

repeated many times, and the ensemble of coordinates is used to reconstruct an image with superior resolution [2, 3].

RNA polymerases have been localized in bacteria using conventional approaches [4]. In our study, we focus on the spatial organization of bacterial transcription sites in *E. coli* at the molecular scale. To reach that goal, we apply high-resolution fluorescence methods, and we will present a refined understanding of structure and function in the bacterial transcription machinery.

#### References:

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#### 371-Pos

##### **A Microfluidics-Based Platform For Identification and Detailed Characterization of Transcription Factor Binding Sites**

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Transcriptional regulation of gene expression is mediated by transcription factors that bind DNA sequence-specifically within gene promoters to activate or repress transcription. However, biochemical data linking transcription factors to their consensus binding sites has traditionally been difficult to obtain, complicating reconstruction of cellular pathways contributing to a transcriptional response.

We have developed a versatile and sensitive microfluidics-based technique for *de novo* identification and subsequent detailed characterization of transcription factor consensus motifs and binding energy landscapes. Our technique offers several advantages over current methods. First, our technique mechanically traps all complexes at equilibrium prior to measurement, allowing detection of weak or transient interactions and providing direct, quantitative measurements of reaction parameters. Second, our technique requires extremely small amounts of reagents, permitting protein production via cell-free transcription/translation of PCR-generated templates and eliminating laborious and time-consuming cloning steps. Finally, our technique allows high-throughput screening of transcription factor binding to all possible DNA 8mers in a single experiment.

To evaluate the performance of our new technique, we probed DNA binding patterns for 30 yeast transcription factors from various families and used a statistical-mechanical model of transcription factor binding to determine preferred consensus motifs. In all cases, the core consensus obtained agreed with previous literature results, validating the utility of our technique for *de novo* identification of transcription factor binding sites. This quantitative data set provides critical information that can be used to revise and refine current models of transcription factor binding interactions.

#### 372-Pos

##### **Model of Transcriptional Activation By MarA in Escherichia Coli**

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The AraC family transcription factor MarA activates ~40 genes (the *marA/soxS/rob* regulon) of the *Escherichia coli* chromosome resulting in different levels of resistance to a wide array of antibiotics and to superoxides. Activation of *marA/soxS/rob* regulon promoters occurs in a well-defined order with respect to the level of MarA; however, the order of activation does not parallel the strength of MarA binding to promoter sequences. To understand this lack of correspondence, we developed a computational model of transcriptional activation in which a transcription factor either increases or decreases RNA polymerase binding, and either accelerates or retards post-binding events associated with transcription initiation. We used the model to analyze data characterizing MarA regulation of promoter activity. The model clearly explains the lack of correspondence between the order of activation and the MarA-DNA affinity, and indicates that the order of activation can only be predicted using information about the strength of the full MarA-polymerase-DNA interaction. The analysis further suggests that MarA can activate without increasing polymerase binding and that activation can even involve a *decrease* in polymerase binding,

which is opposite to the textbook model of activation by recruitment. These findings are consistent with published chromatin immunoprecipitation assays of interactions between polymerase and the *E. coli* chromosome. We find that activation involving decreased polymerase binding yields lower latency in gene regulation and therefore might confer a competitive advantage to cells. Our model yields insight into requirements for predicting the order of activation of a regulon and enables us to suggest that activation might involve a decrease in polymerase binding, which we expect to be an important theme of gene regulation in *E. coli* and beyond.

#### 373-Pos

##### **Transcription Factor Switching Dynamics Regulates Gene Activation**

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Molecular mechanisms controlling the time diffusing molecules take to activate specific target proteins are pivotal for cellular response and signaling. We study the activation dynamics when diffusing ligands switch between various states induced by chemical interactions or conformational changes, while target activation is possible only in a specific state. We find that the activation time is very sensitive to changes of the switching rates, which is a way to modulate cellular signaling. Interestingly, target activation can be fast although the ligand spends most of the time in a non-activating state, which is relevant if activation occurs in a state where the ligand is also prone to degradation. Using a modeling approach and data from FRAP and single particle tracking experiments, we study the switching dynamics of the positive transcription elongation factor b (P-TEFB) inside the nucleus and unravel a novel mechanism of gene regulation. P-TEFB is necessary for the activation of many genes and its motion is controlled by various chemical interactions that alter the state of P-TEFB and its affinity for the DNA.

#### 374-Pos

##### **A Genome-Wide Analysis of Poised Promoters in Bacteria**

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As the first and usually rate-limiting step of transcription initiation, bacterial RNA polymerase binds to double stranded DNA (the closed complex formation) and subsequently opens the two strands of DNA (the open complex formation). Poised promoters in bacteria are sequences where RNAP binds with high binding affinity, but which do not have detectable levels of transcription initiation due to too slow transition from closed to open complex. Existence of a considerable number of poised promoters in genome has been often hypothesized, but poised promoters have not been systematically studied, since a large scale analysis of promoter kinetics is not experimentally feasible. To computationally address promoter poising on a genome-wide scale we use a recently developed biophysical model of transcription initiation [1]. We show that promoter poising is significantly reduced by i) Existence of -35 box interactions ii) Binding specificities of (physically independent) RNAP domains that interact with -10 box single-stranded and double-stranded DNA. We show that the later (dominant) effect is not due to generic properties of protein-DNA interactions, and argue that RNAP is designed to reduce promoter poising in genome. However, despite this reduction, we obtain that the number of poised promoters is still significant, and corresponds to ~30% of strongly bound sequences in bacteria [2]. This number roughly matches with lower bound of reported false positives in RNAP ChIP-chip experiments, which suggests that poised promoters are a major contributor to false positives in searches of bacterial promoters.

[1] M Djordjevic and R Bundschuh, *Biophysical Journal* 94: 4233 (2008). [2] M Djordjevic, submitted (2009).

## **Protein-Nucleic Acid Interactions I**

#### 375-Pos

##### **Dynamic Investigation of DNA Bending and Wrapping By Type II Topoisomerases**

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Type II topoisomerases catalyze DNA decatenation and unwinding which is crucial for cell division, and therefore type II topoisomerases are some of the main targets of anti-cancer drugs. A recent crystal structure shows that, during the catalytic cycle, a yeast type II topoisomerase can bend a 34 base pair DNA segment by up to 150 degrees. Bacterial gyrase, another type II topoisomerase, can wrap an approximately 100 bp DNA segment into a tight 180 degree turn. Bending a stiff polymer like DNA requires considerable energy and could

represent the rate limiting step in the catalytic (topological) cycle. By substituting diaminopurine (DAP) deoxyribonucleotides for dATP in PCR reactions, stiffer DNA fragments have been produced and used as substrates for topoisomerase II-mediated relaxation of plectonemes introduced in single molecules using magnetic tweezers. The overall rate of relaxation of plectonemes by recombinant human topoisomerase II alpha decreased on the stiffer DNA. In addition the ability of recombinant *E. coli* gyrase to wrap DNA also decreased for DAP-substituted DNA in which every base pair has three hydrogen bonds. These dynamic measurements of DNA bending and wrapping by type II topoisomerases are consistent with the hypothesis that DNA flexibility affects the rate determining step for type II topoisomerase activity.

### 376-Pos

#### Characterization of HIV-1 Reverse Transcriptase 3TC Specificity By Conformationally Sensitive Fluorescence Reveals New Insights Into the Kinetic Basis of Inhibitor Discrimination

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HIV-1 Reverse Transcriptase (HIV-RT) is the target of nine Nucleoside Reverse Transcriptase Inhibitors (NRTI's) currently approved by the FDA. Polymerase specificity is best quantified by analysis of the concentration dependence of the rate using single turnover rapid Quench-Flow methods which provide a rate of polymerization ( $k_{pol}$ ) and an apparent dissociation constant ( $K_d$ ) such that  $k_{pol}/K_d = k_{cat}/K_m$ . Analysis of nucleoside analog RT inhibitors (NRTIs) has led to the surprising conclusion that most appear to bind more tightly than normal nucleotides. For example, 3TC-triphosphate binds 10-fold tighter than the correct nucleotide (dCTP). Using a conformationally sensitive fluorophore attached to the fingers domain of the enzyme, we show that nucleotide binding is a two step process involving weak nucleotide ground state binding, followed by a conformational change from an "open" to "closed" state. These steps together define the true  $K_d$  for nucleotide binding at equilibrium. Examining the kinetics of 3TC incorporation, we show that contrary to previously reported findings, the dCTP analog binds 8-fold more weakly to the enzyme than the correct nucleotide. Further, we show that the enzyme's conformational change to the "closed" state is capable of sensing dCTP versus 3TC and results in an increased or decreased binding affinity, respectively. The result is a specificity constant ( $k_{cat}/K_m$ ) of  $9.7\mu M^{-1}s^{-1}$  for dCTP and  $0.7\mu M^{-1}s^{-1}$  for 3TC. The specificity constant for dCTP is determined solely by the rate of nucleotide binding ( $k_{cat}/K_m = K_1k_2$  in the two-step sequence), whereas the slower chemical reaction ( $k_3$ ) for 3TC incorporation allows the binding and isomerization to reach equilibrium so that  $k_{cat}/K_m = k_{pol}/K_1K_2$ . This work provides mechanistic basis for discrimination of 3TC, and corrects how  $K_m$ ,  $K_d$ , and  $K_{d,app}$  must be assigned for NRTIs.

### 377-Pos

#### Coliphage 186 Genetic Switch: A Single Molecule Study

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It is increasingly clear that in most cases genes are regulated by wrapping or looping of DNA on large, cooperatively assembled protein complexes. In most eukaryotic organisms, 150 bp of DNA are wrapped around histone octamers (nucleosomes). Furthermore, interaction between proteins bound at distant sites on the DNA may cause looping out of the intervening DNA and have regulatory significance. The mechanism by which these DNA-protein nanostructures are formed is not clear. The interaction between the bacteriophage repressor 186CI (a disc-shaped heptamer) and its DNA is an ideal model system to study DNA wrapping and looping and to reveal fundamental principles of long-range interactions and regulation by nucleoprotein complexes. Here we report on AFM work aimed at elucidating the 186CI-DNA interaction. We analyzed the structure of the protein DNA complexes revealed by the AFM images and we propose a mechanism that leads to repression of the lytic genes in 186 and regulation of the repressor expression via DNA wrapping around a protein heptamer and protein repositioning along the DNA.

### 378-Pos

#### Using Real-Time, Single-Molecule Experiments To Monitor RecA-Mediated Pairing and Strand Exchange Reactions in Various Nucleotide States Hsiu-Fang Fan<sup>1</sup>, Michael M. Cox<sup>2</sup>, Hung-Wen Li<sup>1</sup>.

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RecA recombinases play a central role in homologous recombination pathway. Once they assemble on single-stranded (ss) DNA, the RecA/ssDNA filament mediates the pairing of homologous DNA sequence and strand exchange

processes. We used tethered particle motion (TPM) experiments to investigate the details of *E. coli* RecA-mediated pairing and strand exchange steps at the single molecule level. TPM experiments measure the DNA tether length change according to the bead Brownian motion. In the "incoming bead" experiment, ssDNA molecules bound with sub-micron sized polystyrene beads were coated with RecA and then paired with homologous duplex DNA tethered on surface. Therefore, the appearance of the bead tether and its Brownian motion amplitude permit the direct observation of RecA-mediated pairing and strand exchange processes in real-time. In the "leaving bead" experiment, surface-bound hybrid duplex DNA molecules were tethered with polystyrene bead, and then reacted with RecA-coated complementary ssDNA. Disappearance of the tethered beads indicates the completion of strand exchange. It was found that pairing and strand exchange steps are more efficient under low pH=6.5 condition in which the strand exchange efficiency of  $0.17 \pm 0.02$ , is higher than that in pH=7.5 ( $0.11 \pm 0.05$ ). The pairing process occurs successfully in both ATP and its non-hydrolyzable analog, ATP $\gamma$ S state, but not in ADP state where the three-stranded intermediate are found to be unstable (half-life time=0.7s). Surprisingly, the strand exchange efficiency under ATP and ATP $\gamma$ S states are similar ( $0.19 \pm 0.03$  and  $0.18 \pm 0.01$  for ATP and ATP $\gamma$ S respectively), suggesting ATP hydrolysis of RecA is not necessary to complete strand exchange step in our experiment. These single-molecule experiments provide new mechanistic details on the RecA-mediated processes.

### 379-Pos

#### Intersubunit Regulation Between Nuclease and Helicase Domains of Recbcd Enzyme

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The *Escherichia coli* RecBCD helicase/nuclease initiates homologous recombinational repair of damaged blunt-end duplex DNA molecules. RecBCD, a multifunctional enzyme complex, contains two DNA motors as well as a nuclease domain to process duplex DNA and generate single-stranded DNA molecules. We used single-molecule tethered particle motion (TPM) experiments to investigate the regulation mechanism between the nuclease domain and two helicase domains of RecBCD enzyme using calcium ions, which specifically inhibit nuclease activity. In the absence of calcium ions, RecBCD translocation rate is found to slow down after recognizing chi sequence. However, in the presence of calcium ions, the rate change in individual RecBCD translocation is abolished, returning similar averaged translocation rate before ( $71 \pm 20$  bp/s) and post ( $81 \pm 36$  bp/s) chi-sequence, under  $30\mu M$  ATP. Furthermore, large portion of individual RecBCD unwinding time courses (13 out of 32) revealed repetitive forward and backward translocation along individual DNA molecules. Compared with the experiments carried out without calcium ions, the processivity of RecBCD also decreases when the nuclease domain is inhibited. About 50 percent of translocating tethers (17 out of 32) stalled within 1.5 Kb DNA used in the presence of calcium ions. Together, these observations suggest that the nuclease domain, located in the RecB subunit, plays regulatory roles not only in RecBCD translocation properties but also in chi-regulated intersubunit interaction in this complex machine of the RecBCD enzyme.

### 380-Pos

#### A Structural Model For RNA Remodeling By a Dimeric Dead Box Helicase Markus G. Rudolph<sup>1</sup>, Dagmar Klostermeier<sup>2</sup>.

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DEAD box helicases couple ATP hydrolysis to RNA structural rearrangements. *T. thermophilus* Hera (heat resistant RNA-dependent ATPase) consists of a helicase core and a C-terminal extension. In single molecule FRET experiments we identified fragments of the 23S rRNA comprising hairpin 92 and RNase P RNA as substrates for Hera. RNA binding requires the C-terminal extension. Both substrates switch the helicase core to the closed conformation and stimulate the intrinsic ATPase activity of Hera. ATP-dependent unwinding of a short helix adjacent to hairpin 92 of 23S rRNA suggests a specific role for Hera in ribosome assembly, in analogy to the *E. coli* and *B. subtilis* helicases DbpA and YxiN. In addition, the specificity of Hera for RNase P RNA may be required for RNase P RNA folding or RNase P assembly.

Hera forms a stable dimer in solution, setting it apart from other helicases. Crystal structures show that the C-terminal extension is bipartite, forming a highly flexible dimerization motif with a novel fold and an additional RNA-binding module that adopts the fold of a degenerated RNA recognition motif (RRM). Comparison with RRM/RNA complexes suggests an RNA binding mode similar to that of the spliceosomal protein U1A. The structure-based model for the complete Hera dimer bound to RNA reveals a likely binding