

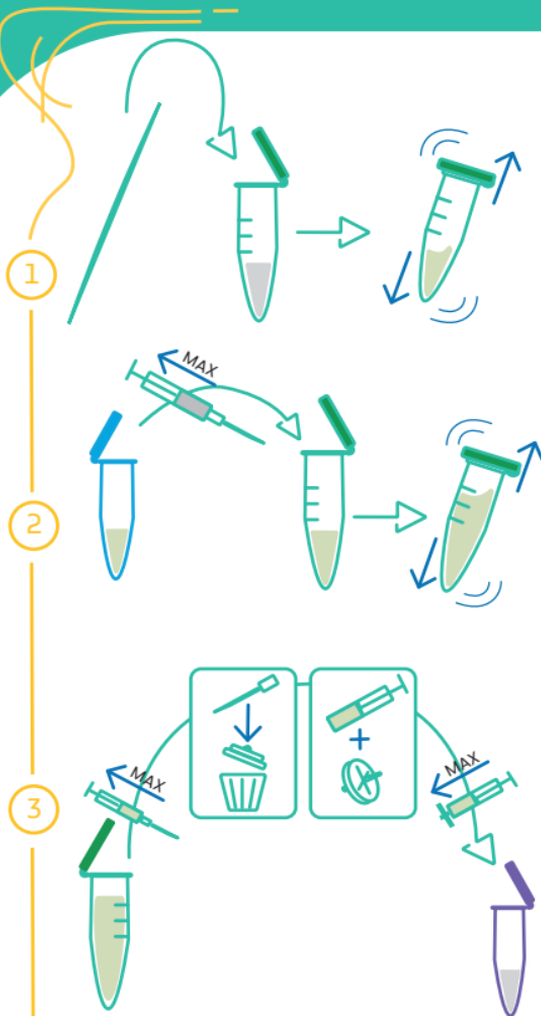
RAPID USER GUIDE



Switch on and program the instrument.



Before opening the tubes, make sure that all the liquid and lyophilized pellets are at the bottom of the tubes.



Take the equivalent of half a grain of semolina from faeces using the **pick** provided, then immerse and spin the **pick** in the **green tube**.

Shake the **green tube** vigorously.

Use the **syringe** and **needle** to transfer entirety of liquid from the **blue tube** and dispense it in the **green tube**. Shake vigorously.

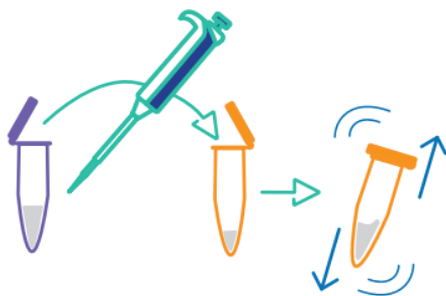
With the same **syringe** and **needle**, draw the entirety of the **green tube**.

All the foam formed previously does not need to be drawn.

Remove the **needle** and fit the supplied **filter** on the **syringe**.

Dispense the filtrate into the **purple tube** carefully.

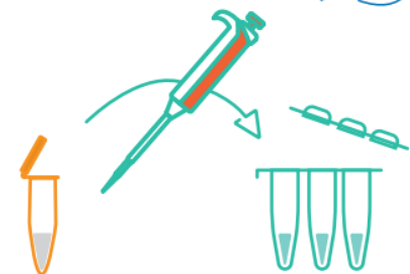
4



Draw liquid from the **purple tube** with the **blue pipette** fitted with filter-tip.

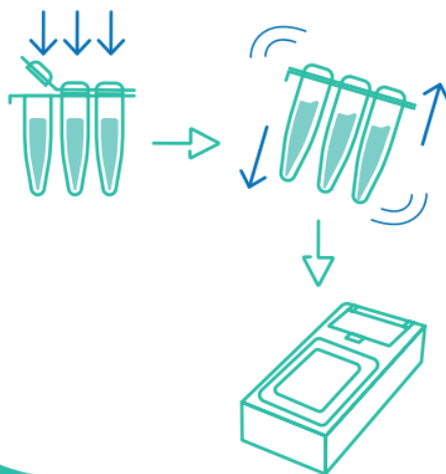
Transfer the liquid to the **orange tube** then shake it vigorously.

5



Transfer liquid from the **orange tube** with the **orange pipette** fitted with filter-tip and dispense it in each **minitube**.

6



Replace caps tightly then shake the **minitubes** to ensure the pellets have dissolved and no air bubbles are present.

Insert the **minitubes** into the **instrument** (place the end tab on the left) then immediately close the lid of the **instrument** and press ok to start the analysis.



Never reopen the minitubes once they have been inserted in the instrument, or after the end of the reaction.

Never leave the minitubes inside the instrument after the reaction has ended.

Refer to the user manual for more detailed information:

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