# Coordination to Support Fisheries Management in the Western and Central Mediterranean. CopeMed Phase II COPEMED



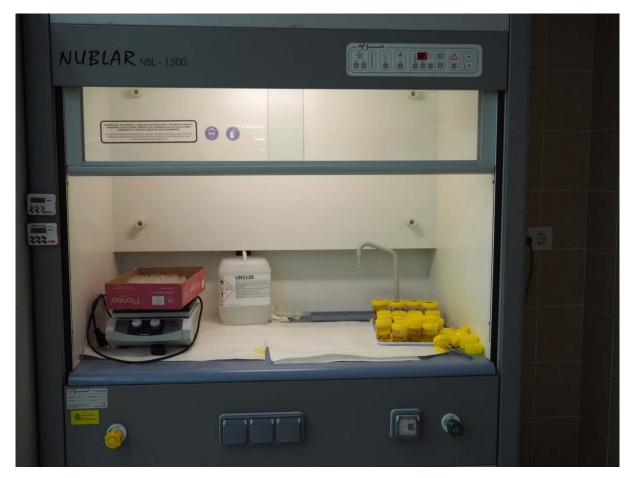
### TRAINING COURSE ON ICHTHYOPLANKTON

# Laboratory equipment, sample preparation, ichthyoplankton sorting, preserving and storing fish eggs and larvae

by

J.M. Rodríguez and A. García

Fuengirola (Málaga), Spain 22-26 February 2016 Spanish Institute of Oceanography (IEO) The first step in handling ichthyoplankton samples is to eliminate the preserving liquid, in most cases formalin. For this we need a **fume hood** 



This allows us avoid to respire the formalin steam that is highly carcinogenic

#### We follow:

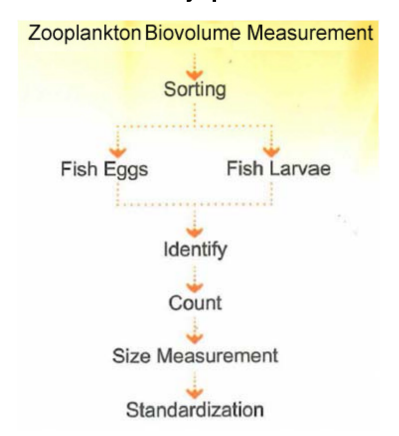
- Draining samples through a net (mesh size the same as the plankton net)
- •Returning the preserving liquid to the original sampling jar

The above two steps are done inside the **fume hood** 

- Washing the dry plankton sample with fresh water
- •Pouring the free formalin plankton sample in a container with fresh water
- Stirring the solution softly with a glass rod
- Pouring a small quantity of the plankton sample into a petri-dish
- Sorting fish eggs and larvae from the petri-dish under a dissecting microscope at a magnification of about 10x
- •Placing eggs and larvae in separated and labelled petri-dishes
- •Precaution must be taken to prevent fish egg and larva damage when handling them
- Counting the number of fish eggs and larvae
- Recording the total number of fish eggs and larvae removed from the sample
- •Storing fish eggs and larvae in separate labelled vials with 70% ethanol\* or a solution of 5% formalin and fresh water
- •The label of each vial, must include information on sampling date, station number, sampling site, sampling method (oblique, horizontal or vertical tow), and the plankton net used

<sup>\*</sup>Eggs must be not preserved with ethanol because it dehydrates and wrinkles eggs, becoming not identifiable

### Laboratory procedure



## Zooplankton biovolume measurement

(the biovolume is an estimate of the zooplankton biomass)

- Remove non-plankton organisms, such as adult / juvenile fish and large plankton organisms (individual volume > 5 cm<sup>3</sup>) such as jelly fish and tunicates
- Determine the total volume
- Remove the preserving liquid by filtering through a mesh of the same size that that of the plankton nets
- Determine the volume of the removed preserving liquid
- Total volume preserving liquid = biovolume of zooplankton
- Record the biovolume
- Return the sample to the original preserving liquid

### The standardization

Fish egg and larva counts can be standardised either, to number of individuals per unit of volume (m³, 10 m³, 100 m³ or 1000 m³) or to number of individuals per surface unit (generally 10 m²). The formula for calculating the number of individuals per surface unit is:

no. of egg/larvae x 10 square meters = 
$$\frac{n.p}{v}$$
. 10

Where n is the eggs or larval counts, p is the depth reached by he net during the sampling and v is the volume of water filtered by the net

# **Size measurement**

Image capture

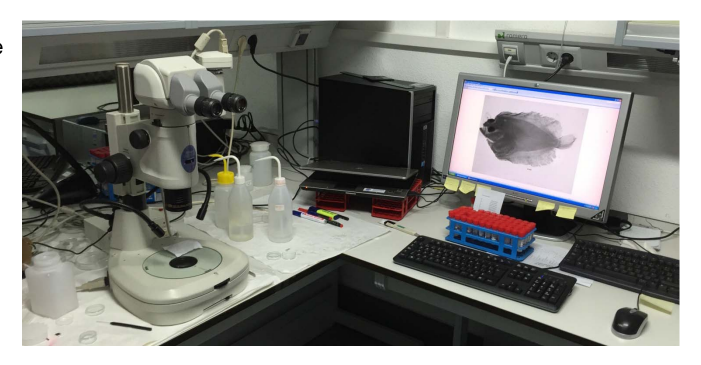


Calibration of microscope magnification



Image measurement

To capture an image we need a digital camera connected to a trinocular stereoscopic microscope and to a computer



The **calibration factor** tells us how many pixels correspond to a millimetre for each microscope magnification

To calibrate microscope magnification we use use graph paper

Calibration Factor		
5x	66,40	pix/mm
10x	102,20	pix/mm
15x	153,80	pix/mm
20x	206,60	pix/mm
25x	257,75	pix/mm
30x	316,33	pix/mm
35x	359,67	pix/mm
40x	414,50	pix/mm

Calibration Easter

For egg and larval fish measurements we use of the free software **IMAGEJ 1.45s** (available at http://imagej.nih.gov/ij/)

Now we are going to show you how to measure a fish larva with the **IMAGEJ 1.45s**