CRISPR-Cas systems allow bacteria to memorize prior infections as a means to combat the same invader if it attempts another attack in the future. While the underlying mechanisms of this bacterial immunity have been intensely studied over the past decade, little attention has been paid to CRISPR defense at the single-cell level. In their recent work, Brouns and colleagues (McKenzie et al., 2022) track memory acquisition and defense in individual cells and find a wide range of temporal dynamics that shape how a cell population experiences and combats an active infection.

Mol Syst Biol. (2022) 18: e11011
See also: RE McKenzie et al (April 2022)

CRISPR-Cas systems, well-known as the source of widely used genome editing tools, naturally protect bacteria and archaea from foreign invaders such as bacteriophages or mobile plasmids (Hille et al., 2018). Protection starts with each system storing small fragments of DNA derived from the invader within the genome of the host. The stored fragments, called spacers, are then used as templates to transcribe CRISPR RNAs, which direct cleavage of the invader DNA. This mode acts much faster than naive acquisition (Fineran et al., 2014). Naive acquisition usually incorporates spacers derived from stalled replication forks, but this mode tends to be extremely slow. Primed acquisition is triggered when the CRISPR machinery encounters a spacer target that has accumulated mutations. While these mutations prevent a full-fledged attack by the CRISPR machinery, the mutated spacer target can drive the acquisition of an adjacent fragment of invader DNA. This mode acts much faster than naive acquisition (Fineran et al., 2014).

The authors set out to measure the timing of acquisition and defense at the single-cell level in a growing population. Using a fluorescent reporter plasmid encoding a perfect or mutated spacer target in a bacterial strain with inducible expression of the CRISPR-Cas system, the authors were able to follow the loss of the plasmid over time via time-lapse microscopy (Fig 1). By tagging one component of the CRISPR machinery with another fluorescent protein, the authors could also evaluate how plasmid loss correlated with single-cell abundance of the machinery. In addition, this approach allowed concomitant assessment of other factors, such as a cell’s doubling time and the fate of progeny with a shared parent (sister cells) or grandparent (cousin cells). Finally, by incorporating stochastic modeling, the authors interrogated mechanistic features difficult to probe through experimental means.

This set of approaches led to intriguing insights that help inform how CRISPR-based immunity plays out across a population. For one, clearance of the plasmid with the perfect spacer target was rapid (1–3 cell doublings) and relatively uniform across the population, showing that direct interference can offer robust protection to an entire cell population. In contrast, clearance of the plasmid with the mutated spacer target was extremely variable, spanning 2–30 cell doublings. This variability was traced to the process of primed acquisition and the ranging time scales in which a new spacer could be acquired. Because direct interference commences shortly after a new spacer is acquired, the plasmid was much more likely to be cleared at the same time by sister cells than by cousin cells or any other cell in the population. The authors found that the cellular features they tracked had different effects on plasmid clearance and primed acquisition. For example, high levels of the CRISPR machinery correlated with faster clearance but not primed acquisition. Separately, rapid cellular growth was associated with faster clearance but also slower acquisition. In addition, modeling indicated that variability in the levels of the CRISPR machinery could accelerate primed acquisition across the population due to a highly non-linear relationship between abundance and acquisition. These insights paint a complex picture of the factors influencing CRISPR-based defense, leading to a hyper-variable response if defense occurs through primed acquisition. Such a response would leave most of the cell population susceptible...
to invader attack, leading to few survivors. The upside is that these survivors carry robust immunity against the invader if a second attack occurs again in the future.

Notwithstanding the authors’ advances, there are some limitations when extrapolating these findings to natural protection by CRISPR-Cas systems. For instance, the use of an inducible system to express the CRISPR machinery contrasts with natural systems that are constitutively expressed under infection conditions. The use of a plasmid as a target makes it difficult to extrapolate the findings to infection by bacteriophages that rapidly propagate and lyse the cell. Finally, the CRISPR-Cas system in *E. coli* has remained an oddity because it is insufficiently expressed under any natural growth conditions to enact immune defense (Pul et al., 2010; Westra et al., 2010). While these limitations are justifiable given the experimental setup and the benefits of using a well-characterized system, the authors’ findings raise important questions that will drive further research into CRISPR-based defense. What are the single-cell dynamics during an active infection by lytic bacteriophages often targeted by CRISPR-Cas systems? How do the dynamics and underlying mechanisms change across the rich diversity of CRISPR-Cas systems? And how do these dynamics influence the evolution of CRISPR-Cas systems and invaders, out to circumvent the defense system? By bringing single-cell techniques into the realm of CRISPR biology, we can expect to learn much more about the functions of these ingenious immune systems and possibly translate these insights into new and improved CRISPR technologies.

**References**

Fineran PC, Gerritzen MJH, Suárez-Diez M, Künne T, Boekhorst J, van Hijum SAFT, Staals RHJ, Brouns SJJ (2014) Degenerate target sites mediate rapid primed CRISPR adaptation. *Proc Natl Acad Sci USA* 111: E1629 – E1638

Hille F, Richter H, Wong SP, Bratovic M, Ressel S, Charpentier E (2018) The biology of CRISPR-Cas: backward and forward. *Cell* 172: 1239 – 1259

Levy A, Goren MG, Yosef I, Auster O, Manor M, Amitai G, Edgar R, Qimron U, Sorek R (2015) CRISPR adaptation biases explain preference for acquisition of foreign DNA. *Nature* 520: 505 – 510

McKenzie RE, Keizer E, Vink J, van Lopik J, Büke F, Kalkman V, Fleck C, Tans S, Brouns S (2022) Single cell variability of CRISPR-Cas interference and adaptation. *Mol Syst Biol* 18: e10680 https://doi.org/10.15252/msb.202110680

Pul U, Wurm R, Arslan Z, Geissen R, Hofmann N, Wagner R (2010) Identification and characterization of *E. coli* CRISPR-cas promoters and their silencing by H-NS. *Mol Microbiol* 75: 1495 – 1512

Westra ER, Pul U, Heidrich N, Jore MM, Lundgren M, Stratmann T, Wurm R, Raine A, Mescher M, Van Heereveld L et al (2010) H-NS-mediated repression of CRISPR-based immunity in *Escherichia coli* K12 can be relieved by the transcription activator LeuO. *Mol Microbiol* 77: 1380 – 1393

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