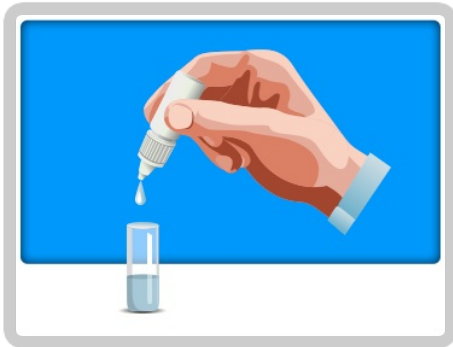
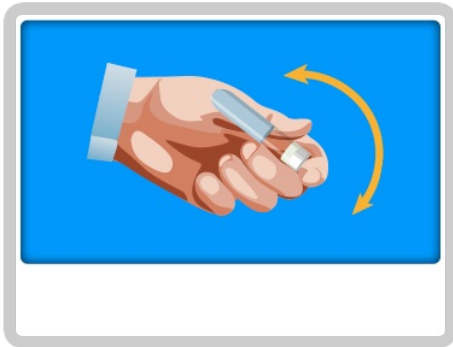


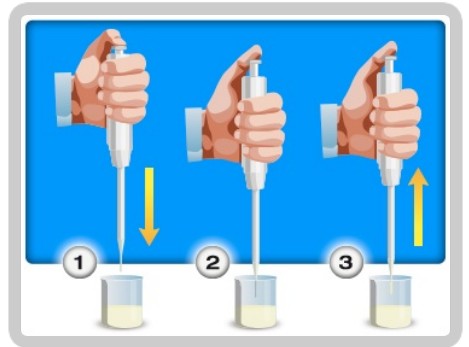
PROCEDURE



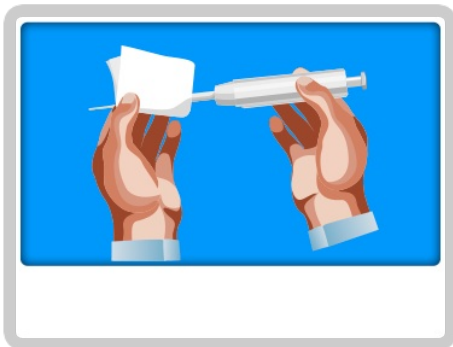
1. Add 1 drop of R1a to the cuvette containing R1 and shake.



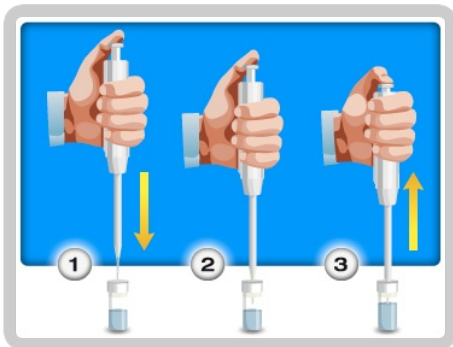
2. Homogenize the sample before collection.



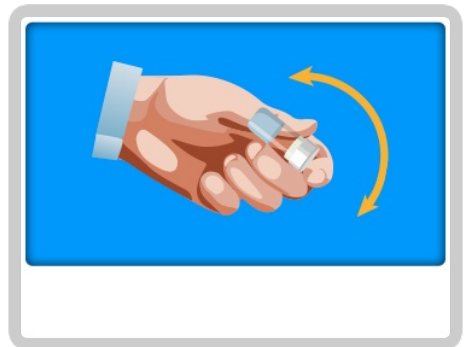
3. Collect 100 μ L of sample with a pipette. Use a new tip for each test to prevent contamination from previous samples.



4. Carefully clean the outside of the pipette tip with blotting paper, avoiding contact between the extremity of the tip and the paper.



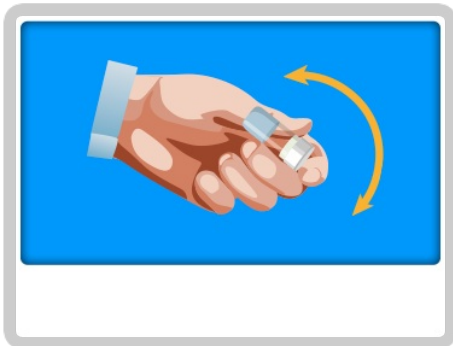
5. Place the sample in the cuvette. Keeping the tip immersed in reagent, press and release the piston of the pipette several times.



6. Gently shake the cuvette 2-3 times.



7. Place the cuvette in one of the incubation cells and start the timer by pressing the timer icon.



8. Gently shake the cuvette 2-3 times.



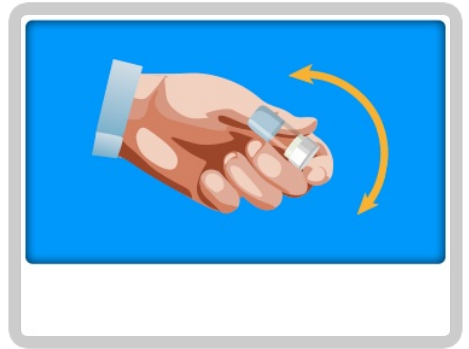
9. Select the cuvette to be read from the display. Place the cuvette in the cell marked with the blue light and press READ.



10. Add 1 drop of R2 to the cuvette containing R1 + sample and shake.



11. Place the cuvette in one of the incubation cells and start the timer by pressing the timer icon.



12. Gently shake the cuvette 2-3 times.



13. Place the cuvette in the cell marked with the blue light and press READ to perform the photometric reading.