

FOR FAECAL CONCENTRATION OF

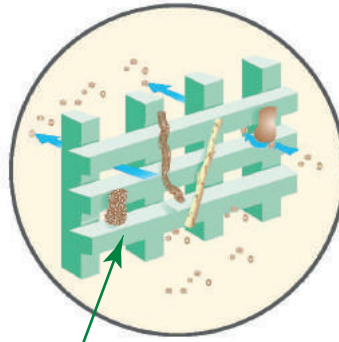
HELMINTH OVA AND LARVAE / PROTOZOA CYSTS AND OOCYSTS

APACOR

Mini Parasep® SF
FAECAL PARASITE CONCENTRATOR



MESH DETAIL



SELF STANDING
SAMPLE CHAMBER

MIXING CHAMBER

INTEGRAL
SPOON

Filter

A two stage filtration matrix. Large particles are rejected without occluding the 425µm pores. Recovery rate with Parasep® is comparable to traditional sieve method, ie: Ridley-Allen. The vertical filter enclosed design is patented.

Debris Trap

Rejected particles are trapped to prevent extrusion into the Sedimentation Cone during centrifugation.

Air/Liquid Seal and Safety Lock

The 'seal' prevents the release of biohazardous material. The 'lock' ensures the Mixing Chamber and Filter are removed together for safe disposal.

Fat Dispersion Chamber

A perforated fat dispersion chamber removes the smaller faecal debris and separates the fat content so that it can be efficiently removed from the resulting sediment without the use of ether or ethyl acetate.

Sedimentation Cone

Sediment forms in the base of the cone allowing examination for the presence of helminth eggs or larvae and protozoa cysts or oocysts.

Health and Safety Benefits

- Totally enclosed/sealed process
- Reduced reagent volumes
- No cleaning required
- Single use, no sample contamination
- Ready to use systems available

Performance Benefits

- Optimum sample recovery
- Enhanced sample clarity
- Rapid four step process
- Human resources optimised
- Easy patient identification
- Fits all 15ml centrifuge buckets



PARASITOLOGY

SINGLE USE IN VITRO DIAGNOSTIC DEVICE



FAECAL PARASITE CONCENTRATOR

Mini Parasep® SF

Procedure

STEP 1 - SAMPLE PREPARATION

Fresh Samples

For empty Parasep®, unscrew lid and add 3.3ml of fixative and one drop of surfactant (eg: Triton X-100) to the mixing chamber.

Alternatively use the reagent ready Mini Parasep® SF.

Introduce a scoop of faecal sample using the spoon on the end of the Mini Parasep® SF filter.

Mix in thoroughly with the Mini Parasep® SF spoon. If the sample is hard, break it up with the end of the spoon.

Preserved Samples

Shake or vortex the incoming preserved sample to thoroughly mix. Transfer 0.3ml = 0.3g of sample into the Mini Parasep® SF mixing chamber. Add 3.0ml of 10% Formalin/water plus 2 drops of Triton X.

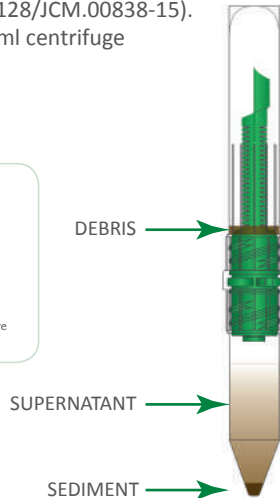
STEP 3 - CENTRIFUGATION

Invert the Mini Parasep® SF and centrifuge at 200g for two minutes or 400g for two minutes (J. Clin. Microbiol. doi:10.1128/JCM.00838-15). Mini Parasep® SF fits all 15ml centrifuge buckets.

NOTE: TO CALCULATE THE REQUIRED RPM FOR ANY CENTRIFUGE.

$$RPM = \sqrt{\frac{g}{1.12r}} \times 1000$$

RPM - rotor speed in revs/min.
g - centrifugal force (max 1000g)
r - radius, horizontal distance between sedimentation cone tip and spindle centre measured in mm.



STEP 2 - EMULSIFICATION

Seal the Mini Parasep® SF by screwing in the filter/ sedimentation cone unit.

Vortex or shake to emulsify with the sedimentation cone pointing upwards.

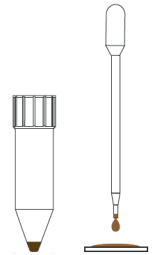


STEP 4 - EXAMINATION

Direct Method

Unscrew and discard the filter and mixing tube. Pour off all the liquid above the sediment.

Pipette one drop of saline or Lugol's Iodine solution onto a slide, add one drop of deposit to the saline or Lugol's Iodine, mix sample and cover with cover-slip.

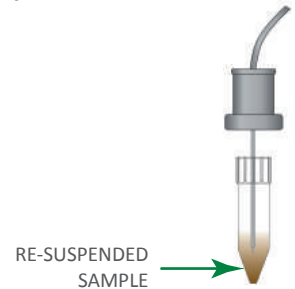


OR

Semi-automated System - ParaSys™

Unscrew and discard the filter and mixing tube. Pour off all the liquid above the sediment.

Press 'Dilute' to add Saline to the sediment. Shake or vortex to re-suspend sample. Insert Aspirator into suspension and press SAMPLE to draw 100µl into the ParaSlide™. (Refer to ParaSys™ instruction manual).



See label for storage conditions and expiry date. Please adhere to the following guidelines when handling Mini Parasep® SF. To avoid cross contamination the Mini Parasep® SF device should remain closed at all times except when introducing the sample or when retrieving the final concentrated sample for examination.

Mini Parasep® SF is available empty or reagent ready
Please ask for details

Products can be ordered direct from Apacor or from an appointed distributor
Visit our website for all the latest information www.apacor.com or email on: sales@apacor.com



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