

base-complementarity with a region of TER, and extends the DNA using TERT's reverse transcriptase activity. Structural, functional, and kinetic studies of telomerase have been limited by its poor expression in vivo and in heterologous systems, as well as its structural heterogeneity. For this reason, this is an ideal system to study at the single molecule level, as single molecule assays do not require large amounts of material and can parse out different sub-populations of molecules. Our lab has developed a single-molecule Förster resonance energy transfer (smFRET) assay in order to monitor structural changes in the telomerase enzyme during its activity. We have used this assay in combination with several telomerase mutants to conduct structural and mechanistic studies of telomerase function. Our results demonstrate that a conserved telomerase N-terminal domain (TEN domain) stabilizes duplex formation between telomerase RNA and its DNA substrate. This TEN domain functions by altering the equilibrium between the duplex (active) state and a previously unknown alternative (inactive) binding conformation. Furthermore, we demonstrate that mutants that stabilize the alternative conformation have severe defects in telomerase activity. The discovery of this inactive alternative state suggests that stabilizing this conformation could be a useful target of telomerase inhibitors for future cancer treatment.

2174-Plat

Targeting and Degradation of Viral DNA by the CRISPR-Cas System of *Escherichia Coli*

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Bacteria and archaea maintain a history of viral infections by integrating small fragments of foreign DNA into specialized genomic loci called clustered regularly interspaced short palindromic repeats (CRISPRs). Subsequent infections trigger an adaptive immune response, which relies on both CRISPR RNAs (crRNAs) and CRISPR-associated (Cas) proteins to identify and destroy invading DNA. In *E. coli*, the crRNA/Cas ribonucleoprotein surveillance complex is referred to as Cascade. Cascade recognizes foreign DNA targets via crRNA-DNA base pairing and subsequently recruits a trans-acting nuclease/helicase, Cas3, for degradation of the targeted DNA. To fully dissect the mechanistic features of target recognition and degradation by Cascade/Cas3, we have used a combination of single molecule and bulk methods. We reveal crucial features in DNA that Cascade senses to locate and recognize complementary target sequences within the larger context of genomic DNA. Further, we uncover the key elements necessary for Cascade-dependent recruitment of Cas3. Finally, by directly visualizing Cas3 as it unwinds and cleaves viral DNA, we characterize the degradation machinery in exceptional detail, providing fundamental insights into the mechanism of CRISPR-based immunity. Importantly, by comparing these results with our recent findings for another CRISPR-Cas system, Cas9 in *S. pyogenes*, we can begin to understand the evolution of DNA targeting machineries in related RNA-based adaptive immune systems.

2175-Plat

Selective Acetylation Reveals Distinct Roles of Histones H3 and H4 in Nucleosome Dynamics - a FRET Study

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Histone tails and their posttranslational modifications play a crucial role in controlling genetic activity through alterations of nucleosome structure. Whether or not histone tails regulate DNA accessibility independently of each other or in a concerted fashion is currently under debate. Here we studied the structure-defining properties of selective histone acetylation and point mutations in the H4 tail in a combined bulk FRET - single molecule FRET assay. Nucleosome unwrapping was monitored by FRET between the linker ends of the DNA, while FRET experiments at an internal DNA site in the H2A/H2B binding region reported on nucleosome disassembly.

By analysis of nucleosome unwrapping, structural heterogeneity during salt-induced disassembly and dimer exchange between nucleosomes we show that histones H3 and H4 assume significantly different roles in controlling nucleosome architecture. H4-acetylation opposes destabilization by H3-acetylation and reduces linker DNA unwrapping and dimer exchange at higher ionic strength, whereas its influence on nucleosome structure at physiological salt is minute. We found no increase in unwrapping when H3 and H4 were

acetylated simultaneously, which challenges the idea of cooperativeness between tails that was observed for truncated H3 and H4. Our data suggest that the effect of lysine acetylation is not cumulative in nature but shows strong histone specificity. The specific role of the H4 tail was finally probed by comparing the effect of point mutations or acetylation of selective lysine residues at positions 5,8,12 and 16.

Regardless of the state of acetylation nucleosomes disassemble via an intermediate state, which is suppressed at higher nucleosome concentration, confirming our proposed model of step-wise disassembly.

Platform: Calcium Fluxes, Sparks, and Waves

2176-Plat

Decomposition of a Calcium Spark in Cardiac Myocytes

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Ca²⁺ sparks are the elementary SR Ca²⁺ release events in heart muscle. Spontaneous and triggered Ca²⁺ sparks are usually centered on a junctional sarcoplasmic reticulum (jSR) across a 15 nm "subspace" gap from a transverse tubule (TT) at the Z-disk. The jSR-TT junction contains ryanodine receptor (RyR2) clusters of a variable number of RyR2s and diverse organizations. To determine if Ca²⁺ sparks may reveal a substructure that reflects the jSR-RyR2 organizational variability, we have examined rabbit ventricular myocytes and imaged [Ca²⁺] in living myocytes using a confocal microscope. The cytosol was loaded with the [Ca²⁺]_i indicator rhod-2 and the SR loaded with the low affinity [Ca²⁺]_{SR} indicator fluo-5N. This method has allowed the detection of Ca²⁺ blinks (the local SR Ca²⁺ depletion during a Ca²⁺ spark) and Ca²⁺ sparks simultaneously. Furthermore, this method makes it possible to image the recently described set of small, sub-spark events, the quirky SR Ca²⁺ release or QCR. The spatial profiles of Ca²⁺ sparks and Ca²⁺ blinks has thus revealed a sub-structure within a single Ca²⁺ spark. The images suggest that Ca²⁺ sparks could be composed of a central Ca²⁺ release site with surrounding small release site elements (RS-elements). These RS-elements may be asymmetrically distributed around the central site. That high concentrations of EGTA (30 μM) prevented the activation of the RS-elements suggests that these RS-elements or QCRs are activated by Ca²⁺-induced Ca²⁺-release (CICR) triggered from the central site. How the underlying structures may change in diverse diseases may be important in broadening our understanding of both arrhythmogenesis and heart failure, diseases widely associated with altered Ca²⁺ signaling, [Ca²⁺]_i stability and altered cellular ultrastructure.

2177-Plat

Examination of Single Channel RyR Behavior from Long-Lasting Ca²⁺ Sparks

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Physiology & Pharmacology, University of Bristol, Bristol, United Kingdom. Calcium sparks are crucial to cardiac excitation-contraction coupling and are due to calcium-induced calcium release via ryanodine receptors (RyRs) located on the sarcoplasmic reticulum (SR). A calcium spark results from the concerted opening of a cluster of RyRs, which generally closes by a robust mechanism as shown by stereotypic spark duration (~40 ms). However, much longer calcium sparks (up to seconds) can be seen in cells isolated from an animal model of heart failure (1) and when RyRs have been pharmacologically inhibited (2). The mechanism underlying such long-lasting events is unclear. It has been suggested that a lack of local SR calcium depletion (from a combination of decreased release flux and increased SR calcium buffering and/or refilling) is important (2), however it has also been suggested that the activity of RyR sub-clusters (located 1 ms ≈ 1 μm apart) may also sustain release (1). We examined calcium sparks in cells treated with tetracaine to partially inhibit RyRs. Spark-like and long-lasting events occurred at the same location in a line-scan image, suggesting a single release site is able to exhibit both types of behavior. Analysis of long-lasting release events (mean duration = 315 ms) yielded results consistent with a mechanism that involves a reduction in RyR availability and there was no detectable change in the location of the origin of release. These conclusions are supported by a computer model that was able to reproduce long-lasting events when the number of stochastically gating RyRs available was reduced and SR calcium was increased. Large alterations in RyR cluster geometry were not required, but the geometric organization of RyRs in the cluster plays an important role.

References

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