CompactDry “Nissui” AQ Illustration Manual
Weigh 50g solid sample and add 450mL Buffering Solution to the sample. It is recommended to use a stomacher bag with filter to eliminate risks of carry over of tiny pieces of foodstuffs into the surface of the medium.

Homogenize this mixed sample by a blender
Open aluminum bag, and take out a set of 4 plates.
Take off the cap of the plate

Write the appropriate information on the memorandum section.
Pipette 1ml of homogenized specimen (to be further diluted if necessary) in the middle of dry sheet of Compact Dry AQ.

Specimen diffuses automatically and evenly into all over the sheet (total medium of 20 cm²) to transform it into gel within seconds.
Inoculate 1 ml of wiping solution (to be diluted if necessary), which is obtained from cotton swab,
Turn over the plate capped put in an incubator.
Incubate for 44 + 4 hours at 36 + 2 °C.
Another plate is incubated for 68 + 4 hours at 22 + 2 °C.
Colonies grown CompactDry AQ are almost red colored.

Detection limit of Compact Dry AQ is between 1 – 300 cfu/plate.

From backside of the plate, count the number of colored colonies appeared in the medium. White paper placed under the plate can help to count colonies easier.