



Apr 21, 2020

COPAS wormsorter

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Works for me

dx.doi.org/10.17504/protocols.io.bfc9jiz6

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ABSTRACT

Protocol for dispensing adult worms using the COPAS 500 flowpilot

Prepare equipment

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COPAS wormsorter indicating key components

Turn on the compressor at the wall – it should show a pressure of 40psi after switched on


- 2 Turn on COPAS machine with switch on the left hand side
- 3 Turn on the lasers (488 laser sufficient if using unmarked animals). Add in picture of lasers.
- 4 Turn on the computer
- 5 Discard waste contents that are in the recovery cup (small shallow cup on the left-hand side of the machine)
- 6 Check that there is water in the sheath. If the water is low, fill up with MQH₂O (not M9).
- 7 Make sure that the recovery cup and sample cup are securely tightened so that there are no leaks in the system


Prepare software

- 8 Open dbgvie – should always be running in the background
- 9 Open FlowPilot software and a prepared experiment with a set gate for eg Adults. :
- 9.1 File -> Load Experiment
- 9.2 File -> Load sample

Clean system

- 10 Maintenance -> Flush Sample
- 11 Click 'Refill Sample' – the sample cup pressure should decrease. You can see this in the software on the left hand size (include screenshot).

 Sometimes the sample cup pressure doesn't decrease and in fact increases. You can still unscrew the sample cup but if this persists there may be a blockage.
- 12 Unscrew sample cup and replace with falcon filled with cleaning solution (pink in colour)
- 13 Once securely replaced click 'Done refill'
- 14 Check 'Sample on' and 'mixer on' – cleaning solution should now pass through the system; allow a 2-3 ml to pass through (make sure sheath is unchecked)

 You will get a warning about contaminating the flow cell, this normal and you can click 'Yes'
- 15 Uncheck 'Sample on' or click Abort to stop sample flow.
- 16 Repeat steps 11-15 with water

Load sample

- 17 Repeat steps 11-13 with sample.
- 18 **Turn mixer ON.** If you do not do this you may lose all your worms that have settled to the bottom of the tube!!!
- 19 Maintenance -> Prime Flow Cell; to flush sample through the system and remove air bubbles
- 20 Maintenance -> Flush sample
- 21 Check 488nm (and 568nm) laser boxes
- 22 Check 'Use sort gate' for stored sort gate – include screenshot of software here

- 23 Click 'Acquire' – sample should pass through the system and number of events per second will be shown:
- Aim for 10-20 events per second
 - If too few/too many events increase/decrease 'Sample cup pressure' so that it is between 1.5-2psi
 - To ensure only one event per droplet go to Setup->Coincidence, select 'Pure, no double'. This increases accuracy in the number of worms dispensed but the time to dispense may increase.

Test dispensing

- 24 Click on the plate icon on the top bar
- 25 Select number of objects to sort
- 26 Select the wells you would like to fill (for testing we use a spare 60mm plate and fill wells A1, A2, B1, B2)
- 27 Select which gate to use
- 28 Apply
- 29 Place 60mm plate in front left corner of left-hand stage with A1 in the left corner.
- 30 Click 'Fill plate'
- 31 Keep an eye on the number of events per second
- 32 Ensure the 'Diverter pressure' is checked
- 33 Check under microscope that the correct number of objects were dispensed per 'well'
- 34 If too many objects, decrease sample cup pressure and repeat steps 8-11 or select Pure no double to increase accuracy.

Fill plate

- 35 Click on the plate icon on the top bar
- 36 'Clear plate'
- 37 Select number of objects per well and click 'Apply to All' or select which wells you would like to fill.
- 38 Apply
- 39 Place 96 well plate in left-hand stage
- 40 Ensure 'Diverter pressure' is checked'; if it is not then liquid comes out of the dispenser constantly and you get flooding.
- 41 'Fill plate'
- 42 Keep an eye on the number of events per second still and monitor how much sample fluid is coming through the system

Clean system

- 43 Repeat steps 11-15

- 44 Keep sample cup with water secured so that the system is air-tight and closed
- 45 Turn off all equipment (Computer, lasers, compressor, worm sorter).



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