Electroporation of M. smegmatis

Materials and Reagents:

- 1. Frozen or freshly made 400µl aliquot of electrocompetent M. smegmatis
- 2. Ice in ice bucket
- 3. Pipettman, 1000µl, 200µl, 10µl
- 4. Sterile pipette tips, 1000μl, 200μl, 10μl
- 5. GenePulser electroporation cuvette with a 0.1 cm electrode gap (Invitrogen Catalog no. 65-0030)
- 6. GenePulser electroporator (Bio-Rad catalog no. 1652660 through 1652668)
- 7. Kimwipes
- 8. Sterile LB broth
- 9. 15ml conical Falcon tubes (VWR catalog no. 21008-918)
- 10. 37°C shaking incubator
- 11. LB plates with appropriate antibiotics
- 12. 37°C incubator
- 13. Plastic bags
- 14. Allegra 6 R tabletop centrifuge
- 15. Autoclave bags

	Autoclave tape Autoclave
Protocol:	
1	Thaw a 400μl aliquot of electrocompetent <i>M. smegmatis</i> on ice.
2	Once thawed, swirl transforming DNA (0.1µg) into the cells and incubate on ice for 5 min.
3	Transfer cells and DNA to a pre-chilled GenePulser electroporation cuvette with a 0.1-cm electrode gap (Invitrogen Catalog #65-0030) and rap smartly against the benchtop in order to draw all cells to the bottom. Return to ice.
4	Flip the power switch on the right side of the GenePulser electroporator to "ON". A menu will appear.
5	Choose "#1 Exponential Protocol" by pushing "Enter".
6	Fill in $1.25 kV$, $25 \mu F$ (capacitance), 1000Ω (resistance), 1mm (electrode gap size) by navigating the menu with the arrow buttons. Once finished push "Enter". A "P" will begin to flash in the bottom right corner of the screen. This indicates that the machine is ready to pulse.
7	Cap the cuvette and wipe all moisture from the outside of the cuvette. Any remaining moisture could cause the machine to arc.
8	Place the cuvette into the safety chamber of the electroporator. Orient the cuvette so that the metal plates are aligned with the electrodes. Push the cuvette into the chamber until it is sealed between the electrodes and the base of the chamber. Close the lid on the safety chamber.
9	Press and hold the large red "Pulse" button until the machine beeps and the screen changes to show your exponential curve.
10.	Record the time constant and voltage applied. A time constant of 1.0-10s is optimal. A voltage of $1000\text{-}1500\Omega$ is optimal.
11.	Immediately transfer the cuvette to ice and gently resuspend the cells in 1 ml of sterile LB. The number of transformants declines as the cells remain without media.

12 Transfer the cells and media to a sterile 15ml conical Falcon tube (VWR catalog no. 21008-918). Loosen the cap on the tube to allow proper aeration and secure with tape to prevent it being shaken off.
13 Place the tube in the 37°C shaking incubator for 4 hours. This allows the cells to begin expressing antibiotic resistance genes.
14 Warm two LB-antibiotic plates to room temperature.
15 Spin down the cells in an Allegra 6 R tabletop centrifuge for 10 min at 3000 rpm, 4°C. Pipette off supernatant. Resuspend cells in 500 μl sterile LB.
16 Plate 50μl of resuspension on one of the plates and 100μl on the other. Allow plates to dry, invert, and place in plastic bag. Place this in the 37°C incubator. Flat, crinkled, white colonies should appear in 3-4 days.
17 Autoclave and discard all wastes generated, including the electroporation cuvette.