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Antibiotic Resistance: Can We Ever Win?

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Part I – Measuring Resistance

Katelyn was excited to start her summer job in her microbiology professor's research laboratory. She had enjoyed Dr. Johnson's class, and when she saw the flyer recruiting undergraduate lab assistants for the summer, she had jumped at the opportunity. She was looking forward to making new discoveries in the lab.

On her first day, she was supposed to meet with Dr. Johnson to talk about what she would be doing. She knew the lab focused on antibiotic resistance in Staphylococcus aureus, especially MRSA (methicillin-resistant S. aureus).

She still remembered the scare her family had last year when her little brother, Jimmy, got so sick. He'd been playing in the neighborhood playground and cut his lip when he fell off the jungle gym. Of course he always had cuts and scrapes—he was a five-year-old boy! This time though his lip swelled up and he developed a fever. When her mother took him to the doctor, the pediatrician said the cut was infected and had prescribed cephalothin, an antibiotic related to penicillin, and recommended flushing the cut regularly to help clear up the infection.

Dr. James Johnson

Two days later, Jimmy was in the hospital with a fever of 103°F, coughing up blood and having trouble breathing. The emergency room doctors told the family that Jimmy had developed pneumonia. They started him on IV antibiotics, including ceftriaxone and nafcillin, both also relatives of penicillin.

It was lucky for Jimmy that one of the doctors decided to check for MRSA, because that's what it was! MRSA is resistant to most of the penicillin derivatives. Most cases of MRSA are hospital-acquired from patients who are already susceptible to infection, but the ER doctor explained that community-acquired MRSA was becoming more common. The doctor then switched the treatment to vancomycin, a completely different kind of antibiotic, and Jimmy got better quickly after that.

Katelyn had dropped Jimmy off at swimming lessons just before coming to work at the lab. As she waited in the hallway for Dr. Johnson, she hoped that she would be at least a small part of helping other people like Jimmy deal with these scary resistant microbes. She was surprised when the professor burst out of the lab, almost running into her.

"Hi Katelyn, I'm really sorry but I have to run to a meeting right now—they sprung it on me last minute. There are a bunch of plates in the incubator right now that need their zones of inhibition measured. I'll be back in a few hours," Dr. Johnson said as he rushed down the hallway with a stack of folders.

Katelyn dug out her old lab notebook to look up what she was supposed to do. She found the lab where she and her fellow students had examined the antimicrobial properties of antibiotics using the Kirby-Bauer disk diffusion technique. Looking at the plates Dr. Johnson had told her about, she saw they had all been "lawned," or completely coated with microbes to make a thick hazy layer over the agar surface. She could also see paper disks with letters on them, and some of the disks had clear zones around them where the microbe had been inhibited (Fig. 1). Her notebook explained how to measure the zone of inhibition around the disks (Fig. 2).

"Antibiotic Resistance" by Maureen Leonard

Interested in studying microbial

antibiotic resistance?

We're looking for undergraduate lab assistants!

Johnson@fictionaluniv.edu

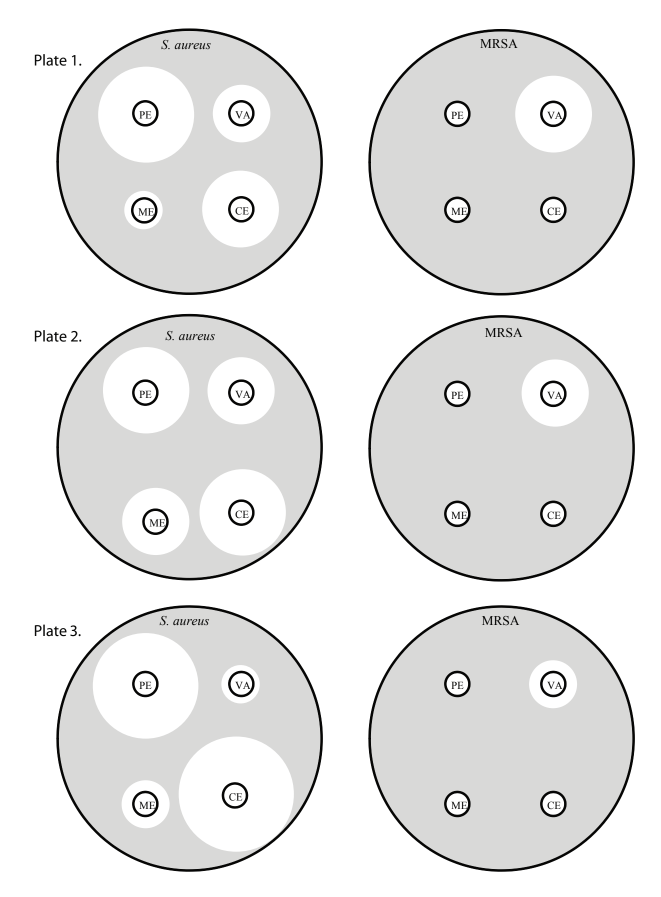


Figure 1. Agar plates of S. aureus or MRSA lawns with antibiotic disks placed on them.

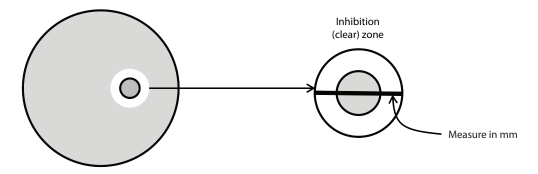


Figure 2. Katelyn's diagram of how to measure a zone of inhibition from her microbiology lab notebook.

Exercise1

Measure the zones of inhibition for each antibiotic on the plates shown in Figure 1 and note the measurements in the spaces in Table 1 below. (*Note:* The Kirby-Bauer method is standardized so that no zone of inhibition is scored as a 0, and all others include the disk as part of the zone.)

Plat	e	S. aureus	MRSA
1	PE		
	ME		
	CE		
	VA		
2	PE		
	ME		
	CE		
	VA		
3	PE		
	ME		
	CE		
	VA		

Key: PE = penicillin, ME = methicillin, CE = cephalothin, and VA = vancomycin

An average, or mean (\bar{x}) , is a measure of central tendency in the data, or what value occurs in the middle of the data set. The mean is calculated by adding up all the values for a given set of data, then dividing by the sample size (n).

Average

$$\overline{x} = \frac{\sum_{i=1}^{n} x_i}{n}$$

Standard deviation measures the spread of the data—as in how variable the data set is. The standard deviation (*s*) is calculated by the following:

Standard deviation
$$s = \sqrt{\frac{\sum (x - \overline{x})^2}{n - 1}}$$

Standard error measures the difference between the sample you have taken and the whole population of values. The standard error (SE) is calculated as follows:

Standard error $SE = \frac{s}{\sqrt{n}}$

Exercise 2

In Table 2 below calculate and record the averages and standard errors for each antibiotic in *S. aureus* and MRSA.

	S. au	reus	MRSA	
	Average	SE	Average	SE
PE				
ME				
CE				
VA				

Exercise 3

Now, redraw Tables 1 and 2 into a single, more organized table. Be sure to label the table appropriately.

Exercise 4

Graph the results from Table 2. Be sure to label the figure and the axes correctly.

Questions

- 1. What do you think the experimental question is?
- 2. What hypotheses can you come up with to answer the experimental question?
- 3. If your hypothesis is correct, what would the plates look like (i.e., what predictions would you make for each hypothesis)?
- 4. Is the experiment you just collected data for an appropriate test of the experimental question you came up with in your answer to Question 1?
- 5. Which antibiotics where most effective against *S. aureus*? Against MRSA?
- 6. When comparing the antibiotics effective against both, were there differences in effectiveness?
- 7. What other questions do the data shown in Figure 1 make you think of?

Part II – Resistance

Among the first antibiotics used on a large scale was penicillin, which was discovered in 1929 by Alexander Fleming. It was finally isolated and synthesized in large quantities in 1943. Penicillin works by interfering with the bacterial cell wall synthesis. Without a cell wall, the bacterial cells cannot maintain their shape in changing osmotic conditions. This puts significant selective pressure on the microbes to evolve, as they cannot survive the osmotic stress. Any microbe that can resist these drugs will survive and reproduce more, making the population of microbes antibiotic resistant.

The specific mechanism of penicillin is the prevention of cell wall synthesis by the β -lactam ring of the antibiotic (Fig. 3), which binds and inhibits an enzyme required by the bacterium in this process.

The enzyme is called penicillin-binding protein (PBP), even though it is an enzyme involved in cell wall synthesis. Normally enzymes have names that indicate what they do and end in the suffix *-ase*, like lact*ase*, the enzyme that breaks down lactose. Figure 4 is a representation of PBP and its active site.

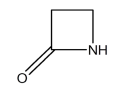


Figure 3. The β -lactam ring common to the penicillin family of antibiotics.

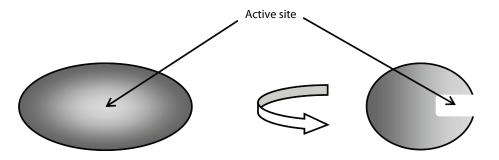


Figure 4. PBP (penicillin-binding protein) active site is a groove allowing formation of cross-links in the bacterial cell wall.

Bacterial cell walls are layered structures, where each layer is made of peptidoglycan, a sugar and protein polymer. Each layer is cross-linked to the next, strengthening the wall and allowing the cell to resist osmotic pressure. The way the enzyme PBP works is to form those cross-bridges by joining strings of amino acids together in the active site, which is a groove in the protein (Fig. 5).

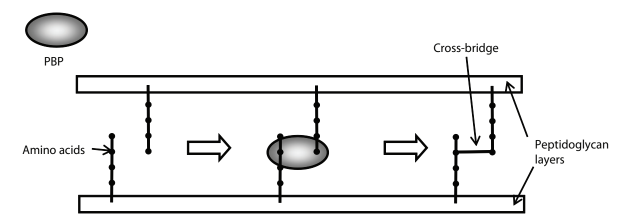


Figure 5. Cross-link formation in bacterial cell walls by PBP (penicillin-binding protein).

The PBP takes amino acid residues attached to peptidoglycan layers and forms bridges between them within the active site groove. This cross-linking, or cross-bridging, stabilizes and strengthens the cell wall. β -lactam antibiotics interfere with the PBP enzyme by binding to the active site, blocking the site from the amino acids (Fig. 6).

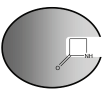
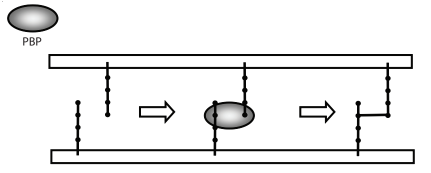
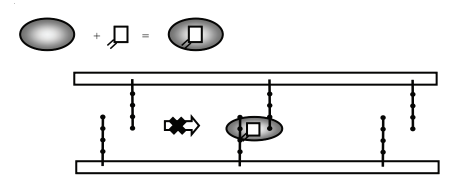


Figure 6. Inhibition of PBP (penicillin-binding protein) by β-lactam blocking the active site.

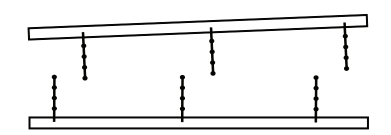
There are over 80 natural and semi-synthetic forms of β -lactam antibiotics, including cephalothin and methicillin. Vancomycin also interferes with cell wall synthesis, but its mechanism of action is to bind directly to the cell wall components (Figs. 7 and 8).



a. Normal PBP binding and cross-bridge formation

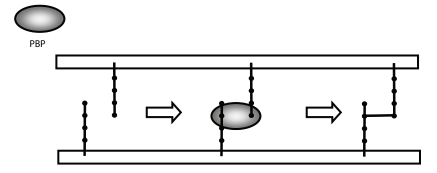


b. PBP inhibited by β -lactam antibiotic

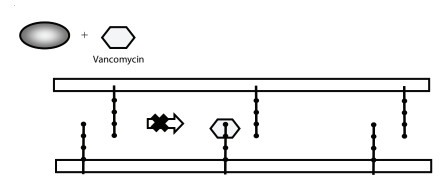


c. Cell wall does not form properly

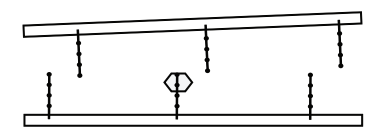
Figure 7. PBP (penicillin-binding protein), the enzyme that allows the bacterial cell wall to form cross-bridges, is inhibited by the β -lactam family of antibiotics. This prevents proper cell wall synthesis and the bacterium will succumb to osmotic stress.



a. Normal PBP binding and cross-bridge formation



b. Vancomycin binds PBP substrate



c. Cell wall does not form properly

Figure 8. Vancomycin inhibits cell wall synthesis a different way by binding PBP's substrates and preventing cross-bridging. This prevents proper cell wall synthesis and the bacterium will succumb to osmotic stress.

The first MRSA case was discovered in 1961 in a British hospital, and was the result of a mutation in the enzyme normally inhibited by the β -lactam ring of methicillin. The site where the antibiotic would bind no longer allowed access to the ring, so the enzyme continued to function normally. The microbe acquired a new gene that, when made into protein, was a different version of PBP, one that couldn't be inhibited by penicillin.

Questions

- 1. Describe what is happening in Figures 7 and 8 in a complete sentence of your own words.
- 2. What are the differences in how β-lactam antibiotics and vancomycin work?
- 3. What other mechanisms might arise to allow resistance to the β -lactam antibiotics?
- 4. Could resistance arise to vancomycin? Why or why not?

Part III – Restoring Susceptibility

Katelyn had been working for Dr. Johnson for a month, and while she had become quite good at measuring inhibition zones, she didn't know why she was doing all this work. She had gotten very curious after she began doing all the measurements on a new set of antibiotics. This experiment involved infecting mice with MRSA and tracking how the MRSA grew over time.

Data were collected by counting the cells of MRSA taken from fluid samples from the mice. The cells were measured by taking one gram of the fluid and spreading it over plates, but now Katelyn counted the colonies that grew on the plate after 24 hours. Because there were so many, she actually measured the colonies as "log CFU/g." A CFU is a colony forming unit, or essentially a cell that will divide into a colony that can be seen. Because there can be so many, Katelyn measured them on a logarithmic (log) scale. The raw data in her lab notebook looked like the following:

		Treatment		
Trial	Control	FtsZ inhibitor	Imipenem	FtsZ inhibitor + imipenem
1	9.11	7.55	6.98	2.21
2	8.25	8.12	8.12	4.55
3	9.05	9.27	9.01	7.98
4	9.37	8.02	8.33	5.64
5	8.80	7.65	7.64	1.25
6	9.25	8.3	7.77	9.98
7	9.41	7.99	8.21	6.78
8	9.11	7.71	7.98	3.45
9	8.61	8.22	7.68	2.45
10	9.12	8.11	8.21	1.01

Table 1. Effect of treatment on MRSA in mice after 24 hours of drug treatment as lo	g CFU/g.
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Questions

- 1. What do you think the experimental question is?
- 2. What hypotheses can you come up with to answer the experimental question?
- 3. What predictions would you make for each hypothesis?
- 4. Looking at the data in Table 1, what do these numbers mean? (Keep in mind a log value means each integer increase is actually a ten-fold increase in the number of cells.)
- 5. What do you think FtsZ inhibitor and imipenem are?

Next, Katelyn further analyzed the data she collected by calculating the average and standard error.

Table 2. Average effect of treatment on MRSA in mice after 24 hours of drug treatment (log CFU/g).

		Treatment		
	Control	FtsZ inhibitor	Imipenem	FtsZ inhibitor + imipenem
Average	9.008	8.094	7.993	4.53
SE	0.114	0.153	0.169	0.954

Question

6. Does Table 2 change your interpretation of the experimental data from Question 4? Why or why not?

She then made the following graph (Fig. 9):

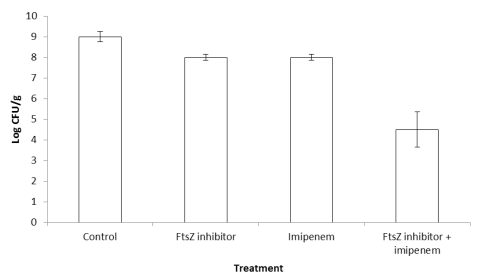


Figure 9. Effects of treatments on MRSA numbers in mice. Samples were taken at 24 hours post-infection. (Figure modified from Tan et al. 2012).

Katelyn was very excited by the results, but she didn't know what an FtsZ inhibitor was, or what imipenem was. She decided to ask Dr. Johnson what his research was all about.

"Dr. Johnson, look at these results I got from the last round of plates," Katelyn said as she handed him a copy of the results above. "What exactly are we testing here?"

Dr. Johnson looked at the results and smiled. "These are great! This could really change the way we deal with antibiotic resistance.

"To answer your question, β-lactam antibiotics are still the most heavily used antibiotics, though resistance is a big problem. Most treatments have changed to using multidrug regimens in the hopes of allowing the antibiotic to still function while at least slowing down the resistance mechanism.

"Another approach involves looking for other proteins that could be inhibited, and looking for existing inhibitors to make into drugs. Instead of looking just for new antibiotics, we're looking for new targets."

Dr. Johnson handed Katelyn a few papers to read. In them she learned that the protein, FtsZ, helps "pinch off" the new cells at the end of cell division. This involves interacting with the cell wall as it is synthesized, and if FtsZ is interfered with, cell wall synthesis stops too. This prevents cell division and the microbe can no longer reproduce.

Dr. Johnson tested the new target idea by using a recently discovered inhibitor of FtsZ to see what effects that had on a MRSA infection. As part of the study, the inhibitor was tested by itself and in combination with imipenem, a β -lactam antibiotic, resulting in the data above.

Questions

- 7. How effective was the FtsZ inhibitor alone? Imipenem alone?
- 8. How effective was the combination of the inhibitor and the β -lactam antibiotic?
- 9. How would you explain these results?
- 10. What questions would you pursue next?

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