimental affects on ONs bound to the
surface of the array or were unable to remove bound protein.

**Competitive assay**

The method was essentially the same as above except that both His$_6$-CspB and SsoSSB proteins were added to the binding reaction mix at the specified molar ratio. The binding reaction was carried out as before. The array was then incubated for 1 h in a humid chamber at 4°C with 100 µl of diluted (1:100 in blocking buffer) Alexa 532-conjugated polyclonal antibody to His$_6$ (Molecular Probes). After incubation, the array was washed (3 ×) with PBS and 0.05% Tween-20 and once with PBS for 3 min each. Excess water was removed from the slide surface (by flicking), which was allowed to dry before scanning.

**Microarray analysis: data collection**

All microarray slides were scanned using an ArrayWorx microarray scanner at a range of laser settings, the highest of which produced a saturated signal for the majority of spots. The Alexa-532 (Cy3 equivalent) fluorophore was excited at 532 nm and the emission was recorded at 570 nm. The Cy5 fluorophore was excited at 633 nm and the emission was recorded at 675 nm. The data were filtered initially using a series of quality-control criteria so that only high-quality spots were used in our analysis. For each array we removed any flagged spots, these were spots that had dust flakes, scratches and irregular spots (spots that outmatched the average size). The average size of a spot is 135 µm ± 15% in size, any spot that did not correspond to this size constraint was excluded from the data. This size constraint also provided a crude method of approximating the DNA concentration of each spot, which allowed only spots with an optimum DNA concentration into the data collected. All microarray TIF images were quantified using Imagene Version 5.0 software.

**Microarray analysis: data processing**

The extent of background fluorescence was initially determined from an experiment using BSA. The level of background fluorescence from the spots and array surface was found to be similar. Therefore, the average fluorescence between spots on the array surface was used as the background value throughout the experiments, which was minimal in comparison to the average signal intensity. Background subtracted median intensities were calculated for each spot on the microarray and the data was normalized according to the total signal intensity, so that the average spot intensity was the same for each replicate slide (×3). The normalized data of each competitive array experiment was used to generate a list of the high-intensity sequences/spot (the highest to lowest intensity), i.e. spots which were above a threshold level of intensity (Supplementary Data, Figure Aa). High-intensity spots/sequences which occurred in all three replicates were carried forward for further data analysis, this procedure minimized the occurrence of any false positives* or negatives* (*False positives = spots which fluoresce highly on one array but not on all three arrays. Total = 6.5%; False negatives = spots which did not fluoresce on one array but fluoresced highly on two arrays. Total = 2%) in the overall data collection. The average intensity was calculated and the sequences were ranked accordingly. The list of sequences generated were condensed to include only the best binding sequences, these were spots that had intensity above 55% normalized fluorescence and at least 6 standard deviations away from the global mean intensity (Figure Ab, Supplementary Data). The final list contained a total of 50 high affinity-binding sequences for His$_6$-CspB.

**Isothermal titration calorimetry (ITC)**

ITC experiments were carried out as described previously (21) with minor modifications. Four oligonucleotides were used for the experiments (Figure 2B), both possibilities of the consensus-binding sequence (ITC1 and ITC2), a positive (ITC3(26)) and negative control (ITCcontrol). Each exothermic heat pulse (Figure 2A, upper panels) corresponds to an injection of 5 µl of each oligonucleotide (100 µM) into the cell containing 5 µM CspB at 28°C. Integrated heat data (Figure 2A, lower panels) constitutes a differential binding curve, which was fitted to a single-site binding model to give, the stoichiometry of binding (N), binding affinity ($K_d$) and enthalpy of binding ($\Delta H$) for each heptanucleotide.

**RESULTS AND DISCUSSION**

**Cy5 labelled SsoSSB binds ss DNA**

SsoSSB was covalently labelled with the mono-functional dye Cyanine 5 (Cy5; Amersham). Electrophoretic mobility shift assays (EMSA) were conducted to verify that SsoSSB retained its DNA-binding activity subsequent to labelling with Cy5. A single-stranded 25-mer (ONc: 5′-dATCTCTACTGATTGGCCAAGGTGCTG-3′), labelled at the 5′-end with γ-$^{32}$P, was used to compare the binding affinity of unlabelled and Cy5-labelled SsoSSB. Figure Cc (Supplementary Data) shows a gel-shift experiment performed with ONc in the presence of increasing amounts of unlabelled (lanes 1–4) or labelled (lanes 5–8) protein. The similarity in the EMSA for both unlabelled and Cy5-labelled protein suggests that labelling did not significantly affect the binding affinity of the protein. The double banding seen (Figure Cc, Supplementary Data) is a result of the gradual dissociation of labelled DNA from SsoSSB. The fact that previous SsoSSB-ss DNA-binding studies using ITC have shown that SsoSSB bound to ss DNA with a ratio of 5 nt/monomer (2) suggests that the double banding seen in the gel is probably a result of the EMSA technique used and the on/off-rates of SsoSSB-ss DNA complex formation.

**SsoSSB-Cy5 binds non-specifically to all ss DNA sequences**

A generic chip (Figure D, Supplementary Data) was constructed, which contained all possible 4096 ss hexadoxynucleotide sequences found in DNA and incorporated into a general construct, 5′-NH–A$_{10}$–N$_p$–X$_C$–3′ (X = G, C, A, or T and the stretch of 9 Ns is composed
of random bases). The binding of SsoSSB-Cy5 to the generic chip was analysed and all the spots on the array fluoresced with similar intensities, consistent with non-specific binding of SsoSSB-Cy5 (2).

**CspB and His₆-CspB have similar affinity for ssDNA**

A ss 25-mer (3) (ONc: 5'-dATCCCTACTGATTGGC CAAGGGCCTG-3'), labelled at the 5'-end with γ-³²P was used to compare the binding affinities and specificities of His₆-CspB and CspB. Figure 1A shows a gel-shift experiment performed for ONc in the presence of decreasing amounts of His₆-CspB (lanes 2–6) and non-His₆-tagged CspB (lane 7). Lanes 3, 4 and 5 show decreasing migration patterns for the His₆-CspB ss DNA complex as less protein molecules bind. This is most likely a result of deviation in the affinity of CspB for specific binding sites within ONc. The data show that the addition of the His-tag does not significantly affect the binding affinity or specificity of the protein. The effect of flanking DNA at the 3' end of the oligonucleotides on CspB binding, was also examined by EMSA using a series of oligonucleotides that were structurally consistent with the oligonucleotides found on the array; the only difference was that a varying number of bases were added.

**Figure 1.** (A) Electrophoretic mobility shift assay of His₆-CspB and its complex with ONc. 15–20 pmol of 5'-end γ-³²P-labelled ONc was mixed with 0, 500, 300, 100, 50 and 25 pmol (lanes 1–6) of His₆-Csp. Lane 7 contains 500 pmol of CspB (untagged) as a positive control. Protein-DNA complexes were separated on a 20% native polyacrylamide gel. (B) Electrophoretic mobility shift assay of SsoSSB, CspB and the complexes formed with ON1. 15–20 pmol of 5'-end γ-³²P-labelled ON1 was used at all times. Lanes; (1) 300 pmol SsoSSB; (2) 500 pmol CspB; (3) 300 pmol SsoSSB & 100 pmol CspB; (4) 300 pmol SsoSSB and 200 pmol CspB; (5) 300 pmol SsoSSB and 300 pmol CspB; (6) Unbound ON1 probe. Protein-DNA complexes were separated on a 10% native polyacrylamide gel. (C) Competitive microarray assay. Two individual microarray results for the competitive protein binding assay between His₆-CspB (specific binding protein) and SsoSSB (non-specific binding protein). The fluorescent signal is a result of His₆-CspB being detected on the surface using alexa 532-conjugated polyclonal antibody to His₆. Microarray images shown here were results from two different batches of arrays and proteins. Sub-array three from each competitive protein-binding experiments has been enlarged to highlight the repeated pattern seen on all arrays. The oligonucleotide, GCACCTT, appears as a bright spot on both arrays. The microarrays have been artificially coloured as a visual aid only. **Abbreviations.** SA = Sub-array (D) CspB consensus binding site. A 3-D bar chart showing the relative base distribution of each base within the proposed heptanucleotide-binding site. (E) Sequence logo (generated using the EnoLOGOS program), corresponding to the frequency of each base within the heptanucleotide-binding site. Both graphic visualizations show the resulting consensus sequence, 5'-GTCTTTG/T-3'.
to the 3' end. The results from these experiments show that flanking bases at the 3' end or a lack of them did not seem to affect the binding of CspB (T1 and Figure B, Supplementary Data).

**CspB is competitive with SsoSSB**

A competitive EMSA was used to show that His<sub>6</sub>-CspB binds more strongly than SsoSSB to the previously reported high affinity Y-box-binding motif (27) (ATTGG). A 25-mer, ON1 (5'-dA19-GATTGG-3'), which contains the Y-box-binding motif, was labelled at the 5'-end with γ<sup>32</sup>P. ON1 is similar in composition to the oligonucleotides found on the generic chip. Figure 1B shows the result of a gel-shift experiment performed with ON1 in the presence of varying amounts of His<sub>6</sub>-CspB and SsoSSB proteins. An intermediate band can be seen in lane 5, corresponding to the formation of the His<sub>6</sub>-CspB-SsoSSB-ON1 complex. This band occurs only when the SsoSSB:CspB proteins are in a molar ratio of approximately 1:1.

**The competitive binding assay can be transferred to microarray format**

The fluorescent signal from His<sub>6</sub>-CspB bound to the chip was analysed (Figure E, Supplementary Data). About 20% of the spots had signals greater than threshold level (which was set at 40% of the maximum fluorescent signal). To eliminate weakly bound CspB, an equimolar mixture (as determined by gel-shift, Figure 1B) of a competitor SsoSSB, and His<sub>6</sub>-CspB was incubated with an oligonucleotide chip and bound His<sub>6</sub>-CspB was detected (Figure 1C). High-intensity fluorescence spots were observed in repeated patterns on the arrays, which is indicative of selective binding affinity (25).

**High-affinity motifs identified by microarray analysis, are validated by electrophoretic mobility shift assay and isothermal calorimetry**

An EMSA was carried out to confirm the binding-site data generated from the oligonucleotide chip analysis. The high affinity-binding motif, GCACCTT, was chosen from the data (Figure 1C) to examine if the His<sub>6</sub>-CspB could compete successfully with the SsoSSB protein for this binding site. A 25-mer, ON-Microarray test (ON-Mt:5’-dAAAAAAAAGACTTTAAAAAA-3’), containing the high affinity-binding motif was labelled at the 5'-end with γ<sup>32</sup>P. Figure F (Supplementary Data) shows a gel-shift experiment performed with ON-Mt in the presence of varying amounts of His<sub>6</sub>-CspB and SsoSSB proteins. An intermediate band (His-Csp-SsoSSB-ONc complex) can be seen in lanes 3–11 for ON-MTest, indicating that the CspB protein competed with SsoSSB for binding to the (microarray determined) high-affinity-binding site, GCACCTT.

The highest intensity spots identified from the microarray analysis indicate that the strongest CspB-binding sites are pyrimidine rich (Figure Ga, Supplementary Data). The high incidence of thymine bases within the high-affinity 6-mer sequences agrees with previous reports that CspB has a preference for T-rich stretches of ss DNA (28). The presence of a stretch of 10 adenines in the linker region of each oligo (Figure Dd, Supplementary Data) is likely to lead to hairpin formation for T-rich sequences and may be expected to down weight the occurrence of poly T sequences. There is indeed a low intensity for TTTTTT sequences, which may in part be caused by the formation of such hairpins. Despite this effect, the averaging procedure still generates a T-rich consensus sequence. The standard motif alignment method (Genedoc) was used to align the resulting top fifty (as described in Materials and Methods) high-affinity CspB binding sequences (Figure Gb, Supplementary Data). A sequence alignment window 10 bases in length was used; only seven (coloured columns) out of those 10 positions are significantly (>40%) populated. Analysis of the relative distribution of each base within this proposed heptanucleotide-binding site gives a CspB consensus-binding sequence of 5’-GCTTTTG/T-3’ (Figure 1D and E).

Analysis of the microarray binding results indicates that CspB can accommodate the binding of a heptanucleotide with a strong binding preference for cytosine at position 3 and thymine at positions 2, 4 and 6. This is in agreement with another recent study (26), where the sequence-specific binding of heptapyrimidines to CspB was analysed by tryptophan fluorescence quenching experiments. Interestingly, the microarray results show that neither the Y-Box recognition motif (22), 5’-ATTGG-3’, nor its reverse complement 5’-CCAA-3’, bind strongly to CspB. The ATTGG sequence has recently been shown to bind with a low affinity for CspB at 15°C (K<sub>d</sub> = 5.3 μM (29)), which is similar to the results described here for CspB binding to sequences containing ATTGG at 4°C.

Both variants of the preferential binding sequences (ITC-1 and ITC-2) were analysed by ITC and gave binding constants in the low nanomolar range (Figure 2B). There is an order of magnitude difference in the K<sub>d</sub> for the oligonucleotides described here and the K<sub>d</sub> described previously (26), for similar/identical oligonucleotides. This is likely due to a difference in temperature, buffer conditions and the method used (tryptophan fluorescence quenching). The control sequence (TTCTTTT-ITC3), used here and in the previous study allows us to compare and scale the results from the two methods.

Thus, both the ITC and EMSA results confirm that the microarray assay does indeed select and identify tight binding sequences. This screening procedure could, therefore, be used as a general method for the rapid identification of high-affinity binding sites for SNABPs.

**CspB-cytosine model explains the preference for a cytosine at position 3**

The X-ray structure for CspB has been reported previously (24). Figure 2C shows the positively charged face of CspB, highlighting amino acid residues known to be involved in DNA binding (3). Molecular modelling of CspB with the consensus oligonucleotide ITC1, using the
Figure 2. (A) Interaction of CspB with ss DNA templates as monitored by ITC. Titration data for 27 injections of 5µl of each oligonucleotide (100µM) into a cell containing 5µM CspB. Oligonucleotides used for ITC are listed below. All data have been corrected for heats of dilution. (B) Thermodynamic parameters for CspB-ss DNA interactions. Non-detectable = $K_d \leq 5 \mu M$. (C) A molecular model of a potential cytosine-binding pocket on CspB. The electrostatic surface potential of CspB and the backbone trace of the bound consensus heptanucleotide and the hydrogen bonds (depicted as dotted lines) between CspB and a cytosine nucleobase in the proposed binding pocket.
programme WITNOTP, identified a pocket in the centre of the DNA-binding face, which provides an ideal shape and hydrogen-bonding complementarity for binding cytosine. Three hydrogen bonds are formed between the docked cytosine base and amino acid side chains Ser31, His29 and the backbone of Phe27 (Figure 2C), providing specificity for cytosine over the other bases.

In the recently published structure of CspB in complex with hexahymidine (26), the ligand binds to two protein molecules. The nucleobase of T2, T3 and T4 make contact with one protein molecule, T5 bridges between protein molecules and T6 binds to the next protein molecule. The nucleobase of T2, T3 and T4 make contact with one protein molecule, T5 bridges between protein molecules and T6 binds to the next protein molecule. The contacts made by hexahymidine provide the necessary scaffold for the complex to crystallize but the sequence fails to bind across the face of the CspB protein, as this sequence lacks the key cytosine nucleobase at position 3 (26), which, as we have shown here, is required for optimum ss DNA docking (Figure 2).

SNAPs have been reported to activate transcription by binding to a specific recognition sequence upstream (30) or within (31) a promoter, resulting in activation or repression of transcription. In the present study, CspB of B. subtilis which is capable of binding single-stranded nucleic acids (21) and affects expression of over 100 genes under cold shock conditions (32), was used as the test protein. Database analysis of the B. subtilis genome reveals that 89 copies of the consensus-binding sequence (5′-GTCTTTT-3′) exist within potential SNAP promoter regions (100 bases upstream of the ATG start for all genes), of which only 24 have an assigned function. The use of an unbiased genomic assay to identify optimal-binding sequences for SNAPs in vitro, may provide insight into their role in regulating cellular functions. The full implications of these sequences on gene expression and binding of CspB remain to be determined.

SUPPLEMENTARY DATA
Supplementary Data is available at NAR Online.

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